Antioxidants Reverse Age-Related Collateral Growth Impairment

Steven J. Miller a, b  Brian J. Coppinger b  Xiaosun Zhou a  Joseph L. Unthank a, b

Departments of a Surgery and b Cellular and Integrative Physiology, Indiana University School of Medicine, Indianapolis, Ind., USA

Key Words
Apocynin  Arteriogenesis  Nitric oxide  Tempol

Abstract
Aging is a major risk factor for the development of cardiovascular diseases, including arterial occlusive disease. Oxidant stress increases with age, and may be a significant factor contributing to vascular dysfunction and disease. We have shown that aging and hypertension impair collateral growth, the natural compensatory response to arterial occlusive disease, and that antioxidants restore collateral growth in young hypertensive rats. The aim of this study was to test the hypothesis that oxidant stress mediates collateral growth impairment in nondiseased, aged rats. Ileal arteries were induced to become collaterals via ligation of adjacent arteries. Growth was assessed at 7 days by repeated in vivo measurements and comparison to same-animal control arteries. Collateral diameter enlargement did not occur in aged rats, but luminal expansion was stimulated by pretreatment with tempol. Co-administration of L-NAME with tempol prevented collateral development. Expression of p22phox mRNA was increased in aged versus young rat arteries, suggesting NAD(P)H oxidase as a source of reactive oxygen species. Treatment with apocynin increased collateral growth capacity, whether administered prior to, or 7 days following, arterial ligation. The results suggest that antioxidant treatment may be useful in promoting collateral growth to compensate for age-related arterial occlusive disease.

Introduction

The prevalence of cardiovascular disease, including arterial occlusive disease, increases with age [1, 2]. A normal compensatory response to arterial occlusion is the flow-mediated enlargement of preexisting vessels, or collaterals, that bypass the site of occlusion and supply blood to distal ischemic tissues [3]. This process, referred to as collateral growth or arteriogenesis, is inhibited in both humans and animals during aging [4–6]. Although functional and structural changes are known to occur in the vasculature with age, the mechanisms responsible for impaired collateral growth are unclear. Increased oxidant stress, due to elevated reactive oxygen species (ROS) production or to decreased function of natural antioxidant pathways, has been shown to increase with age [5, 7, 8] and may be a factor in inhibiting compensatory remodeling. Nitric oxide (NO) is required for both normal vascular function and collateral growth [9, 10], and its bioavailability may be negatively impacted by oxidative stress via scavenging by superoxide [11] or by other related mechanisms, such as endothelial NO synthase (eNOS) uncoupling [12]. Preclinical [8, 13] and human [14, 15] studies have demonstrated that antioxidants may be used to reverse age-associated endothelial dysfunction, consistent with a role for redox regulation of NO-dependent vascular function during aging.

Potential effects of age-related oxidant stress on collateral growth have not been extensively studied. Previous work in our laboratory [6] has demonstrated collat-
eral growth impairment in a rat model of aging. We have also demonstrated that the impairment in collateral growth in young spontaneously hypertensive rats (SHR) could be completely reversed by antioxidant treatment, including tempol and apocynin [16]. SHR have metabolic features similar to aging in that oxidative stress is increased [8, 17]. The objective of this study was to test the hypothesis that oxidant stress mediates the age-related collateral growth impairment and that anti-oxidant treatment administered subsequent to an arterial occlusion can reverse the impairment. This approach was pursued because vascular disease typically is in progress before treatment begins. Age-related and antioxidant effects on collateral growth were assessed with our established mesenteric model [18]. Tempol and apocynin were used as antioxidants that act through different mechanisms. Our results demonstrate that oxidative stress is present in aged resistance arteries and that pretreatment with either tempol or apocynin partially reverses the age-related impairment in collateral growth. These data establish that excessive levels of ROS are in part responsible for the inhibition of compensatory vascular remodeling. We also demonstrate that the administration of apocynin subsequent to arterial ligation reverses the collateral growth impairment in aged rats. Thus, a form of antioxidant therapy may be useful for treating age-related arterial insufficiency in the presence of ongoing occlusive disease.

