Quantification of Adventitial Vasa Vasorum Vascularization in Double-injury Restenotic Arteries

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Abstract

Background: Accumulating evidence indicates a potential role of adventitial vasa vasorum (VV) dysfunction in the pathophysiology of restenosis. However, characterization of VV vascularization in restenotic arteries with primary lesions is still missing. In this study, we quantitatively evaluated the response of adventitial VV to vascular injury resulting from balloon angioplasty in diseased arteries.

Methods: Primary atherosclerotic-like lesions were induced by the placement of an absorbable thread surrounding the carotid artery of New Zealand rabbits. Four weeks following double-injury induced that was induced by secondary balloon dilation, three-dimensional patterns of adventitial VV were reconstructed; the number, density, and endothelial surface of VV were quantified using micro-computed tomography. Histology and immunohistochemistry were performed in order to examine the development of intimal hyperplasia.

Results: Results from our study suggest that double injured arteries have a greater number of VV, increased luminal surface, and an elevation in the intima/media ratio (I/M), along with an accumulation of macrophages and smooth muscle cells in the intima, as compared to sham or single injury arteries. I/M and the number of VV were positively correlated ($R^2 = 0.82, P < 0.001$).

Conclusions: Extensive adventitial VV neovascularization occurs in injured arteries after balloon angioplasty, which is associated with intimal hyperplasia. Quantitative assessment of adventitial VV response may provide insight into the basic biological process of postangioplasty restenosis.

Key words: Angioplasty; Micro-computed Tomography; Restenosis; Vasa Vasorum

Introduction

Percutaneous transluminal angioplasty is important for the management of occlusive atherosclerotic lesions in humans. However, restenosis following angioplasty occurring in up to 50% of patients limits the long-term effectiveness of this treatment. Postangioplasty restenosis pathophysiology has not been well-defined. Early studies to elucidate the mechanisms of restenosis focused on intimal de-endothelialization as a primary cause, while more recent studies indicated a potential role for adventitial vasa vasorum (VV) in the initiation and/or progression of atherosclerotic lesions and vessel restenosis. Adventitial VV is a network of microvasculature providing oxygen and nutrients to the outer layers of the arterial wall. VV disruption may result in impaired oxygen transformation, vessel wall hypoxia, accumulation of oxidized metabolites, and nutritional deficiencies in the vessel wall. VV can also serve as conduits for the recruitment of inflammatory cells, including macrophages, and noncellular inflammatory components. These effects can lead to angiogenic and mitogenic factor expressions, including adhesion molecules and enzymes, in turn leading to smooth muscle cell (SMC) migration/proliferation, neointimal formation, vascular remodeling, and restenosis following angioplasty. Even though quantification of VV vascularization is reported in undiseased arteries following balloon overstretching responses of adventitial VV to angioplasty in severely diseased arteries, where angioplasty is normally performed in human patients, have not been examined or the relationship of these microvessels to changes in the intima. In the present study, we utilized a double-injury rabbit model: Primary lesions induced in carotid arteries by perivascular manipulation and balloon injury after 4 weeks. High-resolution three-dimensional (3D) volumetric data are suitable for the visualization and quantification of the entire VV microvasculature. Micro-computed tomography (CT) has emerged as the preferred method for this purpose. In the present study, we used micro-CT combined with histological and immunohistochemistry methods to quantify the responses of VV and assess whether there is an association with neointimal formation in experimentally injured arteries following balloon dilation.
Methods

All animal procedures were reviewed and approved by the Institutional Animal Care and Use Committee at Shanghai Jiao Tong University medical division. Male New Zealand white rabbits (2.0–2.5 kg; 10–14 weeks old) were used in the present study. Two weeks prior to the first surgery, high cholesterol diet (2% cholesterol + 6% peanut oil + 92% normal chow diet) was begun in all animals, which was discontinued on the day of the second surgery (angioplasty) as previously described. Animals were divided into three groups in order to demonstrate the various stages of vascular lesions: Group 1, sham operation group (n = 6) where animals underwent a sham operation; however, the artery was not impaired; Group 2, single-injury group induced by a constrictive thread loop (n = 9); and Group 3, double-injury group induced by angioplasty (n = 10).

Induction of the focal lesion in the right common carotid artery rabbits were anesthetized with 3% pentobarbital sodium using ear vein administration. Through a medial cervical incision, right common carotid arteries were exposed, and a 5–0 antibacterial (polygalactin 910) suture (Ethicon Ltd., Scotland, UK) was placed around the artery. Prior to ligation of the thread, a sterile metal tube with an external diameter of 0.7 mm was inserted in the suture collar parallel with the carotid artery, followed by withdrawal of the tube. Ampicillin (50 mg/kg) was administered intramuscularly (i.m.) immediately following surgical palliation. Carotid angioplasty 4 weeks after focal carotid lesion induction, the right carotid artery was exposed, and the thread collar was found to be nearly completely absorbed in both single-injury and double-injury groups. Carotid angioplasty was performed in the double-injury group. Two vascular clips were placed on the carotid artery in order to prevent bleeding; one was placed onto the proximal common carotid artery, and the other was placed onto the internal carotid artery. A standard 3.0 mm × 17 mm percutaneous transluminal coronary angioplasty balloon was inserted in a retrograde fashion via the external carotid artery and was positioned at the point of ligation. For dilation, the balloon was inflated to six atmospheres for 2 min and then deflated and retracted, followed by ligation of the external carotid artery. All rabbits received a bolus of 100 μg/kg heparin intravenously (i.v.), and ampicillin (50 mg/kg) i.m.

Polymer injection and specimen dehydration after 4 weeks, all animals were sacrificed using a fatal dose (>100 mg/kg) of pentobarbital sodium, i.v. The right carotid was cannulated using a 2-mm-diameter plastic catheter. In order to clear the remaining blood from the carotid artery, we infused 500 ml of heparinized saline (0.9% sodium chloride containing 10,000 units of heparin) at 100 mmHg. We then infused low viscosity, radiopaque polymer compound (Microfil® MV-122, Flow Tech, Carver, MA, USA) through the catheter until the injected effluent flowed freely from the collateral common carotid vein. The tissues from each region were then removed from the neck and immersed in 10% neutral buffered formalin at 4°C overnight to polymerization occur. On the following day, the carotid artery was placed in 95% alcohol for 48 h. At 24-h intervals, the glycercin concentration was raised to 30%, 50%, 75%, and finally 100% glycercin in order to completely dehydrate the carotid segments. The specimen was then rinsed with acetone, left at room temperature for 24 h, and embedded in a paraffin mold for 3D micro-CT imaging.

Micro-CT imaging and reconstruction to preserve VV connectivity, the arteries were scanned in 2-cm increments along the arterial lumen axis without physically cutting the carotid artery. In the present study, the micro-CT scanner was configured so that the dimension of the cubic voxels was 20–25 μm (16-bit grayscale). The 3D images (an average number of 1000 slices per 2-cm carotid artery segment) were imaged using a high-resolution (8–36 μm isotropic voxel size) micro-CT imaging system (μCT80, Scanco Medical; Bassersdorf, Switzerland) and analyzed using the accompanying software.

Morphometric analysis

All specimens were traced, and cross-sections were analyzed at 2 mm intervals (excluding branching points) in order to yield 8–10 micro-CT cross-sections per 2-cm specimen. Vessel wall boundaries were defined using a radius of twice the distance from the arterial lumen to the outer adventitia. Blood vessels within this boundary were determined as VV. VV area within the vessel wall area was differentiated from nonvascular structures by setting the lower threshold values for an intensity range of interest that yielded the best identification of VV regions.

Two anatomically different types of VV have previously been defined. First-order VV originated from the main carotid arterial lumen and ran longitudinally to the carotid artery. Second-order VV were smaller, and arose from branches of first-order, forming circumferential arches around the vessel wall. VV were manually counted and measured, and yielded the following morphometric variables for each cross-section: (a) VV density (VV per mm² vessel wall area) was calculated manually counting the VV and dividing by total area of the vessel wall; (b) the ratio of the number from second- to first-order VV; (c) the ratio of VV luminal volume (Volvv) to the total volume of contrast agent (Voltotal) within the arterial wall. VV luminal volume obtained was manually traced and measured on each cross-section, and (d) VV luminal surface fraction (the sum of VV luminal surface areas per mm² VolCA, mm²/mm³). The VV luminal surface area was calculated using the formula of the side surface area of a cylinder (2×π×radius×height).

