# Circulation Research



# **HYPERTENSION COMPENDIUM**

# Obesity, Adipose Tissue and Vascular Dysfunction

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ABSTRACT: Cardiovascular diseases are the leading cause of death worldwide. Overweight and obesity are strongly associated with comorbidities such as hypertension and insulin resistance, which collectively contribute to the development of cardiovascular diseases and resultant morbidity and mortality. Forty-two percent of adults in the United States are obese, and a total of 1.9 billion adults worldwide are overweight or obese. These alarming numbers, which continue to climb, represent a major health and economic burden. Adipose tissue is a highly dynamic organ that can be classified based on the cellular composition of different depots and their distinct anatomical localization. Massive expansion and remodeling of adipose tissue during obesity differentially affects specific adipose tissue depots and significantly contributes to vascular dysfunction and cardiovascular diseases. Visceral adipose tissue accumulation results in increased immune cell infiltration and secretion of vasoconstrictor mediators, whereas expansion of subcutaneous adipose tissue is less harmful. Therefore, fat distribution more than overall body weight is a key determinant of the risk for cardiovascular diseases. Thermogenic brown and beige adipose tissue, in contrast to white adipose tissue, is associated with beneficial effects on the vasculature. The relationship between the type of adipose tissue and its influence on vascular function becomes particularly evident in the context of the heterogenous phenotype of perivascular adipose tissue that is strongly location dependent. In this review, we address the abnormal remodeling of specific adipose tissue depots during obesity and how this critically contributes to the development of hypertension, endothelial dysfunction, and vascular stiffness. We also discuss the local and systemic roles of adipose tissue derived secreted factors and increased systemic inflammation during obesity and highlight their detrimental impact on cardiovascular health.

**Key Words:** adipose tissue ■ cardiovascular diseases ■ hypertension ■ obesity

he worldwide prevalence of obesity has tripled since 1975, with a parallel trend in type 2 diabetes. 1,2 Globally, over 1.9 billion adults were overweight or obese in 2016 and >60% of people with obesity live in developing countries.3 Today, about 2 out of 3 adults (69%) are overweight or obese in the United States, and current projections suggest that nearly 50% of adults in the United States will be obese by 2030.4 Predictions made in 2008, estimated up to 3.3 billion individuals to become overweight and obese by 2030, if adjusted for secular trends.<sup>5</sup> Nonadjusted predictions for 2030 generated by the same study predicted only 1.35 billion overweight and 573 million obese individuals for 2030,5 a number that was outdated already by 2016.3 While it is well documented that genetic and epigenetic factors contribute to obesity, environmental factors such as diet, physical activity, and environmental toxins also play a major role in the increased prevalence of this disorder (Figure 1). For example, the increase in obesity in the United States and

other industrialized nations is closely related to increased consumption of high fructose corn syrup and saturated fat and to reduced physical activity<sup>3,4</sup> (Figure 1). Further, there is emerging evidence that consumption of high fructose corn syrup diets by pregnant women programs the offspring for the subsequent development of obesity and associated cardiometabolic and cardiovascular diseases (CVD) in later life (Figure 2).6 These maternal influences seem to be mediated through adverse effects of metabolic factors such as impaired insulin signaling, dyslipidemia, and altered blood supply on placental function and resultant fetal nutrition as well as epigenetic influences that originate from maternal obesity.6

There is considerable evidence that overweight and obesity and their comorbidities, hypertension and insulin resistance, increase CVD and overall morbidity and mortality rates. 7-12 Indeed, a positive association even exists between a progressive increase in body mass index (BMI) within the normal and overweight range and the

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For Sources of Funding and Disclosures, see page 962.

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#### **Nonstandard Abbreviations and Acronyms**

ΑΜΡΚα1 5'AMP-activated protein kinase catalytic

subunit alpha 1

Ang 1-7 angiotensin 1-7 Ang II angiotensin II **ApoE** apolipoprotein E AT adipose tissue **BAT** brown adipose tissue

ВМІ body mass index

 $EBP\alpha$ enhancer binding protein alpha

CVD cardiovascular disease EC. endothelial cells

**EnNaC** endothelial Na2+ channel **eNOS** endothelial nitric oxide synthase

FABP4 fatty acid-binding protein

HIF1α hypoxia-inducible factor 1 alpha

interleukin

KDM5C lysine-specific demethylase 5C

NO nitric oxide

**PDGFR**α platelet-derived growth factor receptor

PRDM16 PR domain containing 16 ΡVΔΤ perivascular adipose tissue

RAAS renin-angiotensin-aldosterone-system SGK-1 serum/glucocorticoid regulated kinase 1

TNFα tumor necrosis factor alpha UCP1 uncoupling protein 1

**VSMC** vascular smooth muscle cell

WAT white adipose tissue

risk of CVD.67 In this regard, an analysis of the Framingham Heart Study showed a positive association between overweight (BMI, 25-29.9 kg/m²) and the relative risks of hypertension and CVD.8 In addition, the presence of childhood obesity has been shown to increase the risk for development of type 2 diabetes, hypertension, dyslipidemia, and atherosclerosis and related CVD in adulthood.9-11 This review discusses the various factors that promote vascular dysfunction and CVD in obesity, with a focus on the role of dysfunctional adipose tissue.

#### TYPES OF ADIPOSE TISSUE

# **Functionally Distinct Adipose Tissue Depots in Mice and Humans**

Adipose tissue (AT) is a dynamic organ distributed throughout the body with an almost unlimited capacity to expand during obesity. Several distinct depots can be defined by their location, size, cellular composition, and function. While many functions of AT are conserved

between mouse models and humans, their location and abundance can vary broadly. Mammals possess 2 major types of AT: white and brown (Figure 3). White adipose tissue (WAT) represents the largest proportion of wholebody AT and can be found around major organs and blood vessels in the abdominal cavity and subcutaneously (Figure 4). WAT stores excess energy in the form of triglycerides, and increased accumulation of WAT, particularly in visceral depots, is a key determinant of the relative risk for cardiometabolic disorders, hypertension, and CVD.12-17 To this point, fat distribution dictates CVD risk such that individuals with higher visceral AT and ectopic fat deposition have an increased prevalence of cardiometabolic disorders including hypertension, 18,19 dyslipidemia, and insulin resistance 15-17 compared with equally obese individuals with less visceral AT and relatively more subcutaneous fat. Thus, measurements limited to the determination of BMI do not reflect the actual risk for CVD conferred by obesity.

In contrast to WAT, brown AT (BAT) represents only ≈4.3% of all AT in adult humans and can be found in cervical, supraclavicular, axillary, paraspinal, mediastinal, and abdominal depots<sup>20-22</sup> (Figure 4). In addition, newborns possess interscapular BAT that decreases in size over time and is no longer detectable in adults.<sup>23</sup> BAT protects animals from hypothermia by dissipating energy as heat, via a process called nonshivering thermogenesis, and has more recently been found to also have anti-obesity and anti-diabetes properties and to confer broad cardiometabolic health benefits in humans.<sup>24</sup>

The main functional cell type of AT is the adipocyte or fat cell. White adipocytes contain a single large lipid droplet (unilocular) and only possess a small number of mitochondria. Brown adipocytes, on the other hand, have multilocular lipid droplets and contain a large number of cristae-dense mitochondria, which uniquely express UCP1 (uncoupling protein 1) in the inner mitochondrial membrane (Figure 3). UCP1 uncouples oxidative phosphorylation from adenosine triphosphate production, ultimately resulting in the generation of heat.25 More recently, several UCP1-independent thermogenic mechanisms have also been described.<sup>26</sup>

In addition to developmentally preformed brown adipocytes, mice and humans also have inducible brown adipocytes, referred to as beige or brite adipocytes. These multilocular fat cells come from a distinct developmental lineage and tend to be interspersed within WAT but also express UCP127 (Figure 3). At baseline or during thermoneutrality, beige adipocytes display a more white-like phenotype with large lipid droplets and low expression of thermogenic genes, 28 but activation by cold exposure, beta-adrenergic stimulation, or exercise results in the robust upregulation of a thermogenic program in a process commonly called "browning." While these cold-inducible brown-like adipocytes were first described almost 40 years ago,<sup>28-32</sup> their developmental