**Methods**

**Animals and Groups**

All procedures performed in this study were approved by the Indiana University School of Medicine Institutional Animal Care and Use Committee. Young (−200 g, 2–3 months) and old (retired breeder) male Wistar (>600 g, 8–12 months) and Wistar-Kyoto (WKY, 355–390 g, 8–12 months) rats were obtained from Harlan (Indianapolis, Ind., USA) and acclimated for a minimum of 3 days prior to use. Since retired breeder Wistar and WKY rats are approaching 50% of their mean survival age, we consider them to be a suitable model of early aging. Apocynin (acetovanillone) was obtained from Fisher Scientific (Hampton, N.H., USA); 4-hydroxy-2,2,6,6-tetramethylpiperidine-N-oxide (tempol) and N^\text{G}-nitro-l-arginine methyl ester (l-NAME) were purchased from Sigma (St. Louis, Mo., USA). For pretreatment, all drugs were administered in drinking water 3 days prior to model creation and maintained until vessel diameters were obtained. For posttreatment, apocynin was administered in drinking water starting 7 days following model creation and maintained for an additional 7 days when vessel diameters were measured. Concentrations/doses were based upon studies that have demonstrated effectiveness in reducing oxidative stress [19–21], normalizing arterial ROS and NO concentrations [22], or affecting arterial remodeling [16]. Concentrations in drinking water were 5.0 mM for tempol (58.8 ± 3.22 mg/kg/day) and 3.0 mM for apocynin (32.5 ± 1.36 mg/kg/day). In an additional set of tempol-treated rats (n = 4), a low dose of the eNOS inhibitor l-NAME (0.1 mM; 1.6 ± 0.07 mg/kg/day) was added along with tempol in drinking water.

**Model of Collateral Growth and Its Assessment**

Experiments utilized a well-established model of mesenteric artery collateral growth as previously described [18]. This model uses sequential ligation of mesenteric arteries to create a low-dependent collateral pathway. It has the advantage of easy and rapid access, a well-defined two-dimensional pathway, and allows repeated assessments of arterial diameters over time. It also enables comparison of collateral arteries to in-animal control arteries that have experienced similar manipulations, but not increased flow. Briefly, a laparotomy was performed and the terminal ileum exteriorized into a heated tissue support chamber. The bowel and mesentery were immersed in phosphate-buffered saline (PBS) or covered with plastic wrap at all times. Several sequential ileal arteries were ligated such that a region of bowel containing ~45 microvascular perfusion units was dependent upon collateral arteries for perfusion. Digital images of collateral and same-animal control arteries were acquired under conditions of maximal dilation (1.0 mM adenosine and 0.1 mM sodium nitroprusside) with a dissecting microscope (Leica MZ 9.5) and camera (Spot Insight 4 Firewire). The bowel was returned to the abdominal cavity and the incision closed in two layers with 4–0 suture. After 7 (pretreatment) or 14 (posttreatment) days, the laparotomy and acquisition of digital images was repeated, inner arterial diameter was determined by measuring the red cell column with image analysis software (Image-J), and the percent change in diameter was calculated.

**Quantitative RT-PCR**

Relative differences in mesenteric artery mRNA expression were determined using real-time quantitative PCR. For vessel isolation to obtain RNA, the abdominal aorta of an anesthetized rat was cannuled above the iliac bifurcation. After ligating both renal arteries and the proximal aorta, the mesenteric circulation was perfused with 30 ml of cold PBS followed by 10 ml of RNA-later™ (Ambion, Austin, Tex., USA). Mesenteric arteries were isolated, excised, and preserved in RNA-later at ~20°C prior to RNA isolation. Tissues were weighed, disrupted using a bead homogenizer (FastPrep System®; QBIogene, Carlsbad, Calif., USA), and total RNA was purified using an RNaseasy Fibrous Tissue Mini Kit (Qiagen, Valencia, Calif., USA). RNA sample concentration was determined using A260/A280 (NanoDrop ND1000 Spectrophotometer; NanoDrop Technologies, Wilmington, Del., USA) and RNA integrity was assessed by analysis with an Agilent 2100 Bioanalyzer (RNA 6000 Nano Chip Kit; Agilent, Santa Clara, Calif., USA). Aliquots of purified total RNA (0.5 μg) were enzymatically treated to remove contaminating genomic DNA (DNAfree™; Ambion) and then reverse transcribed using Ready-To-Go You-Prime First-Strand Beads (GE Healthcare/Amersham Biosciences, Piscataway, N.J., USA) with random decamer priming. For PCR, aliquots of cDNA (5.0 μl, 1:50 dilution) were combined with primers and probes for either p22^Prx or β-actin endogenous control (TaqMan® Gene Expression Assays; Applied Biosystems, Foster City, Calif., USA) in the presence of a PCR mastermix (Quantitect™ Probe PCR Kit; Qiagen). Reactions were run in triplicate on an Applied Biosystems 7500 Real-Time
PCR System using relative quantification (ddCt) with dual-labeled (FAM/MGB) probes as the product detection method. Standard 7500 PCR cycling conditions were used. Differences in PCR product yields between groups were determined by comparing the fold differences between target mRNA after normalization to \(\beta\)-actin.

**Statistics**
Statistical analyses were done by one- or two-way repeated-measures ANOVA (SigmaStat 3.0). When significance between groups was detected, multiple pairwise comparisons were performed with the Holm-Sidak method. Data are presented as group averages with SE of the mean. The criterion for significance was \(p < 0.05\).

**Results**

**Tempol Reverses the Impairment in Collateral Growth**
We have previously shown [6] that collateral growth was impaired in retired breeder Wistar rats compared to young rats (\(\sim 2–3\) months). To test the possible involvement of oxidant stress in the suppression of collateral growth, Wistar retired breeder rats were pretreated with the superoxide dismutase mimic tempol 3 days prior to, and for 7 days following, model creation. Collateral growth was assessed 7 days after ligation as described in the Methods. Similar to previous results, collateral artery diameters significantly increased (33.0 \(\pm\) 4.3\%, \(p < 0.05\)) in young rats (data not shown), whereas no significant increase in collateral diameter was detected in untreated retired breeders (fig. 1). Collateral artery diameters in tempol-treated aged rats increased 17.6 \(\pm\) 5.8\% \((p = 0.03)\) compared to same-animal controls. Inclusion of a low (non-hypertensive) dose [23] of the NOS inhibitor \(\text{L-NAME} (0.025\text{mg/ml})\) completely inhibited tempol-assisted collateral growth, showing that the process is dependent on NO generation (fig. 1).

**NAD(P)H Oxidase as a Source of Oxidant Stress**
Vascular oxidant stress has been shown to increase with aging [5, 7, 8] and NAD(P)H oxidase is thought to be a major source of vascular superoxide [7, 8, 24, 25]. We examined mRNA expression of an NAD(P)H oxidase subunit in mesenteric arteries of young and retired breeder rats. Levels of \(p22^{phox}\) mRNA were found to be increased 1.5 \(\pm\) 0.2 times in control arteries of aged compared to young rats (fig. 2), suggesting an increase in NAD(P)H oxidase activation, and thus production of superoxide.

Since increases in superoxide due to NAD(P)H oxidase activation could limit the bioavailability of NO [11] and inhibit collateral growth in aged animals, we tested the ability of the NAD(P)H oxidase inhibitor and antioxidant apocynin [26] to promote collateral formation in aged Wistar rats. Apocynin was administered prior to model creation and continued until vessel diameters were obtained. Chronic apocynin treatment restored collateral growth (fig. 3; 18.3 \(\pm\) 2.2 vs. 3.0 \(\pm\) 0.4\%, control) equivalent to that previously found with tempol. Similar increases in collateral growth capacity promoted by both tempol and apocynin confirm NAD(P)H oxidase as a primary source of oxidant stress during vascular aging.

**Antioxidant Posttreatment**
To determine if antioxidant treatment could stimulate collateral growth if administered following, rather than prior to, an arterial ligation, the mesenteric collateral model was created in retired breeder WKY rats and the animals were allowed to recover for 7 days. Apocynin was then administered for an additional 7 days, and control and collateral artery diameters were determined. As shown in figure 4, apocynin posttreatment of the aged WKY significantly stimulated collateral growth (19.9 \(\pm\) 6.3 vs. \(-0.02\) \(\pm\) 2.3\%, same-animal control). Similar to Wistar rats (fig. 1), collaterals of aged WKY rats without treatment exhibited no significant enlarge-
Oxidant Stress Impairs Age-Related Collateral Growth

Discussion

This study is the first to demonstrate that antioxidant treatment administered both before and substantially after arterial occlusion can significantly reverse collateral growth impairment during aging in rats. The results from initial experiments using tempol to decrease oxidative stress and promote collateral growth in aged rats, along with the observation of increased p22phox expression, suggest that excessive vascular superoxide is an important factor in the suppression of collateral growth during aging. This conclusion is consistent with other studies indicating that excessive concentrations of ROS result in impaired collateral development [16, 27, 28]. While some studies have questioned the role of NO in collateral growth [28–30], other studies [9, 31–34] have established an intact NO system to be required for shear-mediated outward remodeling, including collateral growth. The inhibition of the effect of tempol on collateral growth by L-NAME suggests that redox regulation of NO production is involved in the aging-related impairment in collateral growth.

The ability of tempol, a superoxide dismutase mimetic, to promote collateral growth may be related not only to a reduction in superoxide but also to increased hydrogen peroxide. Previous studies have shown that peroxide may act as a dilatory factor during vascular disease [35], and we have recently shown in SHR that peroxide mediates the flow-mediated production of NO [22]. Thus, peroxide could be one form of ROS involved in age-related effects on compensatory remodeling, and merits further investigation.

While increased expression of p22phox mRNA in the retired breeder rat mesenteric arteries (fig. 2) does not itself indicate increased protein expression and NAD(P)H oxidase activation, it is consistent with many other studies that have shown NAD(P)H oxidase upregulation to be a primary source of vascular oxidant stress in cardiovascular disease [36–39] and during aging [7, 8, 25]. Briones et al. [7] demonstrated increased p22phox protein expression in mesenteric arteries of 22- to 24-month-old Sprague-Dawley rats, and our results establish that upregulation of this NAD(P)H oxidase subunit has occurred at an earlier age in Wistar rats. The ability of chronically administered apocynin to stimulate collateral growth is consistent with an involvement for NAD(P)H oxidase in the aging-related impairment. The fact that collateral growth was stimulated to the same extent with either tempol (fig. 1), to remove oxygen radicals, or apocynin (fig. 3), to suppress oxygen radical...
formation, predicts that superoxide overproduction is an important suppressor of growth.

Recent results from Heumüller et al. [40] suggest that apocynin does not inhibit NAD(P)H oxidase in endothelial cells because myeloperoxidase (MPO) is not present, and thus dimers essential for activity cannot form. These results, however, are based on cultured HEK293 cells that were transfected to overexpress NAD(P)H oxidase isoforms. Other studies have shown apocynin dimers to be present in endothelial cells [41] and MPO may be transferred into endothelial cells by a cytokera-
tin 1 pathway [42]. It is also likely that other peroxidases are present in endothelial cells in vivo that may substitute for MPO. Using in vitro vascular preparations, Schlüter et al. [43] have shown a requirement for exog-
enous peroxidase and hydrogen peroxide in order for apocynin to function as an NAD(P)H oxidase inhibitor. Recently, we have found that mesenteric arteries in the SHR [22] and retired breeder WKY [unpubl. observa-
tion] have in vivo hydrogen peroxide concentrations in excess of what was necessary for apocynin-mediated in-
hibition of superoxide production in the Schlüter study. In addition, we showed that apocynin reduced vascular peroxide concentrations to the same extent as the NAD(P)H oxidase peptide inhibitor gp91ds-tat [44]. Thus, although the possibility of nonspecific effects cannot be entirely excluded, it is likely that apocynin functioned as an NAD(P)H oxidase inhibitor in the aged mesenteric arteries.

While the results of this study do point to NAD(P)H oxidase as a major source of oxidant stress in aging ves-
sels, with the caveats mentioned above, possible involve-
ment of other ROS-generating pathways cannot be ex-
cluded. Tempol penetrates the mitochondrial membrane [45, 46], and other studies have implicated mitochondria [47] and xanthine oxidase [37, 48, 49] as vascular super-
oxide sources. However, it has been suggested that NAD(P)H oxidase influences xanthine oxidase as a mas-
ter regulator [50] and can stimulate mitochondrial pro-
duction of ROS [51]. Thus, such data are consistent with a role for NAD(P)H oxidase as the primary regulator of oxidant stress in the aged artery.

Previous results in our laboratory [16] showed that young WKY rats had the same mesenteric collateral growth capacity as young Wistar rats. However, it was not known if collateral growth in the WKY was impaired with aging. This must be established empirically, because mesenteric collateral growth is affected by aging to dif-
f erent extents depending on the rat genetic background [52]. We wanted to establish this impairment both to al-
low comparison of aged data with previous results using
the WKY, and also because the WKY is an inbred strain which is more genetically consistent, an important con-
sideration for future molecular studies. Results from the
retired breeder WKY experiments (fig. 4), previous stud-
ies [16], and preliminary data with apocynin pretreat-
ment [unpubl. results] establish that this strain is analo-
gous to the Wistar strain in terms of mesenteric collat-
eral growth capacity.

In the majority of preclinical studies, pharmacological
treatments are initiated at the time of or prior to model
creation, which does not reflect the clinical reality of pre-
xisting arterial insufficiency. The ability of apocynin to
promote collateral growth when administration was be-
gun 7 days following an arterial ligation (fig. 4) more
closely models clinical situations of chronic arterial oc-
cclusion. The positive results suggest that antioxidant
therapy could be clinically useful to stimulate collateral
growth and improve tissue perfusion in patients with pe-
ripheral vascular disease. These results are particularly
significant in light of the failure of growth factor therapy
for peripheral arterial disease to improve patient out-
come [53]. Although previous attempts to treat arterial
insufficiency in humans with antioxidants have had
highly variable outcomes, these results may be due to sev-
eral factors, including subject age and a lack of prescreen-
ing for oxidant stress [54]. Recent results suggest that
such prescreening can lead to significantly improved out-
comes [55].

A limitation of this study was that only retired breed-
er and not older rats were used. This age range was chosen
for study because the Wistar and WKY strains already
have a complete impairment in collateral growth, and
also because these strains are not readily available at old-
er ages. It is possible that antioxidant treatment may not
promote collateral growth in 18- to 24-month-old rats to
the same extent as in retired breeders due to non-enzy-
matic protein crosslinking in the arterial wall [56]. Simi-
lar vascular wall structural alterations may be occurring
at the age of the retired breeder rat, and if so, treatment
with agents to break protein crosslinks, either alone or in
combination with antioxidants, may be effective in re-
versing age-related impairments in collateral growth. A
second limitation is that mean arterial pressures (MAP)
were not measured, and it is possible that treatment-re-
lated alterations in MAP occurred which then influenced
the arterial structure. We consider this possibility un-
likely for three reasons: (1) the treatment duration was
relatively short; (2) no diameter changes were detected in
control arteries of treated rats, and (3) we have previous-
ly shown in SHR that similar pharmacological treatments restore mesenteric collateral growth independent of alterations in MAP [16].

In conclusion, we have demonstrated that treatment with antioxidants can significantly reverse the impairment in collateral growth that occurs with aging. The results are consistent with the hypothesis that overproduction of ROS, perhaps due to NAD(P)H oxidase activation, is a major factor in suppressing compensatory vascular remodeling during aging. Additional studies are warranted to determine the redox mechanisms that regulate collateral growth due to the potential for the treatment of the rapidly increasing incidence of vascular occlusive disease in the aging population.

References


Acknowledgments

The authors would like to thank P. Melanie Pride, Kathryn E. Madren (supported by an American Physiological Society Frontier in Physiology Professional Development Fellowship), and Randall G. Bills for their expert technical assistance with the animal surgeries, and H. Glenn Bohlen for his comments and suggestions. This work was supported in part by NIH grant HL 42898 and funds from the Department of Surgery (Indiana University).