Histological and immunohistochemical staining following micro-CT reconstruction and analysis, the specimens were immersed in 40°C water for 4 h to gently melt the wax embedding, followed by removal from the plastic mold and were cut every 2 mm. The sections were stained using Hematoxylin and Eosin (H and E) and elastic van Gieson...
staining. The sections were then analyzed in order to
determine the minimum luminal area (narrowest segment)
for each carotid artery. This narrowest luminal area was at
the level where the ligature was placed. Once the section
with the smallest lumen from each vessel was determined,
morphometric measurements for luminal, intimal, medial
and total vascular areas were performed using digitized
images of the van Gieson stained sections with a KS400
software package (Axioplan 2 imaging system).

In addition, immunohistochemical analysis was performed.
A primary monoclonal antibody specific for vascular
α-actin (MAB 1522, Chemicon International Inc.) was used
at a dilution of 1:400 for the evaluation of vascular SMCs.

For macrophage detection, a monoclonal antibody specific
for rabbit macrophage CD68 (RAM 11, Dako) was diluted to
1:6500. The immunohistochemistry protocol has previously
been described.[20] Briefly, sections were incubated with the
primary antibody, anti-mouse IgG secondary antibody (biotin
conjugate) and avidin peroxidase. The peroxidase was then
visualized using chromagen. Sections were counterstained
using H and E. A negative control was performed using primary
antibodies replaced by mouse IgG isotype control in
both α-actin and CD68 staining. The gray area of the stained
section was analyzed. The unstained area was categorized
as unclassified cell areas consisting of extracellular matrix
and unstained cells.

Statistical analysis
All data are presented as mean ± standard error (SE). F-test
was utilized to test for equality of variances among samples.
One-way ANOVA, followed by a Tukey–Kramer post-hoc

test with correction for multiple comparisons was used to
identify statistical differences among the groups. Individual
group comparisons were performed using an unpaired
Student’s t-test. Correlations among the continuous variables
were analyzed by linear regression. \( P < 0.05 \) was considered
as statistically significant.

Results
Technical outcomes
All operative techniques were well-tolerated as suggested
by normal food intake and normal movement. Three animals
succumbed during this study. Because of the failure of wire
traversing, two occluded arteries after primary injury were
excluded from further study. Of the 30 rabbits initially
studied, analyses were performed on 25 animals: 6 rabbits
in sham operation group, 9 rabbits in primary lesion group
and 10 rabbits in the balloon angioplasty model group.

Micro-computed tomography imaging
The carotid artery with a primary lesion showed a tendency
for new vessel formation compared to sham, especially
second-order VV with a shift toward second- to first-order
VV [Figure 1 and Table 1] while no significant differences
in VV luminal surface fraction or the ratio of VolvTo
Voltotal were detected in these two groups [Table 1]. Four
weeks following angioplasty, extensive neovascularization
of the adventitial VV with a profound shift in the ratio of
second- to first-order VV was detectable, combined with
a further decline in VV density [Figure 1 and Table 1]
compared to sham. The VV luminal surface fraction and
the VolvTo Voltotal ratio in the double-injury group were
significantly elevated, compared to the sham and primary
injury groups [Table 1].

Microscopic analysis
Animals subjected to constrictive thread placement
showed mild formation of neointimal hyperplasia
compared to sham. Morphometric analyses exhibited a
significant elevation in intima/media ratio (I/M) 8 weeks
following primary lesion [Figure 2a and 2b and Table 2].
At 4 weeks following balloon angioplasty, extensive
intimal hyperplasia was developed with a high rate of
I/M [Figure 2c and Table 2]. The thickened neointimal
exhibited invasion of macrophages and accumulation of
SMCs. The number of macrophages was mildly elevated in
the single injury group (81.623 ± 13.240 vs. 70.944 ± 7.223
Grey/mm\(^2\), \( P = 0.043 \), Figure 2d and e), coinciding with an
increase in α-actin positive SMCs (228.933 ± 81.250 vs.
171.342 ± 28.52 n/mm\(^2\), \( P = 0.037 \), Figure 2g and
2h) compared to sham. Following angioplasty, both
macrophages (123.402 ± 5.121 vs. 70.944 ± 7.223 Grey/mm\(^2\),
\( P = 0.014 \), Figure 2f) and SMCs (352.167 ± 15.942 vs.
171.342 ± 28.523 n/mm\(^2\), \( P = 0.012 \), Figure 2i) were
significantly elevated.