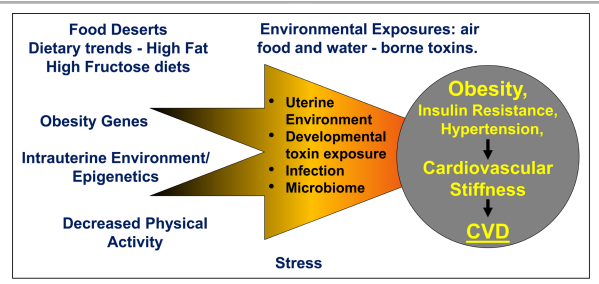


Figure 1. Obesity, vascular stiffness, and cardiovascular disease (CVD): genetic/epigenetic and environment interactions. A food desert refers to an area with limited access to nutritious, affordable food.

origin, molecular properties, and physiological roles have only more recently been investigated. In mice, beige adipocytes are enriched within subcutaneous fat depots and are rarely detected in visceral depots. Intriguingly, due to their temperature-dependent epigenomic plasticity, beige adipocytes also have the capacity to whiten in a warm environment.<sup>33</sup>

In light of their morphological and functional differences, it is not surprising that white and thermogenic brown/beige adipocytes are derived from distinct precursors.<sup>31,34–37</sup> White adipocytes arise from mural precursors that are CD24, CD34,<sup>38</sup> and PDGFRα (platelet-derived

growth factor receptor alpha) positive, <sup>12,39</sup> and subcutaneous and visceral white adipocytes seem to originate from distinct progenitor populations. <sup>40</sup> Developmentally preformed or classical brown fat is derived from a myogenic precursor expressing *Pax7*, *Engrailed-1*, and *Myf5* around embryonic days 9.5 to 11.5 in mice, even before white adipocytes develop. <sup>35,37,41</sup> Beige adipocytes, in contrast, originate from a vascular smooth muscle lineage. <sup>42</sup> Despite their distinct origins, the development of both brown and beige adipocytes is dependent on the transcriptional coregulatory protein PRDM16 (PR domain containing 16). Adult humans also have inducible

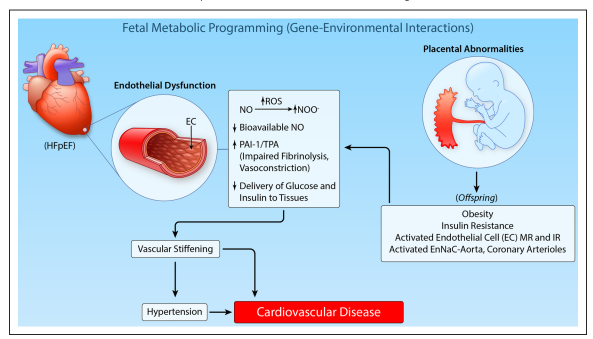


Figure 2. Prenatal programming and epigenetics in the genesis of obesity and cardiovascular disease (CVD) in offspring (Illustration credit: Ben Smith).

EC indicates endothelial cell; EnNaC, endothelial Na2+ channel; HFpEF, heart failure with preserved systolic function; IR, insulin receptor; MR, mineralocorticoid receptor; ROS, reactive oxygen species; and TPA, tissue plasminogen activator.

	White Adipocyte	Beige Adipocyte	Brown Adipocyte
Depots	Visceral and subcutaneous and most PVAT depots	Cervical*, supraclavicular*, axillary*, paraspinal*, renal*, thoracic PVAT*, subcutaneous (only rodents)	Interscapular (human only infants), thoracic PVAT*
Function	Storage of triglycerides, endocrine (secretion of adipokines and vasoactive factors)	Thermogenesis, anti-inflammatory properties, cardioprotective	Thermogenesis, anti-inflammatory properties, cardioprotective
Progenitor	CD24+, CD34+, PDGFRα+	Vascular smooth muscle origin	Myogenic origin Pax7+, En-1+, Myf5+
Changes during obesity	Hyperplasia, hypertrophy, Immune cell infiltration, secretion of vasoconstrictors	"Whitening", loss of UCP1 expression	Potentially resistant to obesity induced inflammation

Figure 3. Function and localization of different adipose tissue depots.

Comparison of white, beige, and brown adipocytes in regard to their localization in specific depots in human and mice. Their major functions and progenitor cells are depicted. Major changes occurring during adipose tissue remodeling in obesity are highlighted. AT indicates adipose tissue; CD, cluster of differentiation; En-1, Engrailed-1; Myf5, myogenic factor 5; Pax7, paired box 7; PDGFRa. platelet-derived growth factor receptor alpha; PVAT, perivascular adipose tissue; and Ucp-1, uncoupling protein-1.

thermogenic adipocytes, and evidence suggests that these cells share properties with both murine brown and beige adipocytes. 23,27,29,36,43 The relative proportion of brown versus beige adipocytes in different human depots in various contexts remains to be fully clarified.<sup>44</sup>

# Stromal Cell Composition of AT and Impact on **Physiology**

Although adipocytes account for most of the volume of AT, they only make up about 50% of the cellular content. 45,46 Other cell types include immune cells such as macrophages, 47-49 lymphocytes, 50-53 eosinophils 54,55 and mast cells,49 as well as fibroblasts, adipocyte precursors, vascular cells, 45 multipotent mesenchymal stem-like cells,56 and nerve processes.57,58 Visceral AT, in contrast to subcutaneous AT, tends to have a higher content of macrophages,<sup>49</sup> regulatory T cells,<sup>52</sup> natural killer T cells,<sup>51</sup> and eosinophils.54 Further, visceral and subcutaneous AT display differences in angiogenesis<sup>59-63</sup> and sympathetic innervation, 58,64,65 which can modulate the propensity for energy storage versus dissipation. Finally, changes in macrophages,66 eosinophils,66-68 and group 2 innate lymphoid cells<sup>69</sup> can regulate browning of AT.

# Perivascular and Epicardial Adipose Tissue

In addition to the well-described white and brown adipose depots, AT is also located around most large blood vessels including the aorta and mesenteric vessels but not the pulmonary and brain vasculature or the

microcirculation<sup>70</sup> (Figure 4). Perivascular adipose tissue (PVAT) is a specialized local deposit of adipose tissue surrounding blood vessels that also provides mechanical protection and regulation of blood vessel tone.71-73 Ex vivo aortic ring experiments revealed a role for PVAT in the relaxation of mesenteric arteries and the thoracic aorta of rats in response to stretch-mediated.74 The contractile response of isolated murine mesenteric arteries toward norepinephrine, on the other hand, is significantly reduced in the presence of PVAT.75 Further, electrical field stimulation assays of mesenteric arteries demonstrated a role for sympathetic nerve activation<sup>76</sup> and sensory neurons<sup>77</sup> in the vasodilatory effects of PVAT. The anticontractile effects of sympathetic stimulation are mediated by the stimulation of  $\beta_3$ -adrenoreceptors in PVAT, and treatment with an antagonist of  $\beta_3$ -adrenoreceptors reduces these effects.<sup>76</sup>

Interestingly, PVAT is itself heterogeneous, with its phenotype strongly location-dependent.78-80 Because of its close proximity to the vasculature and direct contact with the adventitia,81 PVAT is thought to play a role in vascular function and pathology. PVAT surrounding the abdominal aorta and the mesenteric arteries displays a mostly white phenotype in humans82 and mice, with almost no UCP1 expressing thermogenic adipocytes.<sup>28</sup> On the other hand, rodent PVAT surrounding the thoracic aorta has a brown-like phenotype with multilocular adipocytes and UCP1 expression similar to classical brown adipocytes.83-86 This is supported by patterns of BAT detected by positron emission tomography-computed tomography in the para-aortic area and around

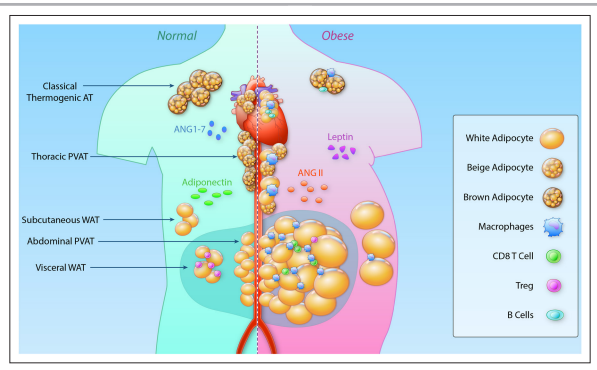


Figure 4. Changes in different adipose tissue depots in homeostasis and during obesity.

