Edward F. Adolph Distinguished Lecture: The remarkable anti-aging effects of aerobic exercise on systemic arteries

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Seals DR. Edward F. Adolph Distinguished Lecture: The remarkable anti-aging effects of aerobic exercise on systemic arteries. J Appl Physiol 117: 425–439, 2014. First published May 22, 2014; doi:10.1152/japplphysiol.00362.2014.—Cardiovascular diseases (CVD) remain the leading cause of morbidity and mortality in modern societies, and advancing age is the major risk factor for CVD. Arterial dysfunction, characterized by large elastic artery stiffening and endothelial dysfunction, is the key event leading to age-associated CVD. Our work shows that regular aerobic exercise inhibits large elastic artery stiffening with aging (optimizes arterial compliance) and preserves endothelial function. Importantly, among previously sedentary late-middle-aged and older adults, aerobic exercise improves arterial stiffness and enhances endothelial function in most groups and, therefore, also can be considered a treatment for age-associated arterial dysfunction. The mechanisms by which regular aerobic exercise destiffens large elastic arteries are incompletely understood, but existing evidence suggests that reductions in oxidative stress associated with decreases in both adventitial collagen (fibrosis) and advanced glycation end-products (structural protein cross-linking molecules), play a key role. Aerobic exercise preserves endothelial function with aging by maintaining nitric oxide bioavailability via suppression of excessive superoxide-associated oxidative stress, and by inhibiting the development of chronic low-grade vascular inflammation. Recent work from our laboratory supports the novel hypothesis that aerobic exercise may exert these beneficial effects by directly inducing protection to aging arteries against multiple adverse factors to which they are chronically exposed. Regular aerobic exercise should be viewed as a “first line” strategy for prevention and treatment of arterial aging and a vital component of a contemporary public health approach for reducing the projected increase in population CVD burden.

artrial stiffness; endothelial dysfunction; oxidative stress; inflammation

Despite reductions in prevalence over the last 3+ decades, cardiovascular diseases (CVD) remain the leading cause of morbidity and mortality in the US and other modern societies.1 The primary risk factor for CVD is advancing age. Indeed, the prevalence of CVD of all causes increases progressively with advancing age, such that the great majority of individuals with CVD are middle-aged and older (MA/O) adults (43). This fact, combined with the projected increases in numbers of older adults worldwide, has led to concern of a new epidemic of CVD in the near future. A recent policy statement by the American Heart Association suggests that such concerns are well founded (31). The report projects that, by 2030, 40% of US adults will have at least one form of CVD, medical costs of CVD will triple, and that these changes will be driven largely by the aging of the population. It concludes that effective prevention strategies will be needed to limit the projected increase in the burden of CVD.

As we seek to identify effective prevention strategies, it is helpful to consider how aging increases CVD risk. Undoubtedly, many factors contribute. However, it is now well accepted that the key intermediary event that drives the majority of the increase in CVD risk with aging is the development of arterial dysfunction (37). Several changes to arteries with aging likely contribute to the increase in susceptibility for clinical CVD as we grow older, but two events, in particular, appear to be important (37). One is the stiffening of the large elastic arteries, i.e., the aorta and carotid arteries. As the nomenclature suggests, these arteries are designed to expand as they accept the left ventricular stroke volume with each contraction of the heart, and then recoil to create the necessary kinetic energy to drive the blood distally to our tissues and cells. The stiffening of these arteries with aging leads to numerous pathophysiologic effects that collectively increase our risk of CVD, including increases in arterial stiffness and pulse pressures, left ventricular hypertrophy (caused by repeatedly ejecting blood out into stiff arteries), and tissue damage as a result of increases in pulsatile flow, especially in high-flow vital organs such as the brain and kidneys (37, 38, 45).

A second clinically important change with aging is the development of endothelial dysfunction (37, 64). The vascular endothelium is a single cell layer at the interface between the flow of blood in the lumen of the artery and the walls of the artery. Once believed to be primarily a physical barrier charged with filtering solutes moving between the blood and arterial wall, the vascular endothelium now is understood to synthesize and release a wide array of biologically active molecules that act in autocrine and/or paracrine fashion to influence the function and health (resistance to disease) of arteries and their surrounding tissues. The most important of these endothelial-derived molecules is nitric oxide (NO), which exerts a provasodilatory and anti-coagulative, -proliferative, and -inflammatory protective effect on arteries (9, 64, 79). Endothelial dysfunction can be defined as any alteration from the healthy endothelial phenotype and typically is associated with reduced NO bioavailability (9, 64, 79).

With this background, we can establish a model in which aging leads to arterial dysfunction, as reflected in part by the development of large elastic artery stiffness and endothelial dysfunction, and this, in turn, increases our risk of clinical CVD. Our laboratory is interested in lifestyle and pharmacological strategies that act to delay or prevent the development of large elastic artery stiffening and endothelial dysfunction with aging, and/or to improve stiffness and endothelial function in MA/O adults with already established (baseline) vascular dysfunction (Fig. 1). In this recap of the 2013 Adolph Lecture, I will summarize the key findings of our work over the

1 This article is the topic of an Invited Editorial by Michael J. Joyner (33a).
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Fig. 1. Aging, vascular dysfunction, and cardiovascular disease (CVD) risk. Aging causes increased risk of developing CVDs in large part through the development of arterial dysfunction, characterized in part by vascular endothelial dysfunction and stiffening of large elastic arteries (aorta and carotid arteries). As such, evidence-based lifestyle, pharmaceutical, and nutraceutical strategies that either delay/minimize/prevent arterial aging, or, in late middle-aged/older adults who already demonstrate features of vascular aging, improve arterial function and reduce the risk of CVD, are of high biomedical research priority.

last 15+ years related to one of the most potent available strategies for preserving vascular function with advancing age: regular aerobic exercise. The ensuing discussion of the effects of aerobic exercise on vascular function with aging will focus on evidence in human subjects, as there are sufficient data to make conclusions in these populations. Discussion of the mechanisms mediating the anti-aging effects of exercise on vascular function will draw from research performed in both humans and rodents, as animal models are required in many cases to provide the most in depth, mechanistic insight. Much of the work discussed has been highlighted in previous reviews of the topic (14, 63-65, 67).

LARGE ELASTIC ARTERY STIFFNESS, AGING, AND EXERCISE

Functional Effects

One important question is whether aerobic exercise can reduce or prevent increases in large elastic artery stiffness with aging, and/or reduce stiffness in middle-aged/older adults who already demonstrate large elastic artery stiffening.

Aortic pulse-wave velocity. The gold standard clinical measurement of stiffness of the large elastic arteries is aortic pulse-wave velocity (PWV), typically measured as carotid-femoral PWV (3, 45, 46). As the term implies, aortic PWV is simply the velocity at which a pulse wave generated by left ventricular contraction travels down the aorta: the faster the velocity, the stiffer the artery. Aortic PWV increases progressively with adult aging, even in the absence of major CVD risk factors or clinical disease (3, 37, 75).

Using this technique, it was first shown that MA/O men who perform vigorous aerobic-endurance exercise (masters endurance athletes) demonstrated lower aortic PWV and systolic arterial blood pressure (SBP) compared with sedentary men of similar age, and closer to values observed in sedentary young controls (78). However, because this study did not report data on a young trained group, it was not possible to determine whether the lower aortic PWV in MA/O adults who perform aerobic exercise was the result of less stiffening with aging (i.e., a smaller age-associated difference) or the same change in stiffness from a lower (young trained group) baseline. Accordingly, we performed a cross-sectional comparison of premenopausal and postmenopausal healthy women who were either sedentary or engaged in endurance exercise training (75). Postmenopausal sedentary women had higher aortic PWV compared with premenopausal controls, but there were no significant differences between premenopausal and postmenopausal endurance exercise-trained women (Fig. 2), suggesting that regular aerobic exercise may prevent/reduce stiffening of the aorta with aging-menopause. The absence of a significant elevation in aortic PWV in the postmenopausal women who exercised was associated with a lack of increase in 24-h systolic arterial blood pressure and pulse pressure (66). Similar cross-sectional observations were later reported for healthy middle-aged men (39), and greater “light” dynamic physical activity was shown to be associated with lower aortic PWV in unfit older adults, suggesting that even a low-intensity aerobic exercise stimulus may exert destiffening effects (25). Importantly, moderate aerobic exercise interventions have been shown to reduce aortic PWV in healthy MA/O men and women (30, 87), indicating that aerobic exercise also can be viewed as a therapy for reducing aortic stiffness in this population.

Carotid artery compliance. The stiffness of the carotid arteries can be assessed using the combination of ultrasound imaging of changes in the internal diameter of one carotid artery throughout the cardiac cycle, with simultaneous mea-
measurement of changes in arterial pressure in the contralateral carotid artery using applanation tonometry (51, 76). Knowing the concurrent changes in diameter and pressure allows the calculation of carotid artery compliance (opposite in direction to stiffness) and the less arterial pressure-dependent measure of carotid \( \beta \)-stiffness (directly proportional to stiffness). Using this approach, we showed that carotid artery compliance decreased (\( \beta \)-stiffness increased) progressively with aging in healthy normotensive men and women, but that these age-associated changes were only \( \sim 50\% \) as great in subjects who engaged in regular aerobic exercise compared with their sedentary peers (51, 76) (Fig. 3A). These findings support the idea that regular aerobic exercise suppresses stiffening of the carotid arteries with advancing age in men and women free of clinical disease. Again, this effect of aerobic exercise appears to be specific for the large elastic arteries, as no differences are observed in the compliance of the peripheral large (femoral) artery between endurance exercise-trained and sedentary MA/O adults (51, 76).

We (51, 76) and others (1) also have shown that moderate aerobic exercise interventions improve carotid artery compliance \( \sim 30\% \) in healthy MA/O men and women (Fig. 3B). These findings suggest that, as with the aorta, aerobic exercise can be used therapeutically to destiffen the large elastic arteries in this group.

For sake of completeness, it is important to emphasize that better maintenance of carotid artery compliance with aging in aerobically exercising adults also has important implications for integrative autonomic-cardiovascular function. Consistent with this notion, preserved carotid artery compliance in older men who exercise regularly is associated with corresponding maintenance carotid baroreflex sensitivity (49).

**Types of aerobic exercise and resistance exercise.** Evidence suggests that several types of aerobic exercise may act to suppress the stiffening of the large elastic arteries with aging. MA/O adults engaged in running, cycling, swimming, or rowing all demonstrate enhanced carotid artery compliance compared with sedentary controls (1, 14, 51, 56, 76) (Fig. 4), and a swimming exercise intervention increases carotid compliance in this population (57). The fact that masters rowers, a group that engages in a combination of aerobic and resistance training activities, exhibit augmented carotid compliance (8), suggests that even an element of aerobic activity may be a sufficient stimulus to inhibit stiffening with aging. This is not an effect of exercise in general, because MA/O individuals who solely perform intensive resistance exercise have unchanged or even reduced carotid artery compliance compared with sedentary adults of the same age (47) (Fig. 4). Again, adding an aerobic component appears to offset the effects of vigorous resistance exercise on large elastic artery stiffness (8, 35). Finally, the influence of aerobic exercise appears to be specific for large arteries, as no differences were observed in the compliance of the peripheral large (femoral) artery between endurance exercise-trained and sedentary MA/O adults (51, 76).

The different types of aerobic exercise that will be discussed in this Review include running, cycling, swimming, and rowing. We have previously shown that these exercise modalities improve carotid artery compliance (51, 76), with a 50% increase in carotid compliance observed in the compliance of the peripheral large (femoral) artery between exercisers and sedentary controls.

**Superscript.** Carotid artery compliance is lower in healthy older men and women compared with young controls. However, the age-associated difference in carotid compliance in aerobically exercising (Ex) adults is only \( \sim 50\% \) as great as that observed in their nonexercising (non-Ex) peers. *\( P < 0.05 \) vs. young. **\( P < 0.05 \) vs. older non-Ex.
elastic arteries, as no effect is observed in the more muscular femoral arteries (8, 56).

**Summary.** In summary, several types of aerobic exercise act to inhibit the stiffening of the large elastic arteries with normal aging in adult humans. Importantly, regular aerobic exercise reduces large elastic artery stiffness in MA/O adults with baseline elevations in stiffness and thus can be viewed as a therapy for age-associated large elastic artery stiffening.

**Mechanisms of Action**

**Stiffness and aging.** The biological mechanisms underlying the inhibitory effect of aerobic exercise on large elastic artery stiffness are largely unknown. Increases in stiffness with aging are believed to be mediated in part by structural changes in the arterial wall characterized by remodeling of the extracellular matrix, including increases in collagen (fibrosis), fragmentation of elastin filaments, and formation of advanced glycation end-products (AGEs), molecules derived from glycolysis that cross-link structural proteins, conferring additional stiffness to the artery (36, 37). So-called “functional” influences involving increases in smooth muscle tone, perhaps linked to abnormal smooth muscle signaling and endothelial dysfunction, also are thought to contribute (44, 69). These processes may be triggered and/or sustained by the development of vascular oxidative stress and inflammation (36, 37, 80, 84).

**Age-associated stiffening and aerobic exercise in humans.** In healthy, estrogen-deficient postmenopausal women, acute systemic infusion of a supraphysiological concentration of ascorbic acid (vitamin C), which scavenges the reactive oxygen species, superoxide, selectively improves carotid artery compliance in sedentary subjects, increasing levels to those observed in endurance-trained women (52). This finding suggests the possibility that the greater carotid compliance in postmenopausal aerobically exercising women is associated with reduced superoxide bioavailability (reduced oxidative stress). However, no effect of vitamin C infusion or chronic oral supplementation on carotid compliance has been observed in sedentary MA/O healthy men (19). The influence of habitual aerobic exercise is not obviously attributable simply to favorable clinical characteristics and lower CVD risk factor burden in the trained state (51, 63, 76).

**Age-associated stiffening and aerobic exercise in rodents.** The mechanisms contributing to the destiffening effects of aerobic exercise on large elastic arteries with aging are difficult to study in humans because of the central location of these vessels in the systemic circulation. This prohibits direct access to/sampling from these arteries. Moreover, administration of agents that might activate or inhibit potential signaling pathways involved in modulating the stiffness of large elastic arteries must not alter vascular resistance, or the measurements could be confounded by baroreceptor counterregulatory responses. Peripheral arteries cannot be used as surrogate vessels, because they do not demonstrate age-associated stiffening. As a result, animal models must be employed to provide mechanistic insight. Unfortunately, few data are available at present.

In rats, collagen content was shown to be greater and elastin lower in aorta of older compared with young adult sedentary animals (55). The age-related difference in aortic stiffness, as determined by the elastic modulus calculated from stress-strain responses, was found to be smaller in animals that underwent 4–5 mo of swim training. However, this effect of swimming was not associated with differences in collagen or elastin protein expression, or calcium or polar amino acid content of the elastin layer.

More recently, we studied the effects of wheel running as a model of voluntary aerobic exercise on carotid artery stiffness in mice (22). Sedentary old animals demonstrated greater carotid stiffness than young controls. The age-related stiffening was associated with several changes in arterial wall composition, including increases in collagen I and III in the adventitial (outer) layer of the artery (Fig. 5), which were, in turn, associated with corresponding increases in the profibrotic cytokine, transforming growth factor-β1 (TGF-β1), and an in-
crease in smooth muscle α-actin, a marker of a myofibroblast or collagen-synthesizing phenotype. Elastin was reduced in the medial (middle) layer of the carotid artery, and this was accompanied by decreases in lysyl oxidase, a prosynthetic elastin enzyme, and increases in metalloproteinase 2, an elastin-degrading enzyme. There also was a small increase in arterial calcification in the arteries of the old mice, although no differences were observed in fibronectin, a glycoprotein involved in extracellular matrix remodeling. Voluntary wheel running for 10–14 wk initiated late in life reversed the carotid stiffening in old animals. This was associated with reductions in collagen I (Fig. 5) and III, TGF-β1, smooth muscle α-actin, and calcification, whereas elastin and its modulating enzymes were unaffected. In vitro experiments linked increases in TGF-β1 to oxidative stress-associated stimulation of collagen in adventitial fibroblasts. Recent preliminary studies from our laboratory using this mouse model suggest that wheel running also reduces stiffness in the aorta (aortic PWV), which is associated with reduced superoxide-related oxidative stress and accumulation of AGEs.

Summary. In summary, the limited mechanistic data available suggest that voluntary aerobic exercise may destiffen large elastic arteries with aging, at least in part, by reducing oxidative stress and inducing a remodeling of the extracellular matrix. The latter appears to involve reductions in adventitial collagen I, the primary load-bearing structural protein in the arterial wall, associated with reduced TGF-β1, a reversal of the adventitial myofibroblast phenotype, a decrease in calcification, and a reduction in stiffness.

**VASCULAR ENDOTHELIAL DYSFUNCTION, AGING, AND EXERCISE**

**Functional Effects**

As with large elastic artery stiffness, the key questions here are if regular aerobic exercise can slow, reduce, or even prevent the development of vascular endothelial dysfunction with aging and/or improve endothelial function in MA/O adults who already demonstrate baseline endothelial dysfunction.

**Assessment of endothelial function.** Vascular endothelial function is most commonly assessed in human subjects by measuring the dilation produced by a stimulus that evokes NO synthesis/release from the vascular endothelial cells (9, 64, 65, 79). This is termed an endothelium-dependent dilation (EDD): the greater the EDD, the greater the endothelial function (and vice versa). Two main types of stimuli are used to induce an EDD: a chemical stimulus (typically acetylcholine) or a mechanical stimulus (typically an increase in blood flow or shear rate). In either case, these stimuli activate the NO-producing enzyme in endothelial cells, endothelial NO synthase (eNOS), which synthesizes NO. The NO diffuses to the vascular smooth muscle cells in the medial layer of the artery, where it induces a guanylate cyclase-mediated increase in cyclic guanosine monophosphate, causing smooth muscle cell relaxation and dilation of the artery/increase in blood flow (EDD). The NO-dependent component of EDD can be assessed by chemically inhibiting NO production by eNOS using the competitive inhibitor Nω-monomethyl-l-arginine or other agents.

**Effects of aging and exercise on forearm blood flow responses to acetylcholine and NO bioavailability.** We (12, 63) and others (74) have used increases in forearm blood flow to brachial artery infusions of acetylcholine for assessing the influence of aerobic exercise on endothelial function in the microvasculature with aging. Among healthy men, sedentary older subjects demonstrate smaller increases in forearm blood flow in response to increasing concentrations of acetylcholine compared with young controls (12, 74) (Fig. 6A). In contrast, the forearm blood flow responses to acetylcholine do not differ in young and older men who regularly perform aerobic-endurance exercise (12, 74) (Fig. 6B), providing support for the concept that regular aerobic exercise may prevent age-associated microvascular endothelial dysfunction. Importantly, EDD in response to acetylcholine is improved by 12 wk of brisk walking in older previously sedentary men (12) (Fig. 6C). Indeed, such a walking program can restore EDD to levels observed in young sedentary and exercise-trained men and in older masters endurance athletes performing chronic, high-intensity endurance exercise training (63) (Fig. 6D). This aerobic exercise program also improves other expressions of endothelial function in older men, such as endothelial fibrinolytic capacity (68). Taken together, these findings provide evidence that aerobic exercise also can be viewed as a treatment for endothelial dysfunction of resistance arteries in MA/O men.

The greater EDD in young men and in older exercising men appears to be mediated by greater NO bioavailability because group differences in EDD are abolished with inhibition of eNOS (74) (Fig. 6D). Moreover, compared with young controls, basal NO production is reduced in sedentary, but not in endurance exercise-trained, healthy older men, and regular moderate-intensity aerobic exercise restores basal NO production to young adult levels in previously sedentary older men (63). It is not known if aerobic exercise has similar effects in postmenopausal women because, to our knowledge, there are no published data on this group.

**Effects of aging and exercise on brachial artery flow-mediated dilation in MA/O men and women.** The interaction between aging, aerobic exercise, and macrovascular endothelial function has been investigated using mechanical (flow-mediated) stimulation of EDD. We have shown that brachial artery flow-mediated dilation (FMD) is only ~50% of young control levels in healthy late MA/O sedentary men, but is well preserved in male masters endurance athletes of the same age (20, 21, 59) (Fig. 7A). In general, higher levels of brachial artery FMD in older aerobically exercising compared with sedentary men have been reported in the literature (23, 24, 60, 61). Recent work by our laboratory extended these cross-sectional observations by establishing that 8 wk of moderate-intensity regular aerobic exercise (i.e., brisk walking) improves brachial artery FMD by ~50% in previously sedentary healthy older men (59) (Fig. 7B). This exercise program restored brachial artery FMD to levels close to those observed in young men and in older endurance exercise-trained men (Fig. 7). Other laboratories have reported improvements in brachial artery FMD in response to aerobic exercise training in middle-aged healthy and/or mildly hypertensive men (62, 85). The results of studies performed in rodent models assessing EDD, generally in the carotid arteries and skeletal muscle arterioles, support these observations in human subjects (17, 70, 71).

The influence of regular aerobic exercise on macrovascular endothelial function with aging in women is far less clear. In the largest cross-sectional analysis of sedentary and endurance exercise-trained healthy MA/O men and women performed to
date \((n = 167)\), we found that brachial artery FMD was \(\sim 50\%\) greater in the exercising compared with sedentary men \((59)\). In contrast, there was no difference between the exercising and sedentary postmenopausal women, all of whom were estrogen deficient. No factor was identified during extensive post hoc analysis of the results that could explain this lack of difference in brachial artery FMD between the two groups. Comparisons of smaller samples of sedentary and aerobically exercising postmenopausal women have reported either similar observations to ours, i.e., no group differences \((7, 53)\), or higher values in exercising women \((5, 28)\). Using an exercise intervention design, we found that the same 8-wk program of brisk walking that produced a 50% increase in brachial artery FMD in MA/O men failed to increase FMD in estrogen-deficient postmenopausal women \((59)\). Other aerobic exercise trials in postmenopausal women have reported either no improvements \((7, 73)\) or significant increases \((2, 72, 86)\) in brachial artery FMD, and a single bout of aerobic exercise has been reported to increase brachial artery FMD in postmenopausal, estrogen-deficient women \((29)\).

In our initial trial \((59)\), absence of increases in FMD in response to aerobic exercise in our postmenopausal subjects may have been linked to their estrogen-deficient hormone status. To address this issue, we recently completed a follow-up intervention trial in which healthy, but sedentary estrogen-deficient postmenopausal women underwent 12-wk of aerobic exercise training with a preceding 12-wk period of either placebo treatment or estrogen replacement \((53)\). Brachial artery FMD was significantly improved in response to aerobic exercise training in estrogen-replaced, but not estrogen-deficient (placebo treatment) women \((53)\). These findings support the possibility that circulating estrogen concentration plays a permissive role for improvements in endothelial function with aerobic exercise in postmenopausal women. It has been suggested that sedentary estrogen-deficient postmenopausal women with lower baseline endothelial function may respond more consistently to aerobic exercise intervention than women with higher baseline levels \((73)\); however, the evidence supporting
in male and female patients with essential hypertension, high-intensity interval aerobic exercise training has been reported to improve brachial artery FMD, whereas continuous, moderate-intensity training was found to have no effect (48). Improvements in endothelial function have been observed in MA/O adults with durations of 30–60 min/day of aerobic exercise (2, 12, 57, 59, 62, 72, 86). Training periods of 8 wk up to 6 mo of aerobic training have been found to improve endothelial function in this group (2, 12, 48, 57, 59, 62, 72, 86), whereas a single bout of aerobic exercise also has been reported to increase brachial artery FMD in postmenopausal women (29). In general, much more insight is needed to establish the necessary aerobic exercise stimulus for preserving/improving endothelial function in older adults.

Summary. In summary, evidence obtained from both cross-sectional and intervention studies support the concept that regular aerobic exercise is associated with enhanced NO-mediated endothelial function in MA/O men. Moreover, experimental findings suggest that aerobic exercise also increases basal NO production and other endothelial functions in this group. Based on either an absence of data (forearm blood flow responses to acetylcholine) and/or inconsistent results (brachial artery FMD), the effect of aerobic exercise on endothelial function in estrogen-deficient postmenopausal women is uncertain at present. It is possible that circulating estrogen concentrations modulate endothelial responsiveness to aerobic exercise in postmenopausal women, and that low baseline function may influence improvements with training more in this group than in older men. Finally, multiple types of moderate-intensity aerobic exercise that can be performed by most MA/O adults appear to be sufficient for improving endothelial function. However, much more information is needed on the minimal and optimal overall aerobic exercise stimulus for enhancing endothelial function in this population.

Mechanisms of Action

In comparison to large elastic artery stiffness, much more is known about the mechanisms underlying the beneficial effects of aerobic exercise on vascular endothelial function with aging. This is because more studies have been performed using preclinical models, as well as the fact that the vascular endothelium is more accessible experimentally in peripheral vasculature of human subjects than is the case with the large elastic arteries. In general, oxidative stress and inflammation are the primary “macro-mechanistic” processes thought to be involved in mediating endothelial dysfunction with aging, and regular aerobic exercise is believed to preserve/restore endothelial function with aging, largely by reversing these mechanisms (14, 63–65, 67) (Fig. 9). Changes in expression and activation of eNOS as a possible contributor to altered NO signaling with aging and exercise training and the bioavailability of other vasodilatory molecules also have been investigated to a limited extent (63–65, 67).

Oxidative stress. Endothelial dysfunction with aging in preclinical models and adult humans is associated with arterial oxidative stress as a result of excessive vascular superoxide production and either unchanged or reduced antioxidant defenses (64, 65). Excessive vascular superoxide production with aging, in turn, is mediated primarily by a combination of increased expression and activity of the oxidant enzyme, nic-
Fig. 9. Mechanisms of aging and aerobic exercise on endothelial function. Aging induces endothelial dysfunction via vascular oxidative stress and inflammation involving excessive superoxide ($O_2^{-}$) production as modulated by nicotinamide adenine dinucleotide phosphate (NADPH) oxidase, superoxide dismutase (SOD), uncoupling of endothelial nitric oxide synthase (eNOS), and other mechanisms (e.g., dysfunctional mitochondria). Excessive superoxide causes peroxynitrite formation (ONOO$^-$) and protein oxidation (nitrotyrosine), as well as activation of nuclear factor-$\kappa$B (NF-$\kappa$B) and induction of inflammatory cytokines.

We have investigated the possibility that regular aerobic exercise prevents or reverses the development of oxidative stress-dependent endothelial dysfunction with aging by influencing one or more of the mechanisms described above. To do so, we have employed an integrative translational research approach combining observations made in preclinical mouse models and clinical studies in human subjects varying in age and aerobic exercise status.

**Nitrotyrosine Abundance.** In our investigations, we have assessed oxidative stress using the cellular marker, nitrotyrosine, which reflects oxidant modification of proteins by depicting the extent of nitration of tyrosine residues. In our initial preclinical study in mice, we found that 10–14 wk of daily wheel running, a model of voluntary aerobic exercise training in humans, started late in life (~26–28 mo), restored NO-mediated EDD in old mice to that of young controls (17). Aortic nitrotyrosine was 200% greater in old compared with young cage control (sedentary) animals, but wheel running reduced levels in old mice to those observed in the young controls (Fig. 10A). We then showed that, in endothelial cells biopsied from the brachial artery of human subjects, nitrotyrosine was markedly greater in cells from older compared with young sedentary men (58). However, endothelial cells obtained from older men who regularly performed aerobic exercise demonstrated levels of nitrotyrosine as low or lower than those observed in young controls (58) (Fig. 10A). Taken together, these results support the view that regular aerobic exercise prevents/reverses the development of oxidative stress with aging in whole arteries of mice, as well as in vascular endothelial cells of healthy adult humans.

**Superoxide.** To determine the potential role of superoxide in these age- and aerobic exercise-related differences in endothelial function, our laboratory and others have performed functional studies in which superoxide bioavailability was experimentally manipulated, and its effects on function measured. In our original study in mice (17), ex vivo administration of the superoxide scavenging compound, TEMPO, restored EDD in old cage control mice, but had no effect on young cage controls or on old mice given access to voluntary running wheels late in life. Similarly, infusion of ascorbic acid (vitamin C), a potent antioxidant when administered at doses/rates that create supra-physiological circulating concentrations, selectively restored EDD in older sedentary men, while having no effect on young controls or on older men who performed regular vigorous aerobic exercise (19, 74). Recently, we found that ascorbic acid infusion improved brachial artery FMD in both sedentary and aerobic exercise-trained estrogen-deficient postmenopausal women, but not in estrogen-replaced women who had undergone aerobic exercise training (53). Collectively, these findings indicate that excessive superoxide bioavailability contributes to endothelial dysfunction with aging, and that the enhanced endothelial function in aerobicically exercising older men and estrogen-sufficient postmenopausal women is the result of reduced superoxide suppression of endothelial function.

**NADPH Oxidase.** A reasonable next question may be the source(s) of the age- and aerobic exercise-associated differences in vascular superoxide production. In mice, we found that NADPH oxidase expression (p67$^{phox}$ subunit) and activity were increased in the aorta with sedentary aging, but that these...
changes were reversed in old animals that ran voluntarily on wheels for 10–14 wk late in life (17) (Fig. 10B). Moreover, we showed that ex vivo inhibition of this enzyme system with apocynin restored EDD in old cage control mice to levels observed in young animals, but had no effect on old mice given access to running wheels (17). Consistent with these preclinical findings, we found that the expression of NADPH oxidase (p47phox subunit) in endothelial cells obtained from the brachial arteries of older sedentary men was much greater than in young controls, but that expression in cells of older aerobically exercising men was as low or lower than in young men (58) (Fig. 10B). These observations support the idea that aerobic exercise enhances endothelial function with aging, at least in part, by inhibiting superoxide production/signaling by the oxidant enzyme, NADPH oxidase.

eNOS UNCOUPLING AND MITOCHONDRIA SUPEROXIDE PRODUCTION. Regular aerobic exercise may inhibit excessive superoxide-associated suppression of NO-mediated EDD through other mechanisms as well. For example, we (21) and others (33) have shown that administration of BH4 to reduce eNOS uncoupling restores EDD in MA/O sedentary adults to levels observed in young controls, and this effect appears to be mediated by restoration of NO bioavailability (33). Similar observations have been reported in skeletal muscle arterioles with aging in rodents (10). In contrast to sedentary aging, we find that BH4 administration has no effect on brachial artery FMD in MA/O men who regularly perform aerobic exercise (21), indicating that habitual exercise preserves BH4 bioactivity with aging, presumably by reducing vascular oxidative stress and consequent oxidation-inactivation of this molecule. Moreover, preliminary data from our laboratory suggest that aortic mitochondrial superoxide production increases with aging in cage control mice, but that this is reversed in old mice engaged in voluntary wheel running. Our preliminary work also suggests that ex vivo administration of mitochondrial-specific antioxidants improves EDD in old sedentary mice, but has no effect on young animals or on old wheel-running mice (27). Collectively, these observations support a role for eNOS “recoupling” and reduced mitochondrial superoxide production in the ameliorative effects of voluntary aerobic exercise on age-associated endothelial dysfunction.

SUPEROXIDE DISMUTASE. We have shown that old mice given access to running wheels late in life demonstrate increases in aortic total, manganese (mitochondrial), and copper-zinc (cytosolic) + extracellular superoxide dismutase activity compared with old cage controls (17) (Fig. 11A). Consistent with these preclinical results, we find that expression of manganese superoxide dismutase in brachial artery endothelial cells obtained from MA/O men who perform regular aerobic exercise is 50% greater than in cells from older healthy sedentary men and equivalent to levels in young controls (58) (Fig. 11B). In addition, we have shown that circulating endothelium-derived extracellular superoxide dismutase activity is reduced with aging in sedentary men, but preserved at young adult levels in MA/O men engaged in habitual aerobic exercise (58) (Fig. 11C). These observations indicate that preserved/enhanced superoxide dismutase antioxidant defenses may represent another mechanism by which regular aerobic exercise inhibits vascular oxidative stress and maintains healthy endothelial function with aging.

eNOS. Finally, upregulation of eNOS also may contribute to preserved NO bioavailability in the setting of regular aerobic exercise and aging. We (17) and others (70) have reported that expression of eNOS is greater in arteries from old mice who regularly exercise in running wheels or on motorized treadmills. In addition, we have shown that activation of eNOS, as indicated by increased phosphorylation at Ser1177, is increased in aorta of old mice given late-life access to running wheels compared with cage control old animals (17). Greater eNOS expression and activation could result in greater constitutive endothelial NO production and, therefore, contribute to enhanced vascular NO bioavailability in the aerobic exercise-trained state with aging.

Inflammation. Inflammation and oxidative stress are mutually reinforcing processes, so it is not surprising that the development of chronic low-grade inflammation is a hallmark of primary vascular aging, i.e., aging in the absence of other major risk factors or clinical disease. This is supported by multiple lines of evidence, including age-associated increases in proinflammatory proteins in the whole arteries of experimental animals and autopsied adult humans, as well as in vascular endothelial cells obtained from peripheral arteries and...
veins of living human subjects (40, 41, 64, 65, 83, 84). This chronic low-grade inflammatory state is associated with up-regulation of the major proinflammatory transcription factor nuclear factor-κB (NF-κB), and the transcription and expression of several of the inflammatory cytokines it regulates, as well as oxidative stress (40, 41, 64, 65). Consistent with these observations, we have shown that inhibiting NF-κB with sodium salicylate in old mice reduces the expression of proinflammatory cytokines in the aorta and restores EDD to levels not different from young mice by reducing vascular oxidative stress (40). These observations demonstrate that activation of NF-κB stimulates vascular inflammation with aging and induces chronic oxidative stress-related suppression of endothelial function.

INITIAL PRECLINICAL WORK WITH EXERCISE. Regular aerobic exercise has long been suggested to have anti-inflammatory effects based in part on selective observations of lower circulating markers of systemic inflammation, such as C-reactive protein, in aerobically exercising compared with sedentary MA/O adults (34, 54). Until recently, however, there was little or no direct evidence on the potential anti-inflammatory effects of aerobic exercise on vascular aging per se. In our initial (preclinical) work on this issue (41), we found that, compared with young (6 mo) animals, old (31 mo) cage control mice demonstrated increased aortic activation of NF-κB, as indicated by greater phosphorylated-to-total ratios of both the inhibitor of NF-κB kinase and the p65 subunit of NF-κB. These changes in NF-κB activation with aging were associated with increased aortic expression of the proinflammatory cytokines interleukin-1β (IL-1β), IL-6, interferon-γ, and tumor necrosis factor-α (TNF-α) (Fig. 12), as well as impaired NO-dependent EDD. Most importantly, 10–14 wk of voluntary wheel running, begun late in life (27–28 mo), normalized the age-related increases in aortic inhibitor of NF-κB kinase, NF-κB activation and proinflammatory cytokine expression (Fig. 12), while restoring endothelial function and NO bioavailability to levels of young animals. This was the first direct evidence to our knowledge of the anti-inflammatory effects of regular aerobic exercise on vascular inflammation with aging.

TRANSLATIONAL STUDIES WITH EXERCISE IN HUMANS. In our translational work on this topic, we first showed increased expression of the proinflammatory markers total and nuclear nuclear factor-κB (NF-κB), and the transcription and expression of several of the inflammatory cytokines it regulates, as well as oxidative stress (40, 41, 64, 65). Consistent with these observations, we have shown that inhibiting NF-κB with sodium salicylate in old mice reduces the expression of proinflammatory cytokines in the aorta and restores EDD to levels not different from young mice by reducing vascular oxidative stress (40). These observations demonstrate that activation of NF-κB stimulates vascular inflammation with aging and induces chronic oxidative stress-related suppression of endothelial function.

Fig. 12. Aging, aerobic exercise, and arterial inflammation. Aortic expression of the inflammatory cytokines interleukin-6 (IL-6), interferon-γ (IFN-γ), and tumor necrosis factor-α (TNF-α) increases with sedentary aging in mice, but this effect is reversed by 10–14 wk of voluntary (Vol) running in old animals. *P < 0.05 vs. the other groups. Data are from Lesniewski et al. (41).
NF-κB, IL-6, TNF-α and monocyte chemoattractant protein-1 (MCP-1) in vascular endothelial cells obtained from healthy MA/O men and women compared with young controls (15). The MA/O subjects also had greater plasma C-reactive protein and IL-6 concentrations and exhibited impaired brachial artery FMD compared with the young subjects. These findings suggested that, consistent with our findings in mice, primary vascular aging in humans is associated with development of a proinflammatory phenotype in the vascular endothelium featuring upregulation of NF-κB. In a follow-up investigation, we were able to show that the age-related increase in NF-κB expression in vascular endothelial cells is not observed in MA/O men who regularly perform aerobic exercise, and that this was associated with preserved brachial artery FMD (58). Preliminary results from our laboratory also suggest that the absence of elevations in NF-κB in endothelial cells of MA/O exercising adults is associated with a lack of increase in endothelial cell expression of IL-6.

Recently, we have attempted to gain insight into the possible role of reduced endothelial NF-κB activation in the enhanced vascular endothelial function associated with regular aerobic exercise in MA/O humans. To do so, we treated groups of young controls and MA/O sedentary and aerobically exercising subjects (~70% male) for 4 days with salsalate, a potent NF-κB-inhibiting agent, or placebo (82). Salsalate reduced NF-κB expression in endothelial cells of MA/O sedentary subjects in the absence of changes in the cells of the young controls or MA/O exercising group. Brachial artery FMD was impaired in the MA/O sedentary subjects compared with the other two groups under placebo conditions, as expected. Salsalate restored FMD in the MA/O sedentary group to levels not different from those of the young controls or MA/O exercising subjects, while having no effects in the latter two groups (Fig. 13). Experiments in which vitamin C was infused at supraphysiological concentrations to acutely reduce superoxide bioavailability revealed that the effects of regular aerobic exercise in inhibiting inflammation-associated suppression of brachial artery FMD with aging are mediated by reduced oxidative stress. Together, these observations suggest that upregulation of NF-κB is associated with oxidative stress-related impairment of EDD with aging in sedentary adults, but that regular aerobic exercise prevents these events to preserve endothelial function.

UPSTREAM OR REINFORCING MECHANISMS OF EXERCISE. We also have sought to obtain initial insight into the upstream or, perhaps, reinforcing events associated with these anti-inflammatory effects of aerobic exercise on aging arteries. It is well established that circulating immune cells, including monocytes, adhere to the vascular endothelium, attracted by chemoattractant proteins such as MCP-1 (11). Some percentage of the immune cells infiltrate the endothelial layer, with the infiltrating monocytes differentiating into macrophages. The macrophages and other immune cells lodging in the arterial wall can release superoxide and proinflammatory cytokines, stimulating inflammation and possibly contributing to a chronic low-grade inflammatory state.

In our initial investigation in mice, we found that the abundance of macrophages and T lymphocytes was increased in the adventitial and perivascular regions of the aorta of old compared with young cage control mice (41). There was no increase in the number of immune cells in the intimal and medial layers of the aorta in the old animals, suggesting either an “outside-in” infiltration pattern of the arterial wall or infiltration from the lumen to the outer regions of the wall with sedentary aging (6). Late-life voluntary wheel running for 10–14 wk reversed the age-related increases in aortic adventitial and perivascular macrophage infiltration and tended to have the same influence on T lymphocyte abundance. These observations suggest that regular aerobic exercise may exert its anti-inflammatory actions on arteries with aging, at least in part, by suppressing immune cell infiltration into the outer layer and surrounding (perivascular) region of the vascular wall. Fewer immune cells would result in less proinflammatory signaling and cytokine release into the arterial wall, reduced vascular inflammation, and enhanced function.

To provide translational insight into the findings in mice, we isolated peripheral blood mononuclear cells (PBMC) from groups of young and MA/O healthy sedentary men and women, as well as in a subset of the MA/O subjects before and after 8 wk of moderate-intensity aerobic exercise (26). Compared with the young controls, PBMC from the MA/O sedentary subjects had higher mRNA expression (PCR) of several proinflammatory/prooxidant factors, including NF-κB, the receptor for AGEs, TNF-α, MCP-1, NADPH oxidase, and the inducible form of NOS (Fig. 14). Regular aerobic exercise improved fitness in the MA/O subjects, and this was accompanied by reductions or a trend for reductions in all of these PBMC proinflammatory markers (Fig. 14). Preliminary findings from our laboratory also suggest that MCP-1 expression is higher in endothelial cells obtained from MA/O healthy sedentary adults compared with young controls, but that MA/O subjects who regularly perform aerobic exercise express MCP-1 at levels at or below those of young adults.

Collectively, these translational observations indicate that circulating immune cells develop a proinflammatory phenotype with sedentary human aging. Increased endothelial expression of chemoattractant proteins could increase infiltration of these cells into the arterial wall, eventually locating in the adventitial and perivascular regions of the arterial wall. Alternatively, greater macrophage accumulation in the outer regions of the wall may originate from the “outside-in” (adventitial-
perivascular regions). In either case, increased abundance of these cells in the arterial wall may contribute to a proinflammatory process featuring increased superoxide and proinflammatory cytokine production, ultimately developing into a state of chronic-low grade inflammation. Regular aerobic exercise appears to inhibit the development of a prooxidant/inflammatory phenotype in the circulating immune cells with aging and prevents the consequent downstream events leading to vascular inflammation.

Summary. In summary, regular aerobic exercise may act to preserve vascular endothelial function with aging via several mechanisms. Translational observations in mice and humans support the concept that aerobic exercise inhibits the development of excessive superoxide-dependent vascular oxidative stress with aging via some combination of suppressing the upregulation of the NADPH oxidant enzyme system, preventing the uncoupling of eNOS, constraining mitochondrial superoxide production, and preserving endogenous antioxidant enzyme defenses. This restraint of superoxide-associated oxidative stress, along with induction of eNOS expression and activation, contribute to greater NO bioavailability and enhanced EDD. Habitual aerobic exercise also exerts a potent anti-inflammatory effect on artery walls by potentially inhibiting several steps in the process of developing chronic low-grade inflammation and its suppression of endothelial function.

VASCULAR PROTECTION FROM “ADVERSE” FACTORS

Historically, much of the influence of regular aerobic exercise for reducing the risk of CVD with aging has been attributed to the secondary effects of exercise on traditional risk factors for CVD. That is, it has been assumed that habitual aerobic exercise favorably modifies risk factors such as blood pressure, plasma lipids, blood glucose control, and body weight/fatness, and that this lower risk factor burden is responsible for the maintenance of vascular health and associated reductions in CVD observed in exercising adults. However, epidemiological data indicate that, at most, only about one-half of the CVD risk-lowering effects of regular aerobic exercise can be attributed to such an influence (50). So clearly some other mechanism(s) are involved. Several years ago, we advanced the complementary hypothesis that regular aerobic exercise may be exerting at least part of its CVD-protective effects not by lowering traditional risk factors, but rather by inducing an increased “resistance” against the potentially harmful effects of existing levels of risk factors or, even more broadly, against a wide array of harmful factors to which arteries are commonly and chronically exposed (67, 81).

In our initial study in humans seeking support for this hypothesis (81), we used plasma low-density lipoprotein cholesterol (LDL-C) as the potential “harmful factor” and brachial artery FMD as the measure of vascular function. We found that having even a borderline elevation in plasma LDL-C was associated with greater impairment of brachial artery FMD in older sedentary adults compared with their peers who had normal/optimal plasma LDL-C. In contrast, among sedentary young adults, borderline elevations in plasma LDL-C had no influence on brachial artery FMD, as if young age exerted a protective effect against the potentially adverse influence of elevated plasma LDL-C on endothelial function. Most importantly, older adults who performed regular aerobic exercise demonstrated the same “resistance” to elevated plasma LDL-C as observed in young adults. A follow-up study in which impaired fasting blood glucose was used as a potential adverse factor (13) confirmed the results of the initial investigation, i.e., that regular aerobic exercise protects aging arteries from the possible harmful effects of circulating stressors (Fig. 15).
More recently, we have used a preclinical mouse model to gain insight into the mechanisms by which regular aerobic exercise confers this apparent increased resistance to stress in aging arteries. In our initial investigation, we found that ingestion of a “Western” diet (40% fat, 19% sucrose) for 10–14 wk started late in life exacerbated age-associated endothelial dysfunction (reduction in ex vivo carotid EDD) in old cage control mice (42). However, old mice given access to running wheels were protected against the adverse effects of the Western diet on endothelial function. The primary mechanism underlying the harmful influence of Western diet on vascular function in the old cage control mice was excessive superoxide-associated oxidative stress and resulting reduction in NO bioavailability. Voluntary wheel running in old mice preserved endothelial function during consumption of the Western diet by preventing superoxide-related reductions in NO bioavailability. More recent preclinical work by our laboratory suggests that the increased resistance to Western diet and other forms of arterial stress conferred by voluntary aerobic exercise is linked to reductions in mitochondrial-derived oxidative stress and improved mitochondrial homeostasis (27).

Summary

Taken together, these translational findings support the novel concept that regular aerobic exercise protects aging arteries, not only by lowering the risk factor burden to which they are chronically exposed, but also by improving the intrinsic resistance of the arteries to existing levels of a variety of potentially harmful factors. Our work also suggests that aerobic exercise confers this protection, at least in part, by preventing exacerbation of vascular oxidative stress in response to these adverse factors, which allows the preservation of NO bioavailability and arterial function.

OVERALL SUMMARY AND CONCLUSIONS

“Aging” of arteries is the major risk factor for CVD, and this will become a progressively greater challenge in the future with projected worldwide increases in the number of older adults. Regular aerobic exercise should be viewed as a “first-line” strategy for the prevention and treatment of arterial aging and, therefore, reducing population CVD risk. Aerobic exercise enhances the compliance of the large elastic arteries (inhibits arterial stiffening) and preserves endothelial function during adult aging. These benefits of aerobic exercise are attributable largely to the suppression of vascular oxidative stress and inflammation. Habitual aerobic exercise preserves arterial health by not only exerting favorable effects on selective CVD risk factors, but also by inducing a direct protective effect to aging arteries against multiple potentially “adverse” factors to which they are chronically exposed. Given the public health implications for prevention of CVD and related age-associated disorders, more information is needed regarding the minimal and optimal types, durations, intensities, and frequencies of aerobic exercise for preserving arterial function with aging.

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DISCLOSURES

No conflicts of interest, financial or otherwise, are declared by the author(s).

AUTHOR CONTRIBUTIONS

Author contributions: D.R.S. conception and design of research; D.R.S. interpreted results of experiments; D.R.S. drafted manuscript; D.R.S. edited and revised manuscript; D.R.S. approved final version of manuscript.

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