Neointimal hyperplasia in histological sections with
adventitial VV neovascularization assessed using
cross-sectional micro-CT images demonstrated a positive
correlation between I/M ratio and the number of VV in
the angioplasty group \( (R^2 = 0.82, \ P < 0.001, \text{Figure 3}) \).
Adventitial VV density declined in both the single- and
double-injured groups [Table 1] as compared to sham.
However, no significant correlation was observed between
the I/M ratio and VV density.

Discussion
This study was designed to identify the response of
adventitial VV to acute vascular injury following angioplasty
and examine potential associations with neointimal
formation. Adventitial neovascularization and extensive
intimal hyperplasia, characterized by accumulation of
macrophages and SMCs, were detected using micro-CT and
histology/immunohistochemistry in injured carotid arteries
following balloon injury.

A rabbit model is commonly used to study responses to
clinical angioplasty in a severely diseased artery.[20] In order
to reproduce the morphological characteristics observed in
patients who have undergone angioplasty, an initial injury
must be induced in order to develop constrictive lesions,
which will undergo angioplasty. The injury methods can
include balloon injury,[21] wire loop,[22] air desiccation,[23] or irradiation.[24] Recently, perivascular
manipulation of the vessel wall using either placement
of constrictive or nonconstrictive cuffs around the vessel or temporary ligation of the vessel rapidly induced a focal atherosclerotic-like lesion, comprised of morphological changes in human atheroma. These alterations included macrophage and SMC infiltration into the arterial subendothelium, foam cell, cholesterol cleft, necrotic core, or fibrous capsule formation.

In the present study, we utilized a two-step injury model of the rabbit carotid artery, featuring a combination of a constrictive loop induced primary lesion and secondary balloon dilation. Manipulation of the outside of the rabbit right carotid artery accompanied by administration of a high cholesterol diet yielded mild intimal hyperplasia with increased densities of macrophage and SMC and decreases in lumen area resembling features seen in early human atherosclerosis. Following the second injury that occurred after balloon dilation, further remodeling of the carotid artery was evident with acute luminal dilatation, plaque fracture and extensive neointimal hyperplasia, reproducing the results of balloon angioplasty in human diseased vessels. We did not utilize the left carotid artery in the same animal with no balloon dilation as control, since after infusing and removing one side of artery for immediate fixation, it is difficult to maintain equal pressure to infuse the other artery. Lower pressure led to compromised resolutions in the micro-CT images and a reduced number of VV in the second infused artery, even following the first and second infused arteries in the same condition (data not shown). To avoid this bias, we used the first infused right carotid artery as our study vessel and compared across all groups.

Kwon et al. reported results of arterial microcirculation following balloon injury in undiseased pig coronary arteries.
These investigators observed that the number of VV vessels, especially in second-order VV, was increased in the injured vessel. The density of newly formed vessels was also increased, correlating with vessel stenosis. In our current study, we observed a similar response of VV neovascularization with significantly increased second-order VV in experimentally injured carotid arteries after balloon angioplasty. New VV vessels were proportional to intimal hyperplasia. Notably, mild VV neovascularization and neointimal hyperplasia were detected in carotid arteries following primary injury. These vessels may serve as further neovascularization following angioplasty-induced injury.

Despite increases in the total number, we have shown that the density of newly formed VV was decreased in injured arteries following balloon dilation. The difference in VV density between Kwon et al. and our studies may likely reflect difference in the stage of restenosis. In the present study, we evaluated restenosis at an early phase, as indicated by a lack of the severe intimal hyperplasia, the surface of the lumen was larger than sham. It is possible that the 4-week time point is not sufficient and a longer period may be necessary for complete remodeling of the injured area and restenosis. Low VV densities after balloon dilation can cause hypoxia, oxidative stress and microinflammation in the balloon-injured artery wall, which could lead to intimal growth. This is supported by micro-CT results of Gössl et al. These authors found that the areas of low VV densities within coronary arteries show decreased oxygenation and increased oxidative stress, which can cause microinflammation and subintimal proliferation and potentially initiate the early atherosclerosis development. It is possible that angiogenic stimulation to enhanced VV neovascularization during the early phase of acute vessel injury can reduce intimal hypoxygen by an increased oxygen supply. Kwon et al. showed a potential for advanced restenosis, as the artery lumen stenosis was about 45% following balloon injuries. Although VV neovascularization can serve as a compensatory mechanism for the delivery of more oxygen and nutrients to injured vessel walls for vascular repair during the advanced stage of restenosis, higher densities of VV may actually aggravate the

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**Table 2: Cross-sectional area of rabbit carotid arteries at the narrowest segment**

| Items       | Sham group  | Single injury | Angioplasty | F    | P    |
|-------------|-------------|---------------|-------------|------|------|
| Lumen       | 0.794 ± 0.126 | 0.660 ± 0.039 | 1.052 ± 0.015 | 1.735 | 0.201|
| Intima      | 0.070 ± 0.011 | 0.500 ± 0.105 | 1.006 ± 0.172** | 9.437 | 0.001|
| Media       | 0.388 ± 0.037 | 0.561 ± 0.085 | 0.739 ± 0.049*  | 5.140 | 0.015|
| I/M         | 0.179 ± 0.023 | 0.885 ± 0.194* | 1.400 ± 0.238*  | 7.609 | 0.003|

*significantly different vs. sham operated group; †significantly different vs. single injury group. I/M: Intima/media ratio.

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**Figure 2:** (a-c) Hematoxylin and Eosin staining of carotid arteries (×5). Mild intimal hyperplasia (surrounded by black dash line) was shown 8 weeks after absorbable suture was wrapped around the carotid artery (b). Pathologic intimal thickening with thrombically active plaque was observed 4 weeks following balloon angioplasty (c). Arteriolar vasa vasorum (VV) filled with polymer (black) in sections; (d-f) Increased invasion of macrophages (brown) (×5) in single and double injury groups, revealed by monoclonal antibodies against macrophage marker CD68; (g-i) Accumulation of smooth muscle cells (×40) in injury groups. Spindle-shaped smooth muscle cells were brown in color with long nuclei stained dark blue.
restenosis. This could function as a conduit for macrophages or inflammatory factor infiltration that could promote the progression of inflammation and restenosis formation. This is supported by Moulton et al.[17] who showed that inhibition of VV neovascularization reduced macrophage accumulation and progression of atherosclerotic plaques. Indeed, along with increased VV formation, the invasion of macrophages was detected in the primary atherosclerotic group and was further enhanced following balloon injury. Two aspects of VV neovascularization in the progression of restenosis may explain why some studies showed that angiogenic cytokines can accelerate reendothelialization, improve endothelial function, and significantly reduce intimal proliferation in models of balloon-induced arterial injury.[29,30] while other studies have shown that angiogenic cytokines can accelerate atherogenesis.[16,31] Further experimental studies are required in order to better understand of the right timepoint for both angiogenesis and anti-angiogenesis/anti-inflammation therapy in order to reduce arterial restenosis.

We found that the luminal surface of VV was higher in the double injured angioplasty group, compared to the sham and primary injury groups. This could indicate elevated exchange between cellular and noncellular components and the vascular wall. This could allow more oxygen and nutrition transported into the vessel wall, which could enhance arterial repair following acute injury. Alternatively, it could promote intimal hyperplasia and restenosis, as newly formed VV are considered to be more fragile, leaky and more prone to rupture.[35]

In conclusion, The double-injury rabbit model of restenosis coupled with perivascular manipulation and secondary balloon injury closely mimics human disease. Marked VV neovascularization occurred in rabbit carotid arteries following balloon injury. The number of VV was increased proportionally to intimal hyperplasia while VV density was decreased in the injured arteries. Additional studies are required to determine whether modifying VV growth could provide therapeutic effects on neointimal hyperplasia and restenosis following arterial injury.

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