In states of normal body weight (left), thermogenic brown and beige adipocytes are found surrounding the thoracic aorta (perivascular adipose tissue [PVAT]) and can be detected in the cervical, supraclavicular, axillary, paraspinal, renal and epicardial area and in infants in the interscapular depot. These cells have a multilocular appearance and due to the high density of mitochondria appear brown. The abdominal aorta and mesenteric vasculature are surrounded by white adipocytes. These unilocular adipocytes are also found in visceral and subcutaneous adipose depots. Adiponectin and angiotensin 1-7 are secreted by adipocytes and have a vasodilating effect on the vasculature. In the lean state, adipose tissue is populated with different immune cells important for homeostasis, which change dramatically during obesity. During obesity (right), T regulatory cells (Treg) are lost in visceral adipose tissue and inflammatory CD8 T cells and macrophages infiltrate the visceral, mesenteric, and to a lesser extent subcutaneous adipose depot. Thermogenic adipose tissue in proximity to the heart and the aorta downregulates thermogenic gene expression and becomes infiltrated with immune cells. Classical brown adipose tissue is potentially protected against obesity-induced immune cell infiltration. Secretion of vasodilatory factors from adipocytes are downregulated whereas leptin and Ang II (angiotensin II) are predominantly secreted, resulting in elevations in blood pressure (Illustration credit: Ben Smith). Ang 1-7 indicates angiotensin 1-7; CD, cluster of differentiation; and WAT, white adipose tissue.

the heart of humans. <sup>87</sup> In addition, autopsy studies of Siberian adults revealed clear UCP1 expression and multilocular and paucilocular appearance of about 40% of mediastinal periaortic vascular AT, with some individuals displaying up to 73%. <sup>88</sup> In mice, long-term moderate cold exposure (16 °C) results in further browning of thoracic PVAT with a markedly increased expression of Ucp1 and  $Pgc1\alpha$  and  $\beta$ . <sup>84</sup>

Thermogenesis of PVAT through cold exposure or genetic manipulation in mice supports a protective role of thoracic PVAT in inflammation and atherosclerosis. Overexpression of the mitochondrial membrane protein MitoNEET induces browning of WAT and thermogenic gene expression. BAT and PVAT prevented mice from an intravascular temperature drop during cold exposure and increased energy expenditure even after removal of interscapular BAT. Further, cold exposure of atherosclerosis-prone ApoE (Apolipoprotein E)-deficient or ApoE-MitoNEET double-deficient mice with removed interscapular BAT.

resulted in reduced atherosclerotic lesion sizes. 84,90 In contrast, lack of PVAT in ApoE-deficient mice with an additional smooth muscle-specific deletion of PPARy (peroxisome proliferator-activated receptor y gamma) increased atherosclerotic lesions and abrogated the protective effects observed after cold exposure. 84 Although the potential contribution of cold-induced browning of WAT was not excluded, these studies imply a contribution of PVAT to whole-body thermogenesis and protection from atherosclerosis.

Several studies in humans have examined the phenotype of perivascular fat surrounding the internal thoracic arteries. While human internal thoracic artery PVAT has been reported to have a white phenotype in one study, it is important to note that 84% of the individuals examined in this study were overweight or obese, which might affect the appearance of their AT.<sup>91</sup> Nevertheless, PVAT of human internal thoracic arteries attenuated the contractile response to the thromboxane A2/prostaglandin H2 receptor agonist U46619 and phenylephrine.<sup>91</sup> Similar effects were observed in PVAT stripped arteries

through the transfer of PVAT-incubated supernatant.91 However, detailed analysis of human thoracic PVAT is limited due to difficulties with sample acquisition and is often isolated from patients with underlying cardiovascular complications, complicating phenotypic assessment.

Despite the close morphological relationship between tp BAT and tPVAT (thoracic PVAT) in mice, proteomics data revealed a depot-specific clustering and an only 43% overlap of their proteome on a standard diet.92 This is comparable to the overlap of 44% of detected proteins between tPVAT with visceral WAT or the overlap of 53% between visceral WAT and BAT, 2 very distinct depots with different functions<sup>92</sup> suggesting a potentially unique PVAT composition. Interestingly, PVAT has been shown to regulate vascular tone83,93 through contact dependent and paracrine functions that are impaired during obesity in mice and humans. 91,94,95 The contractile response of mesenteric arteries to norepinephrine, for example, is reduced in the presence of PVAT but compromised in diet-induced obesity.95 Further, the expression of vasodilatory factors, such as angiotensin (1-7),96-98 adiponectin,75,76 and nitric oxide99 is inhibited during obesity,94,95,99,100 and the expression of the vasoconstrictor Ang II (angiotensin II) is induced in PVAT.70 Finally, a recent single-cell RNA sequencing study demonstrated the existence of 2 main clusters of mesenchymal stem/stromal cells in PVAT of the thoracic aorta of mice.<sup>101</sup> One of the clusters was associated with angiogenic and adipogenic potential, whereas the other cluster was

enriched for genes associated with vascular smooth muscle cell differentiation. 101 Transplantation of those PVAT-derived mesenchymal stem/stromal cells to a vein graft model significantly promoted neointima formation demonstrating a possible role of PVAT in vascular remodeling.<sup>101</sup>

PVAT is an important contributing factor to hypertension, 18,19 endothelial dysfunction, 102 and other vascular abnormalities in obesity71-73,94,103,104 (Figure 5). PVAT normally releases vasodilatory mediators, including adiponectin, 75,76 and yet to be fully characterized molecules often acting on K+ channels, which exert an anticontractile activity and promote vascular relaxation.70 However, in the setting of obesity and insulin resistance, oxidative stress and inflammation are increased in PVAT, thereby resulting in an increase in proinflammatory adipokines including TNF $\alpha$  (tumor necrosis factor alpha), and (ILs) interleukins (IL-6 and IL-8), leading to vascular insulin resistance, impaired relaxation, and vascular stiffness.<sup>71</sup> IL-6 and TNF $\alpha$  also attenuate the vasodilation of mesenteric arteries ex vivo.94 Other cytokines such as IL-18 are thought to have protective effects on PVAT and vascular function, and loss of IL-18 results in elevated blood pressure in mice associated with the whitening of thoracic PVAT.<sup>105</sup> However, the specific impact of IL-18 in PVAT needs to be addressed in AT-specific conditional knock out animals. The Framingham Offspring and Third Generation cohort studies showed that increased PVAT volume is associated with higher thoracic and abdominal

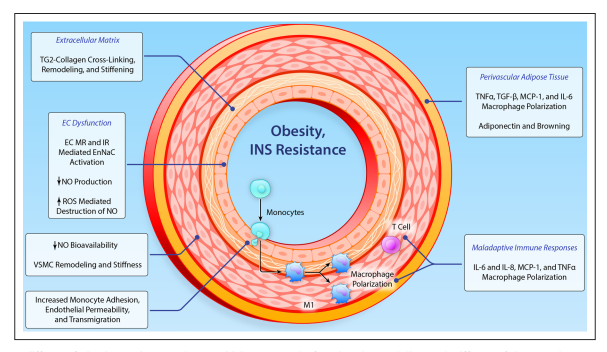


Figure 5. Effects of obesity on the vasculature which promote dysfunctional remodeling and stiffness of the vasculature (Illustration credit: Ben Smith).

Ang II indicates angiotensin II; CRP, C-reactive protein; EC, endothelial cell; ENaC, epithelial Na+ channel; IL, interleukin; IR, insulin receptor; MCP-1, monocyte chemotactic protein-1; MMP, matrix metalloproteinase; MR, mineralocorticoid receptor; NO, nitric oxide; TG2, tissue transglutaminase; TGF-β, transforming growth factor-β; TNF, tumor necrosis factor; and VSMC, vascular smooth muscle cell.

aortic dimensions and increased arterial stiffness, even after adjusting for age and CVD risk factors including BMI and visceral AT volume. 104

The heart is also directly associated with specific AT depots. Epicardial AT is located on the surface of the myocardium in direct contact with the coronary arteries, and pericardial AT is in contact with the pericardial sac. 106 Under physiological conditions, epicardial AT may supply energetic substrates to the heart and has a greater capacity for free fatty acid turnover than other visceral AT depots.<sup>107</sup> Although its cold-induced UCP1 expression does not reach levels of classical BAT,28 human epicardial AT has a thermogenic phenotype and has been suggested to regulate the temperature of the myocardium. 108 Other studies described the portion of epicardial AT surrounding the coronary arteries in humans as a white-like depot despite the expression of some classical brown fat marker genes such as UCP1, PRDM16, and CPT1B.109 The same study found a lower expression of adipogenic marker genes PPARy, FABP4 (fatty acid-binding protein 4) and C/EBPα (enhancer binding protein alpha) but an increased expression of proinflammatory cytokines compared with subcutaneous AT.<sup>109</sup> This discrepancy might be explained by the reported whitening of epicardial AT after birth in humans, with only a distinct subset of multilocular UCP1 positive cells.110 Epicardial AT secretes polypeptides, such as adiponectin<sup>111</sup> and adrenomedullin,<sup>112</sup> which have cardioprotective effects, with low expression of adiponectin in epicardial AT being associated with hypertension.113 Healthy epicardial AT accounts for ≈5% to 20% of the heart weight, 114 and the thickness of epicardial AT is increased in hypertensive individuals. 115-117 Under pathological conditions, epicardial AT becomes infiltrated with immune cells expressing proinflammatory genes (IL-1 $\beta$ , IL-6, and TNF $\alpha$ )<sup>118</sup> and can contribute to structural changes in the heart.119-121 Studies from epicardial AT derived from coronary artery bypass grafts showed significantly lower adiponectin expression compared with other visceral adipose depots and a marked increase in CD45 expression, suggesting increased immune cell infiltration compared with omental AT.122 Studies of mild cold exposure in humans and the analysis of epicardial AT could be beneficial to understanding the role of epicardial AT thermogenesis for CVD.110 Since mice do not have a comparable epicardial AT depot, a mechanistic understanding of how epicardial AT contributes to blood pressure modulation is lacking.

# CARDIOVASCULAR CONSEQUENCES OF OBESITY AND ADIPOSE TISSUE DYSFUNCTION

# Impact of AT on Blood Pressure Regulation

One of the central modes of blood pressure regulation is via the renin-angiotensin-aldosterone system (RAAS).

Its major bioactive component Ang II is produced from its precursor angiotensinogen by the activation of angiotensin-converting enzymes 1 and 2. Angiotensin-converting enzyme 2 can further process Ang II to generate angiotensin 1-7, which has vasodilatory properties.96-98 Ang II<sup>83</sup> and aldosterone are also secreted by adipocytes and can directly activate vascular smooth muscle cells (VSMCs) via the angiotensin type 1 receptor. 123 Ang II is a prominent regulator of vascular tone, 124 and its expression is spatially regulated in PVAT, with higher expression in mesenteric PVAT compared to thoracic PVAT.83 Interestingly, studies in rats have demonstrated that fasting reduces angiotensingen expression in visceral AT, whereas refeeding significantly induces its expression and results in elevated blood pressure. 125 A similar effect can be observed by overexpression of angiotensinogen in mice, which also results in hypertension. 126

All of the components of the RAAS are also secreted by human WAT.127 However, there are conflicting data as to whether the basal expression of RAAS components differ in visceral and subcutaneous AT in lean individuals. One study reported a higher general expression of angiotensinogen, the precursor of Ang II, in visceral AT compared with subcutaneous AT.<sup>128</sup> A more recent, larger study, however, reported no changes in angiotensinogen expression between the 2 depots in lean individuals.<sup>129</sup> Nevertheless, visceral AT expressed higher amounts of renin, angiotensin-converting enzyme 2 and both angiotensin receptor types 1 and 2 in the same study, whereas ACE1 was not changed. 129 In rats, mesenteric PVAT expresses higher levels of Ang II and both angiotensin receptor subtypes than thermogenic thoracic PVAT.83 This is in line with the reported downregulation of angiotensinogen after  $\beta$ -adrenergic stimulation of murine adipocytes in vitro. 130

Thermogenic brown and beige AT is considered to have protective effects on the vasculature, as individuals with detectable thermogenic AT have lower odds for hypertension and coronary artery disease relative to individuals without thermogenic AT.24 Moreover, coding variants in PRDM16, the master regulator of thermogenic AT, are associated with hypertension in humans. 131 Interestingly, components of the RAAS cascade can directly affect AT, and angiotensin 1-7, besides its vasodilatory actions<sup>96-98</sup> also induces BAT and reduces diet-induced obesity in mice. 132,133 Surprisingly, pharmacological activation of angiotensin receptor 2 and Ang II treatment can induce browning of subcutaneous white adipocytes in vivo and stimulation of brown precursor differentiation in vitro. 134,135 This protective impact on BAT is assumed to be either mediated by increased sympathetic nerve activation 135 or through increased conversion of Ang II to angiotensin 1-7. Moreover, deletion of the type 1 angiotensin receptor results in increased appearance of multilocular beige adipocytes. 136 Taken together, it seems that angiotensin 1-7 and activation of the angiotensin

receptor 2 or inhibition of the type 1 angiotensin receptor can stimulate BAT, which in turn has beneficial effects on blood pressure and attenuates development of CVD. Further studies will be needed to investigate the direct impact and molecular basis of the protective impact of thermogenic AT on hypertension.

# **Adipose Tissue Remodeling During Obesity**

Obesity results in a chronic low-grade inflammatory state in adipose tissue. 137,138 Visceral obesity, in particular, is strongly associated with the development of CVD.13,14 Defining and understanding remodeling of different AT depots during obesity is thus of utmost importance to ultimately preventing deleterious sequelae. During obesity, AT can expand by either enlargement of existing adipocytes (hypertrophy) or by increasing the number of adipocytes (hyperplasia) (Figure 3), with the relative importance of either mechanism varying based on depot, sex, and age.31 At baseline, fed a standard diet, neither visceral nor subcutaneous AT exhibit significant new adipogenesis in adult humans or mice. 31,139 Long-term high fat feeding of mice, on the other hand, resulted in increased adipogenesis and hypertrophy in the visceral AT, including mesenteric PVAT, whereas subcutaneous AT adapts to the higher energy intake by hypertrophy.31 The individual impact of hypertrophy versus hyperplasia in the development of the metabolic syndrome is still under debate. 140 Maximum hypertrophy in adipocytes in established obese conditions can result in the exhaustion of the lipid storing capacity in adipocytes, which in turn can induce ectopic storage of fat in other organs such as the liver, supporting the development of the metabolic syndrome.141 On the other hand, visceral AT, despite its ability to expand by hyperplasia is more susceptible to AT inflammation, which in turn contributes to metabolic and CVD outcomes. 142 Sex-dependent differences in AT distribution have been reviewed elsewhere, 143-145 but in short, females most often accumulate AT in the subcutaneous depot, whereas men and postmenopausal women tend to accumulate AT in central visceral depots. 143 Hormone replacement therapy in postmenopausal women prevents this central AT distribution, 146 highlighting the role of sex hormones in fat distribution. However, recent studies, using an elegant separation of gonadal sex and sex chromosomes demonstrated that the XX chromosomal sex results in increased weight gain independent of the gonadal sex.147,148 This was mediated through the X-chromosome-escaped dose-dependent expression differences of the histone demethylase KDM5C (lysine-specific demethylase 5C) in females compared with males, and lowering KDM5C levels in females to the same extend seen in males resulted in weight loss and reduced body fat content.148

In obesity, the immune cell composition of different AT depots demonstrates dynamic changes 70,142,149 (Figure 4). For example, adipose tissue macrophages

increase in obesity and their ablation improves insulin sensitivity and reduces inflammation.<sup>47,150-152</sup> The recruitment<sup>47</sup> and proliferation<sup>153</sup> of proinflammatory macrophages during obesity is greater in visceral than in subcutaneous AT.154,155 Obesity further results in the loss of protective CD4 helper<sup>156</sup> and regulatory T cells<sup>52,157</sup> and in the enrichment of CD8 T cells in visceral AT.53 These variations in immune cell infiltration between visceral and subcutaneous AT result in a low-grade inflammatory environment that can contribute to CVD. 158,159 Recently, eosinophils have gained attention for their role in promoting beige adipocyte activation, 67,68 and their loss during obesity, especially in visceral and mesenteric AT, renders mice susceptible to diet-induced obesity54 and abolishes the anticontractile effect of PVAT to norepinephrine.95 However, some of these findings require further clarification and together with detailed information on PVAT immune cell content and changes during obesity are discussed elsewhere.70

Thermogenic brown and beige fat, on the other hand, have antiobesity effects in humans, 160,161 and depletion of UCP1 itself or ablation of UCP1 expressing thermogenic AT results in weight gain. 162,163 In contrast to WAT, classical BAT of obese mice expresses lower levels of genes associated with immune cells, suggesting that thermogenic AT is resistant to diet-induced inflammation.86 However, other studies have shown that macrophages<sup>164</sup> and B lymphocytes<sup>165</sup> infiltrate thermogenic AT during obesity, and together with increased inflammatory cytokines<sup>109</sup> are thought to suppress UCP1 expression in brown adipocytes. 164 Further, mice fed a high fat diet for 12 weeks show reduced expression of some thermogenic marker genes, and adipocytes shifted from a multilocular to an unilocular appearance with increased lipid accumulation in BAT and thoracic PVAT.92 The increased body and PVAT weight also impairs anticontractile effects of PVAT.91 High fat feeding further results in the upregulation of Notch1 specifically in thoracic PVAT compared with WAT or BAT.92 Genetic adipocyte-specific induction of Notch1 resulted in morphological changes of tPVAT comparable to high fat diet-induced effects.92 This is supported by another study showing that adipocyte-specific overexpression of Notch1 impairs thermogenesis and insulin sensitivity and results in whitening of classical BAT, whereas pharmacological inhibition of Notch1 results in browning of WAT and ameliorates high fat diet-induced obesity. 166

# Remodeling of AT During Obesity and Its Impact on Blood Pressure Homeostasis

Obesity is strongly associated with the development of hypertension,13 a major risk factor for CVD morbidity and mortality.167,168 Compared with normal weight individuals, obese individuals also carry a greater risk for coronary artery calcification, carotid artery intimal media thickening, and left ventricular hypertrophy, even after adjustment for traditional CVD risk factors. <sup>169</sup> Weight reduction significantly improves blood pressure, <sup>19,170,171</sup> and therefore, suggests a direct link between AT phenotype and odds of developing CVD and hypertension. Visceral obesity in rodents and humans is particularly associated with the metabolic syndrome, <sup>172</sup> which consists of several risk factors for CVD, including hypertension. <sup>173</sup> On the other hand, humans with thermogenic AT have lower odds for hypertension, coronary artery disease, and congestive heart failure, even when obese. <sup>24</sup>

Angiotensinogen expression is significantly elevated in obese individuals and is also higher in visceral AT compared with subcutaneous AT128,174,175 (Figure 4). Interestingly, expression of Ang II is increased in subcutaneous AT in obese individuals with hypertension compared with normotensive obese individuals.<sup>128</sup> Diet-induced obesity did not affect angiotensinogen levels in BAT, liver, kidney, or heart in wild-type mice or in mice expressing the human angiotensinogen gene under its own promoter.<sup>175</sup> Importantly, adipocyte-specific deletion of angiotensinogen prevents increased Ang II in the circulation and blocks elevation of BP in obese mice, 176 suggesting a direct impact of AT-derived angiotensinogen on blood pressure. Moreover, angiotensin receptor type 1 inhibition reverses obesity-induced blood pressure elevation in rats.<sup>177</sup> Finally, angiotensinogen levels are negatively regulated by PRDM16, and deletion of PRDM16 and ablation of beige adipocytes results in increased angiotensinogen expression. 178,179 Ablation of BAT in mice results in obesity as well as elevated blood pressure 180; however, whether this is a consequence of obesity induced changes in RAAS or can be directly linked to factors secreted by brown AT needs to be further determined. Aldosterone, another component of the RAAS secreted by adipocytes, 123 also positively correlates with BMI, and weight loss reduces serum aldosterone levels and reduces hypertension.<sup>181</sup> Components of the RAAS can therefore affect VSMC and endothelial dependent regulation of vascular tone, both of which are adversely affected during obesity.

Leptin, an adipocyte-derived hormone that regulates food intake and energy expenditure, is significantly increased in obesity in mice and humans<sup>182,183</sup> (Figure 4). In contrast to angiotensinogen, it may be expressed at higher levels in subcutaneous than in visceral AT,<sup>184-186</sup> and its expression is correlated with adipocyte size.<sup>185</sup> Nevertheless, diet-induced obesity results in elevated leptin levels and attendant increases in heart rate and blood pressure in rodents.<sup>92,187,188</sup> This induction is mediated by a leptin-stimulated increase in sympathetic nerve activity,<sup>189,190</sup> and antibody blockade of leptin or inhibition of leptin receptors on hypothalamic neurons normalized blood pressure in obese rodents.<sup>187</sup> Finally, leptin-deficient mice<sup>191</sup> and humans with loss of function mutations in leptin or the leptin receptor have lower blood pressure

despite severe obesity.<sup>187</sup> It is not well understood how the chronic increase of leptin in obese subjects results in leptin-resistance<sup>192</sup> and whether this affects blood pressure. Based on the abovementioned data, reduced leptin signaling ameliorates blood pressure in mice, and therefore, leptin-resistant obesity should be beneficial in regard to blood pressure. Indeed, leptin also has some vasodilatory effects in healthy rodents, via induction of nitric oxide expression in endothelial cells (ECs)77,193 and in healthy humans by a mechanism independent of nitric oxide. 194 Further, leptin resistance was demonstrated to selectively affect neurons in the hypothalamus that regulate food intake, while affecting other neuronal circuits to a lesser extent, 195,196 which could explain how obese individuals do not have beneficial effects on blood pressure when leptin resistant. In detail, agouti obese mice were resistant to food intake and body weight effects of systemic leptin administration but had a preserved induction of leptin-induced renal sympathetic activation. 196,197 Similar results in diet-induced obese mice showed the preservation of leptin-induced renal sympathetic activation and blood pressure regulation despite the resistance to weight-reducing actions of leptin.<sup>188</sup>

Resistin is enriched in visceral AT,<sup>198</sup> including epicardial AT<sup>199</sup> and PVAT,<sup>200</sup> and is markedly increased during obesity.<sup>200,201</sup> Resistin has an important role in type 2 diabetes and insulin resistance in mice.<sup>201</sup> In humans with type 2 diabetes, resistin expression was only elevated in combination with hypertension and not in patients without hypertension.<sup>202</sup> In hypertensive patients without type 2 diabetes, resistin levels did not correlate with blood pressure indicating a more complex connection of obesity, insulin resistance, and blood pressure regulation by resistin. In mice, resistin treatment induced hypertension through the induction of angiotensinogen.<sup>203</sup> Finally, resistin treatment of isolated human VSMC similar to angiotensin, resulted in increased proliferation.<sup>204</sup>

Visfatin is also expressed in visceral AT, including PVAT,  $^{200}$  and increased through hypoxia-induced expression of HIF1 $\alpha^{205}$  in obesity.  $^{200}$  Hypertensive patients have elevated serum visfatin levels  $^{206}$ ; however, newly diagnosed, nonobese hypertensive men did not show any association of plasma visfatin levels and hypertension.  $^{207}$  Importantly, visfatin is mostly enriched in adipose tissue macrophages in mice  $^{200}$  and humans,  $^{208}$  and therefore, its role in adipocyte specific regulation of blood pressure might be a secondary cause of increased immune cell infiltration in obesity. Nevertheless, it was shown that hypoxic conditions can induce visfatin in murine adipocyte cell lines and its adipocyte specific role in blood pressure regulation should be determined by adipocyte-specific deletion of visfatin.

Adiponectin is another endocrine factor secreted by AT that tends to be reduced during obesity<sup>209,210</sup> (Figure 4). In humans, visceral adiposity inversely correlates with adiponectin secretion, whereas secretion of adiponectin by

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subcutaneous AT is not affected by adiposity.<sup>209</sup> Serum adiponectin levels are reduced in obese individuals with hypertension,211 and lifestyle intervention212 or antihypertensive therapy<sup>211</sup> resulted in increased adiponectin levels and improved blood pressure.<sup>212</sup> In addition, lower adiponectin levels correlate with the risk for development of hypertension in humans, 213,214 independent of body fat distribution.<sup>215</sup> Mice on a standard diet that lack adiponectin display elevated blood pressure despite similar body weight, 6 whereas adiponectin overexpression in obese mice ameliorates elevated blood pressure.210 To understand the direct impact of adiponectin without secondary metabolic effects such as insulin resistance, mice lacking adiponectin were fed a high salt diet. These mice developed hypertension, which could be rescued by adiponectin administration.<sup>210</sup> The observed elevation in blood pressure was associated with reduced eNOS (endothelial nitric oxide synthase) and prostaglandin la synthase,210 indicating a role for adiponectin in EC-mediated vasodilation.<sup>216</sup> Further, ex vivo stimulation of murine mesenteric arteries with norepinephrine was significantly reduced in the presence of PVAT or PVAT-derived supernatant and could be blocked by adiponectin blocking peptide or in vessels derived from adiponectin-deficient mice.95 Adiponectin blocking peptide also blocked electrical field stimulation of mesenteric arteries depending on the presence of PVAT.76 Adiponectin treatment of isolated mesenteric arteries stripped of PVAT restores the anticontractile effects,75,76 depending on the vascular large-conductance Ca2+-activated K+ channel on VSMC.<sup>75</sup> Finally, AMPKα1 (5'AMP-activated protein kinase catalytic subunit alpha 1)-deficient mice secrete less adiponectin, and ex vivo stimulation of thoracic aortic rings from these mice displayed an impaired vasodilatory effect of PVAT after U46619 treatment.217

Another factor enriched in human omental AT and detected in human serum is omentin.<sup>218</sup> Like adiponectin, it is reduced in obese conditions<sup>219</sup> and induced through weight reduction.<sup>220</sup> In rats, omentin treatment ameliorates Ang II or noradrenalin-induced hypertension and reduces blood pressure in normotensive rats.<sup>221,222</sup> Interestingly, omentin suppressed inflammatory mediators in various vascular cell types<sup>222-224</sup> and induced adiponectin levels, which might result in the indirect regulation of blood pressure. This is also the case for adipolin, 225 which is reduced in obese mice<sup>226</sup> and has a protective role in vascular remodeling through the inhibition of VSMC proliferation and macrophage activation,<sup>227</sup> and although associated with protective effects on CVD, its role in regulation of blood pressure needs to be further determined.

Several other factors secreted by different adipose tissue depots have been associated with a role in blood pressure regulation; however, functional and mechanistical proof is still sparse and will be required to understand the independent impact of those AT-derived mediators in the regulation of hypertension. IL-33

(Interleukin-33), for example, plays a pivotal role in the activation of eosinophils, and genetic loss or obesityinduced reduction of eosinophils in PVAT results in a reduced anticontractile response.95 Further, activation of eosinophils by IL-33 treatment rescues obesityinduced high blood pressure to the level of control mice, dependent on an EC and nitric oxide synthase-mediated effect.<sup>228</sup> Of note, patients with pulmonary hypertension showed elevated IL-33 levels, 229 and deficiency of the IL-33 receptor attenuates the progression of pulmonary arterial hypertension in mice.230 Therefore, IL-33 could play a differential role in blood pressure regulation of the vasculature with and without PVAT.

#### Vascular Stiffening and CVD Risk

While vascular stiffening is a normal phenomenon with increasing age, obesity, and associated insulin resistance accelerates this process. To this point, a population study showed that skin-fold thickness is a predictor of arterial stiffness in hypertensive patients.231 Another study found an association between abdominal obesity and increased vascular stiffness. 232,233 Epidemiological studies have demonstrated that hyperinsulinemia or insulin resistance, as present in overweight and obese individuals, is an independent risk factor for vascular stiffening. This vascular stiffening in association with obesity and insulin resistance has been observed in all age groups, including children. 234,235

There is considerable evidence that the vascular stiffening that is increased in obesity is a powerful risk factor for CVD. Data from the Framingham Heart Study have established an increased incidence of CVD events with increasing weight in both men and women,8 and CVD has been strongly associated with vascular stiffness.<sup>235,236</sup> Importantly, arterial stiffening is especially striking in obese and diabetic premenopausal females who tend to lose the normal protection afforded by female sex hormones against vascular disease and show an increase in CVD events relative to lean, nondiabetic, age-matched women.<sup>237</sup> Indeed, vascular stiffness independently predicts heart disease, cerebrovascular disease, and renal disease, as increased vascular stiffness is significantly associated with damage to target organs such as the heart, kidney, and brain.238 For example, stiffening of central arteries increases systolic pressure and decreases diastolic pressure, resulting in increased pulse pressure and afterload leading to an increase in left ventricular mass and myocardial oxygen demand. Further, the decrease in diastolic pressure is associated with reduced coronary blood flow during diastole. These changes have been consistently associated with left ventricular remodeling and fibrosis together with left ventricular diastolic dysfunction and associated heart failure with preserved systolic function<sup>239,240</sup> (Figure 5). While early detection of arterial stiffening in obese individuals certainly helps to

identify a powerful risk factor for CVD, definitive studies on the impact of weight loss on reversal of vascular stiffness have yet to be conducted.

#### **Mechanisms in CV Stiffness With Obesity**

Development of arterial stiffness is a complex process that is driven by the interaction of endocrine factors and AT-derived cytokines, as well as interactions between different vascular cellular components, the ECM (extracellular matrix), PVAT, and immune cells in the vasculature. 6,94 The paragraphs that follow focus on mechanisms involved in CV stiffness in conditions of overnutrition and obesity. This includes a discussion of the role of vascular endothelial abnormalities, which lead to impaired eNOS activation and associated increases in vascular stiffness. We also discuss the emerging role of vascular cell-specific mineralocorticoid and insulin receptor activation in promoting endothelial stiffness via endothelial Na+ channel (EnNaC) activation, and the impact of a decrease in bioavailable nitric oxide (NO) in mediating vascular stiffness in diet-induced obesity (Figure 5).

Arterial stiffness in obesity is associated with structural and functional changes in the intimal, medial, and adventitial layers of the vasculature.<sup>241</sup> Arterial stiffness is regulated by plasma factors such as aldosterone and insulin, as well as factors derived from the different layers of the vascular wall. Moreover, interactive signaling between different cells of the vascular wall modulates structure and function of cellular and noncellular components. Increased arterial stiffness in obese and insulin resistant states has been related to mechanisms related to both EC and VSMC stiffness, leading to the use of such terms as the stiff EC syndrome<sup>241-243</sup> and the smooth muscle stiffness syndrome.<sup>242</sup> In addition to the role of ECs and VSMCs, vascular adipose and immune cell dysfunction and ECM remodeling contribute to obesity-associated arterial stiffness. This underscores the importance of understanding the complex cellular and ECM interactions that contribute to obesity-associated arterial stiffness.243,244

Increased plasma insulin and aldosterone levels lead to heightened activation of vascular MRs and IRs in obesity and insulin resistance states. <sup>239–243</sup> Further, a downstream mediator of mineralocorticoid and insulin receptor activation, the ion channel EnNaC, has recently been identified as a key molecular determinant of endothelial dysfunction and CV fibrosis and stiffening. <sup>239,243</sup> Increased activity of EnNaC results in a number of negative consequences including stiffening of the cortical actin cytoskeleton in ECs, impaired endothelial NO release, increased oxidative stress meditated NO destruction, increased vascular permeability and stimulation of an inflammatory environment. Such endothelial alterations impact vascular function and stiffening through increases in vascular constriction

and stimulation of tissue remodeling including fibrosis. In the case of the myocardium, obesity and associated elevations in aldosterone and insulin are associated with coronary vascular endothelial stiffening and related reductions in bioavailable NO leading to heart failure with preserved systolic function.

Recent studies, conducted in female mice fed a diet high in refined carbohydrates and saturated fat showed increased endothelial and aortic stiffness, impaired endothelial-dependent vasorelaxation, aortic fibrosis, aortic oxidative stress, and increased vascular expression of EnNaC.<sup>239–241</sup> To gain further insight into the vascular role played by EnNaC, we have characterized a mouse model with EC-specific deletion of the  $\alpha$ , pore-forming, subunit of EnNaC.241 Obesogenic diet-induced abnormalities, along with vascular and cardiac remodeling and fibrosis, were all significantly attenuated in mice with deletion of EnNaC.241-243 From a mechanistic standpoint, these studies showed that diet-induced obesity resulted in a heightened inflammatory response that was associated with reduced eNOS activation and NO production and bioavailability. These latter events likely emanated from increased EnNaC activity leading to polymerization of cortical actin fibers, subsequently reducing eNOS activity, and decreasing NO production leading to increased vascular stiffness (Figure 5). This research has further revealed that activation of the endothelial Na+ channel by aldosterone and insulin leads to endothelial cortical stiffening, impaired NO production, and subsequent vascular fibrosis and stiffening in diet induced obesity.<sup>244,245</sup> Additionally, these observations in this obese mouse model also suggest that activation of the endothelial Na+ channel in the coronary vasculature promotes myocardial fibrosis, myocardial stiffening, and impaired diastolic relaxation and heart failure with preserved systolic function, a condition that is especially pronounced in obese and insulin resistant females.

Studies performed in epithelial cells have shown that both aldosterone and insulin increase ENaC activity via activation of the ubiquitously expressed SGK-1 (serum and glucocorticoid regulated kinase 1).246 Very recent work has shown that SGK-1 represents a point of convergence for insulin and aldosterone signaling in ECs.<sup>244</sup> Consistent with this notion, our preliminary studies have shown that aldosterone and insulin induced increases in EnNaC activity are diminished in isolated ECs from SGK-1 global knock-out mice compared with those of wild-type controls.<sup>244</sup> It is also of relevance that evidence exists in humans for SGK-1 playing an important integrative role in the development of the cardiometabolic syndrome. Specifically, an SGK-1 gain of function gene variant that exists in 5% of the population is associated with increased blood pressure and obesity<sup>247</sup> and has a particularly strong effect in increasing blood pressure in states of hyperinsulinemia and obesity.<sup>247</sup> Further, in rodent models, hyperinsulinism sensitizes the blood

pressure to high fructose and salt intake, an effect involving increased activity of SGK-1.248 Indeed, SGK-1-knockout mice are protected against salt-induced hypertension in the context of obesity caused by a high-fat and highfructose diet.<sup>248</sup> Finally, increased SGK-1 activity in obesity and hypertension has also been demonstrated in adipocytes<sup>249</sup> and immune cells.<sup>250</sup> Thus, multiple lines of evidence point toward important contributions of SGK-1 signaling in promoting the cardiometabolic syndrome, vascular stiffness, and associated CVD in obesity.

In summary, obesity is increasing in prevalence, and these increases in obesity are associated with increased consumption of refined carbohydrates and saturated fat and reduced physical activity. These and other environmental factors interact with genetic and epigenetic factors to promote obesity and related CVD (Figure 1). Obesity also negates the CVD protection normally afforded in premenopausal women. The earliest sign of obesity-related CVD is impaired NO-mediated relaxation, which leads to CV stiffness. Recent studies indicate that insulin and mineralocorticoid receptor activation of the EnNaC is important in the pathogenesis of CV stiffness, especially in obese females who lose the protection against CVD normally afforded in premenstrual women.

#### UNANSWERED QUESTIONS AND FUTURE **DIRECTIONS**

While recent research has highlighted key links between obesity, adipose tissue, and vascular function, a number of important unanswered questions remain. From a basic standpoint, a more complete understanding of the developmental origin and cellular and molecular components of perivascular fat is necessary. Moreover, a comprehensive inventory of the secreted polypeptides and metabolites released by adipose tissues in normal physiology and the obese state will help further illuminate how excess adiposity contributes generally to vascular dysfunction and more specifically to the pathogenesis of hypertension and vascular stiffening. Future studies will also need to uncover the role of environment, genetics, epigenetics, and the microbiome on modulating the interactions between adipose tissues and the vasculature.

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#### Sources of Funding

M. Koenen was supported by the Women and Science Initiative at Rockefeller University; P. Cohen was supported by the Sinsheimer Foundation; M.A. Hill and J.R. Sowers were supported by the National Institutes of Health.

#### **Disclosures**

#### **REFERENCES**

- 1. Afshin A, Forouzanfar MH, Reitsma MB, Sur P, Estep K, Lee A, Marczak L, Mokdad AH, Moradi-Lakeh M, Naghavi M, et al; GBD 2015 Obesity Collaborators. Health effects of overweight and obesity in 195 countries over 25 years. N Engl J Med. 2017;377:13-27. doi: 10.1056/NEJMoa1614362
- Gregg EW, Shaw JE. Global health effects of overweight and obesity. N Engl J Med. 2017;377:80-81. doi: 10.1056/NEJMe1706095
- World Health Organization. Obesity and Overweight. Accessed January 11, 2021. https://www.who.int/news-room/fact-sheets/detail/obesity-andoverweight
- 4. Ward ZJ, Bleich SN, Cradock AL, Barrett JL, Giles CM, Flax C, Long MW, Gortmaker SL. Projected U.S. state-level prevalence of adult obesity and severe obesity. N Engl J Med. 2019;381:2440-2450. doi: 10.1056/NEJMsa1909301
- 5. Kelly T, Yang W, Chen CS, Reynolds K, He J. Global burden of obesity in 2005 and projections to 2030. Int J Obes (Lond). 2008;32:1431-1437. doi: 10.1038/ijo.2008.102
- 6. Guanghong J, Hill MA, Sowers JR. Maternal exposure to high fructose and offspring health. Hypertension. 2019;74:499-501.
- Field AE, Coakley EH, Must A, Spadano JL, Laird N, Dietz WH, Rimm E, Colditz GA. Impact of overweight on the risk of developing common chronic diseases during a 10-year period. Arch Intern Med. 2001;161:1581-1586. doi: 10.1001/archinte.161.13.1581
- 8. Hubert HB, Feinleib M, McNamara PM, Castelli WP. Obesity as an independent risk factor for cardiovascular disease: a 26-year follow-up of participants in the Framingham Heart Study. Circulation. 1983;67:968-977. doi: 10.1161/01.cir.67.5.968
- 9. Franks PW, Hanson RL, Knowler WC, Sievers ML, Bennett PH, Looker HC. Childhood obesity, other cardiovascular risk factors, and premature death. N Engl J Med. 2010;362:485-493. doi: 10.1056/ NEJMoa0904130
- 10. Juonala M, Magnussen CG, Berenson GS, Venn A, Burns TL, Sabin MA, Srinivasan SR, Daniels SR, Davis PH, Chen W, et al. Childhood adiposity, adult adiposity, and cardiovascular risk factors. N Engl J Med. 2011;365:1876-1885. doi: 10.1056/NEJMoa1010112
- 11. Cote AT, Harris KC, Panagiotopoulos C, Sandor GG, Devlin AM. Childhood obesity and cardiovascular dysfunction. J Am Coll Cardiol. 2013;62:1309-1319. doi: 10.1016/j.jacc.2013.07.042
- 12. Lee MJ, Wu Y, Fried SK. Adipose tissue heterogeneity: implication of depot differences in adipose tissue for obesity complications. Mol Aspects Med. 2013;34:1-11. doi: 10.1016/j.mam.2012.10.001
- 13. Garrison RJ, Kannel WB, Stokes J III, Castelli WP. Incidence and precursors of hypertension in young adults: the Framingham Offspring Study. Prev Med. 1987;16:235-251. doi: 10.1016/0091-7435(87)90087-9
- 14. Bhupathiraju SN, Hu FB. Epidemiology of obesity and diabetes and their cardiovascular complications. Circ Res. 2016;118:1723-1735.
- 15. Jia G, Martinez-Lemus LA, Sowers JR. Interaction of adipogenesis and angiogenesis in dietary-induced obesity. Diabetes. 2015;64:2326-2328. doi: 10.2337/db15-0202
- 16. Aroor AR, McKarns S, Demarco VG, Jia G, Sowers JR. Maladaptive immune and inflammatory pathways lead to cardiovascular insulin resistance. Metabolism. 2013;62:1543-1552. doi: 10.1016/j.metabol.2013.07.001
- 17. Piché M-E, Tchernof A, Després J-P. Obesity phenotypes, diabetes, and cardiovascular diseases. Circ Res. 2020;126:1477-1500.
- 18. DeMarco VG, Aroor AR, Sowers JR. The pathophysiology of hypertension in patients with obesity. Nat Rev Endocrinol. 2014;10:364-376. doi: 10.1038/nrendo.2014.44
- 19. Hall JE, do Carmo JM, da Silva AA, Wang Z, Hall ME. Obesity-induced hypertension. Circ Res. 2015;116:991-1006.
- 20. Virtanen KA, Lidell ME, Orava J, Heglind M, Westergren R, Niemi T, Taittonen M, Laine J, Savisto NJ, Enerbäck S, et al. Functional brown adipose tissue in healthy adults. N Engl J Med. 2009;360:1518-1525. doi: 10.1056/NEJMoa0808949
- 21. Leitner BP, Huang S, Brychta RJ, Duckworth CJ, Baskin AS, McGehee S, Tal I, Dieckmann W, Gupta G, Kolodny GM, et al. Mapping of human brown adipose tissue in lean and obese young men. Proc Natl Acad Sci USA. 2017;114:8649-8654. doi: 10.1073/pnas.1705287114
- 22. Zhang F, Hao G, Shao M, Nham K, An Y, Wang Q, Zhu Y, Kusminski CM, Hassan G, Gupta RK, et al. An adipose tissue atlas: an image-guided

- identification of human-like BAT and beige depots in rodents. *Cell Metab.* 2018;27:252.e3–262.e3. doi: 10.1016/j.cmet.2017.12.004
- Lidell ME, Betz MJ, Dahlqvist Leinhard O, Heglind M, Elander L, Slawik M, Mussack T, Nilsson D, Romu T, Nuutila P, et al. Evidence for two types of brown adipose tissue in humans. *Nat Med.* 2013;19:631–634. doi: 10.1038/nm.3017
- Becher T, Palanisamy S, Kramer DJ, Eljalby M, Marx SJ, Wibmer AG, Butler SD, Jiang CS, Vaughan R, Schöder H, Mark A, et al. Brown adipose tissue is associated with cardiometabolic health. *Nat Med*. 2021;27:58-65.
- Klaus S, Casteilla L, Bouillaud F, Ricquier D. The uncoupling protein UCP: a membraneous mitochondrial ion carrier exclusively expressed in brown adipose tissue. *Int J Biochem.* 1991;23:791–801. doi: 10.1016/0020-711x(91)90062-r
- Roesler A, Kazak L. UCP1-independent thermogenesis. Biochem J. 2020;477:709-725. doi: 10.1042/BCJ20190463
- Wu J, Boström P, Sparks LM, Ye L, Choi JH, Giang AH, Khandekar M, Virtanen KA, Nuutila P, Schaart G, et al. Beige adipocytes are a distinct type of thermogenic fat cell in mouse and human. *Cell.* 2012;150:366–376. doi: 10.1016/j.cell.2012.05.016
- Waldén TB, Hansen IR, Timmons JA, Cannon B, Nedergaard J. Recruited vs. nonrecruited molecular signatures of brown, "brite," and white adipose tissues. Am J Physiol Endocrinol Metab. 2012;302:E19–E31. doi: 10.1152/ajpendo.00249.2011
- Young P, Arch JR, Ashwell M. Brown adipose tissue in the parametrial fat pad of the mouse. FEBS Lett. 1984;167:10-14. doi: 10.1016/0014-5793(84)80822-4
- Orava J, Nuutila P, Lidell ME, Oikonen V, Noponen T, Viljanen T, Scheinin M, Taittonen M, Niemi T, Enerbäck S, et al. Different metabolic responses of human brown adipose tissue to activation by cold and insulin. *Cell Metab.* 2011;14:272–279. doi: 10.1016/j.cmet.2011.06.012
- Wang QA, Tao C, Gupta RK, Scherer PE. Tracking adipogenesis during white adipose tissue development, expansion and regeneration. *Nat Med.* 2013;19:1338–1344. doi: 10.1038/nm.3324
- Rosenwald M, Perdikari A, Rülicke T, Wolfrum C. Bi-directional interconversion of brite and white adipocytes. Nat Cell Biol. 2013;15:659–667. doi: 10.1038/ncb2740
- Roh HC, Tsai LTY, Shao M, Tenen D, Shen Y, Kumari M, Lyubetskaya A, Jacobs C, Dawes B, Gupta RK, et al. Warming induces significant reprogramming of Beige, but not brown, adipocyte cellular identity. *Cell Metab.* 2018;27:1121–1137.e5. doi: 10.1016/j.cmet.2018.03.005
- Moulin K, Truel N, André M, Arnauld E, Nibbelink M, Cousin B, Dani C, Pénicaud L, Casteilla L. Emergence during development of the white-adipocyte cell phenotype is independent of the brown-adipocyte cell phenotype. Biochem J. 2001;356:659–664. doi: 10.1042/bj3560659
- Timmons JA, Wennmalm K, Larsson O, Walden TB, Lassmann T, Petrovic N, Hamilton DL, Gimeno RE, Wahlestedt C, Baar K, et al. Myogenic gene expression signature establishes that brown and white adipocytes originate from distinct cell lineages. *Proc Natl Acad Sci USA*. 2007;104:4401–4406. doi: 10.1073/pnas.0610615104
- Xue B, Rim JS, Hogan JC, Coulter AA, Koza RA, Kozak LP. Genetic variability affects the development of brown adipocytes in white fat but not in interscapular brown fat. *J Lipid Res.* 2007;48:41–51. doi: 10.1194/jlr. M600287-JLR200
- Seale P, Bjork B, Yang W, Kajimura S, Chin S, Kuang S, Scimè A, Devarakonda S, Conroe HM, Erdjument-Bromage H, et al. PRDM16 controls a brown fat/skeletal muscle switch. *Nature*. 2008;454:961–967. doi: 10.1038/ nature07182
- Rodeheffer MS, Birsoy K, Friedman JM. Identification of white adipocyte progenitor cells in vivo. Cell. 2008;135:240–249. doi: 10.1016/j.cell.2008.09.036
- 39. Berry R, Rodeheffer MS. Characterization of the adipocyte cellular lineage in vivo. *Nat Cell Biol.* 2013;15:302–308. doi: 10.1038/ncb2696
- Gesta S, Blüher M, Yamamoto Y, Norris AW, Berndt J, Kralisch S, Boucher J, Lewis C, Kahn CR. Evidence for a role of developmental genes in the origin of obesity and body fat distribution. *Proc Natl Acad Sci USA*. 2006;103:6676-6681. doi: 10.1073/pnas.0601752103
- Atit R, Sgaier SK, Mohamed OA, Taketo MM, Dufort D, Joyner AL, Niswander L, Conlon RA. Beta-catenin activation is necessary and sufficient to specify the dorsal dermal fate in the mouse. *Dev Biol.* 2006;296:164–176. doi: 10.1016/j.ydbio.2006.04.449
- Long JZ, Svensson KJ, Tsai L, Zeng X, Roh HC, Kong X, Rao RR, Lou J, Lokurkar I, Baur W, et al. A smooth muscle-like origin for beige adipocytes. Cell Metab. 2014;19:810–820. doi: 10.1016/j.cmet.2014.03.025

- Sharp LZ, Shinoda K, Ohno H, Scheel DW, Tomoda E, Ruiz L, Hu H, Wang L, Pavlova Z, Gilsanz V, et al. Human BAT possesses molecular signatures that resemble beige/brite cells. *PLoS One*. 2012;7:e49452. doi: 10.1371/journal.pone.0049452
- Cannon B, de Jong JMA, Fischer AW, Nedergaard J, Petrovic N. Human brown adipose tissue: classical brown rather than brite/beige? Exp Physiol. 2020;105:1191–1200. doi: 10.1113/EP087875
- 45. Silva KR, Côrtes I, Liechocki S, Carneiro JR, Souza AA, Borojevic R, Maya-Monteiro CM, Baptista LS. Characterization of stromal vascular fraction and adipose stem cells from subcutaneous, preperitoneal and visceral morbidly obese human adipose tissue depots. *PLoS One*. 2017;12:e0174115. doi: 10.1371/journal.pone.0174115
- Kumar RK, Jin Y, Watts SW, Rockwell CE. Naïve, regulatory, activated, and memory immune cells co-exist in PVATs that are comparable in density to non-PVAT fats in health. Front Physiol. 2020;11:58.
- Weisberg SP, McCann D, Desai M, Rosenbaum M, Leibel RL, Ferrante AW Jr. Obesity is associated with macrophage accumulation in adipose tissue. J Clin Invest. 2003;112:1796–1808. doi: 10.1172/JCI19246
- Lumeng CN, Bodzin JL, Saltiel AR. Obesity induces a phenotypic switch in adipose tissue macrophage polarization. J Clin Invest. 2007;117:175–184. doi: 10.1172/JCl29881
- Altintas MM, Azad A, Nayer B, Contreras G, Zaias J, Faul C, Reiser J, Nayer A. Mast cells, macrophages, and crown-like structures distinguish subcutaneous from visceral fat in mice. *J Lipid Res.* 2011;52:480–488. doi: 10.1194/jlr.M011338
- Caspar-Bauguil S, Cousin B, Galinier A, Segafredo C, Nibbelink M, André M, