Ultrastructural Characteristics of Experimental Arterial Medial Fibroplasia Induced by Vasa Vasorum Occlusion1,2

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Arterial fibrodysplasia is a pathologic entity of increasing clinical importance and undetermined etiology. Mural ischemia may be of primary importance as a cause of the most common dysplastic lesion, medial fibroplasia [8]. Recent electronmicroscopic studies of fibrodysplastic human renal arteries by the authors associated abnormal vasa vasorum to regions of medial dysplasia. These observations lend indirect support to the tenet that impairment of vessel wall blood supply may be a necessary accompaniment of this disease. The present investigation was undertaken to determine if experimentally induced vasa vasorum occlusion in canine vessels would produce histologic changes of medial fibroplasia. Ultrastructural studies were performed to better define the cellular and subcellular alterations attending this model of arterial fibrodysplasia.

MATERIALS AND METHODS

Sixteen adult dogs, 19 to 42 kg in weight, were utilized for study. Iliofemoral arterial segments approximately 5 cm in length were dissected from surrounding tissues in anesthetized animals (intravenous pentobarbital, 30 mg/kg). Care was taken to avoid unnecessary disturbance of adventitial structures. Vessels were isolated proximally and distally with polyethylene loops and small intervening branches occluded with ligatures. Blood within the isolated arterial segment was aspirated through a polyethylene catheter inserted into the lumen at the occluding loop. In experimental arteries, vessels were infused with a thrombin-gelatin mixture under a pressure of 100 mm Hg. This material was prepared by adding 0.5 ml of bovine thrombin (5000 NIH units) to 5 ml of freshly constituted 10% gelatin solution. After 5 min, the thrombin-gelatin was aspirated. Blood flow was restored following irrigation of the vessel with balanced salt solution. The contralateral iliofemoral artery was treated as a control. An identical protocol was used except for substitution of balanced salt solution for the thrombin-gelatin mixture.

Both control and experimental iliofemoral arterial segments were removed simultaneously for study 1 week to 1 year later. India ink was infused at physiologic pressures into both arterial segments for demonstration of vasa vasorum prior to their harvest. Excised vessels were immediately fixed in a 2.5% cacodylate-buffered glutaraldehyde solution. Transmural tissue specimens, approximately 1.0 mm², were fixed for 3 hr. Tissues for scanning electron microscopy (SEM) were dehydrated, critical point dried, and coated with gold vapor. Tissues for transmission electron microscopy (TEM) were washed in cacodylate buffer for 6 to 8 hr and postfixed in 2% OsO₄ for 2 hr. Specimens were stained en bloc with 0.5% uranyl acetate in malac acid at pH 5.2.

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Following ethanol dehydration, the tissue was embedded in Epon 812. Thin sections were stained with uranyl acetate–lead citrate and coated with carbon. Arterial segments prepared for light microscopy were processed in a standard manner and stained with hematoxylin–eosin, Verhoeff, or Masson stains.

Cellular and subcellular characteristics of the experimental and control arterial walls were compared. Specimens were examined within the first week (three subjects), from 2 to 4 months (seven subjects), from 4 to 7 months (four subjects), and at 12 months (two subjects), postinfusion. Particular care was directed toward examination of those areas most typical of the entire arterial segment.

**RESULTS**

Thrombin–gelatin infusions effectively obliterated nearly all vasa vasorum (Fig. 1A). Reappearance of these nutrient vessels was very unusual during the course of the study. Conversely, normal vasa vasorum, documented with India ink infusions, were present in all control specimens (Fig. 1B). Intraluminal thrombus and perivascular or adventitial inflammation were not encountered in either experimental or control vessels. Gross morphologic alterations, in the form of increased vessel wall thickness, were observed only in the experimental arterial segments. This change became more obvious in vessels examined late following thrombin–gelatin infusion.

Control arteries, irrespective of time studied, exhibited normal cellular and subcellular features. Endothelial cells, subendothelial structures of the intima, as well as adventitial tissues appeared normal. Fibroblasts within the medial tissues were few in number and also normal in appearance. In particular, abnormalities of medial smooth muscle were not observed. Smooth muscle cells of control segments, arranged in regular strata, exhibited certain common features including: close apposition of cytoplasmic processes to neighboring cells, an ovoid nucleus, having deep

![Fig. 1. (A) Experimental specimen of canine ileofemoral artery 200 days postinfusion of thrombin–gelatin. Vasa vasorum were not demonstrated following India ink infusion in any arterial segments treated in this manner. (B) Control specimen of contralateral ileofemoral artery following infusion of buffered salt solution. India ink-filled vasa vasorum (arrows) are identified at the media-adventitial junction.](image-url)
indentations and convolutions, surrounded by cellular organelles (mitochondria, rough endoplasmic reticulum, Golgi complexes, and free ribosomes), orderly appearing myofilaments, predominantly parallel to the longitudinal cell axis, radiodense bodies along the cell wall denoting myofilament attachment to the plasma lamina, as well as basal laminae and peripheral micro-pinocytotic vesicles along the plasma lamina.

Experimental arterial segments having occluded vasa vasorum exhibited distinct morphologic abnormalities. Derangements were less if any vasa vasorum persisted. Although serial sections at different time periods from individual segments were not obtained, the process leading to morphologic alterations appeared anatomically covert the first few months. It was not until 3 months postinfusion that marked changes in smooth muscle with increases in extracellular connective tissue and the appearance of myofibroblasts became obvious.

The intima in experimental vessels was usually unaltered. Occasionally, subendothelial derangements were substantial, yet endothelial cells were invariably maintained (Fig. 2). In vessels having intimal changes, excessive amounts of collagen, elastic tissue, and amorphous ground substances were common. Migratory smooth muscle cells were infrequent accompaniments of the former.

Medial abnormalities were the most obvious consequence of vasa vasorum occlusion. Smooth muscle and connective tissue derangements were clearly present 3 months postinfusion, being most marked in
tissues examined at 1 year. Cellular pleo-
morphism appeared more severe in the
periphery of the media than in tissues
nearer the lumen. A continuum, from ac-
tive smooth muscle cells to distinctly ab-
normal myofibroblasts (Fig. 3–7), was ob-
served in the media of the experimental
vessels.

Two smooth muscle cell abnormalities
were recognized. Rarefication proved to
represent an early type of smooth muscle
degeneration, characterized by areas within
the cytoplasm devoid of myofilaments,
swollen mitochondria, vacuoles, and basal
lamina indistinctness. The second type
abnormality, mummification, seemingly rep-
resented a more advanced form of cell
degeneration. It was characterized by
picnotic nuclei, perinuclear vacuolation,
loss of juxtanuclear organelles, and large
peripheral vacuoles with indistinct myo-
filaments. Whereas rarefication may repre-
sent a reversible process, mummification
is believed a reflection of irreparable cell
damage. Smooth muscle cell fragmentation
with frank necrosis was occasionally ob-
served in these specimens but was not a
dominant feature.

Myofibroblasts were a common finding in
vessels having occluded vasa vasorum.
Morphologic characteristics of these cells
ranged from those of modified fibroblasts
to that of atypical smooth muscle cells. In
contrast to fibroblasts, whose nuclei were

Fig. 3. Smooth muscle cell (S) within media of thrombin–gelatin-infused artery at 200 days. Areas devoid
of myofilament are scattered throughout the cytoplasm (arrows) and are characteristic of rarefication
(see text). Mitochondria (m) are swollen. Intracytoplasmic vacuoles (v) are present, but micropinocytotic
vesicles (mp) are reduced in number. Collagen (c). Original TEM ×15,000.
FIG. 4. Orderly stratum of medial smooth muscle cells (S) 200 days postthrombin–gelatin infusion. Myofilaments exhibit homogeneous indistinct features characteristic of mumification. Perinuclear vacuolation (arrow), loss of subcellular organelles, and peripheral vacuolation with confluence of micropinocytotic vesicles are additional features typical of this degenerative state. Moderate accumulation of ground substance and connective tissue elements surround these cells. Collagen (c). Homogeneous mucoid substance (mu). Original TEM x3000.

usually smooth and ovoid in shape, the myofibroblast nucleus was invariably convoluted with numerous indentations and evaginations. In this respect myofibroblasts resembled contractile smooth muscle cells (Figs. 6 and 7). Other features distinguishing myofibroblasts from fibroblasts were more juxtanuclear loci of cytoplasmic organelles and the existence of cytoplasmic filaments. Basal lamina of myofibroblasts were often indistinct.

The adventitia of vessels subjected to thrombin–gelatin infusions appeared relatively normal and, excepting an absence of vasa vasorum, could not be differentiated from control arteries. Although disrupted vasa vasorum were observed in 1-week studies, later examination failed to document restoration of normal nutrient blood vessels.

DISCUSSION

The pathogenesis of arterial fibrodysplasia has been an elusive subject since this disease entity was first recognized more than 40 years ago. Medial fibroplasia is the most common form of arterial dysplasia [8]. This type of dysplasia represents a continuum. Lesions vary from focal stenoses to multiple constricting lesions in series with intervening aneurysmal out-
pouchings. Two variants of medial fibroplasia exist. A peripheral form is manifest by dense fibroproliferative changes with excesses in collagen, increased ground substance, and loss of smooth muscle in the outer media. Relatively little involvement of the inner media occurs. The second form, diffuse medial fibroplasia, is characterized by more severe disruptions of the media architecture with loss of most recognizable smooth muscle. Medial thinning, alternating with accumulations of fibrous tissue, occurs most commonly with diffuse disease.

In contradistinction to other types of arterial dysplasia, mounting evidence supports the likelihood that several factors contribute to the evolution of medial fibroplasia. Particularly suspect are: (1) mural ischemia, being enhanced by the paucity of vasa vasorum found in vessels most commonly affected, (2) unusual mechanical stresses on arteries most susceptible to fibrodysplastic changes, and (3) hormonal influences, reflected by the fact that more than 90% of patients exhibiting medial fibrodysplasia are female.

Although the present investigation was directed at assessing mural ischemia as a cause of medial fibroplasia, other factors should not be ignored. In fact, two earlier studies have provided indirect support to the mechanical and hormonal hypotheses. Specifically, Leung, et al. demonstrated that repeated stretching of smooth muscle cells in culture potentiated synthesis of collagen and certain acid mucopolysacchariades [2]. Similarly, Ross and Klebanoff were among the first to demonstrate estro-
FIG. 6. Myofibroblast (MF) exhibiting features of both smooth muscle and fibroblast cells. Prominent nuclei are indented and convoluted. Myofilaments appear in the peripheral cytoplasm. Incomplete basal lamina and diffuse distribution of subcellular organelles characterize this cell. Collagen (c). Homogeneous mucoid substance (mu). Original TEM $\times 15,000$.

Impairment of vascular wall blood supply is a common cause of tissue hypoxia. Hypoxia can stimulate smooth muscle cells to produce extracellular connective tissue proteins in vivo [6]. Complex investigations, designed to measure the influence of hypoxia on smooth muscle cells previously stimulated to produce collagen, elastin, and ground substances have not been conducted.

Dominant pathology encountered in the present study was that of the myofibroblast cell. Although speculative, the source of these cells appeared to be vascular smooth muscle cells. Ross demonstrated the capability of smooth muscle cells to synthesize and secrete extracellular proteins in vitro [7]. The stimulus for smooth muscle cells to drastically alter their morphologic appearance and function is unknown. Mural ischemia with tissue hypoxia is assumed the inciting event in the present study, but altered tissue pH, accumulation of metabolites, or some other factors may be more important. No data were evident from the current investigation to suggest that myofibroblasts evolved from proliferation of previously dormant mesenchymal cells. This does not exclude such a possibility. Indeed, medial smooth muscle cells are considered by Wissler to represent multifunctional mesenchymal cells [9]. Gabbiani et al. on the other hand, view the myofibroblast as a contractile fibroblast rather than a modified smooth muscle cell [1].
as a cause of arterial medial fibrodysplasia has received indirect support from a number of earlier studies. Obliteration of vasa vasorum using a thrombin-containing mixture has been described by Nakata [5], Nakata and Shionoya [4]. Although their studies related to the aorta, their light microscopic descriptions of medial changes paralleled those of the current study. Clinical recognition of the peripheral form of medial fibroplasia [8], and the observation that altered smooth muscle cells and myofibroblasts were more common in the outer media of vessels subjected to thrombin-gelatin infusions, may be a reflection of the degree of vessel wall hypoxia. Obliteration of vasa vasorum should have a greater effect in reducing oxygenation and nourishment of cells in the outer media, in comparison to those near the vessel lumen. Wolinsky and Glagov documented the constancy of the avascular inner media in aortae of different mammals [10]. In this regard, limitations in transluminal passage of nutriments may be important in peripheral medial fibroplasia. This would account for inner medial smooth muscle preservation and development of dysplastic disease in deeper layers. The duration and exact degree of ischemia necessary to stimulate myofibroblasts are unknown. Madden, et al. have demonstrated these cells in tissues subjected to limited hypoxic events [3]. They documented active contractile processes in musculature occurring long after inciting stimuli had been removed. In many respects, in the current study the lack of early morphologic changes following vasa

**Fig. 7.** Myofibroblast (MF) in secretory state typical of fibrodysplastic process following vasa vasorum occlusion. Dilated cisternae and prominent Golgi complexes (gc) reflect active secretion. Fibrous bundles (f). Mitochondria (m). Original TEM ×15,000. Inset: Extracellular deposition of proteinaceous matter (arrow) by active exopinocytosis. Original TEM ×60,000.
vasorum occlusion followed by the late appearance of myofibroblasts parallels their experience.

Ultrastructural characteristics of experimental arterial medial fibrodysplasia induced by vasa vasorum occlusion in the current investigation bear a striking resemblance to human fibrodysplastic lesions. Modification of arterial smooth muscle cells to more primitive functioning myofibroblasts appears to be a major if not the primary pathologic feature of this fibroproliferative disease.

REFERENCES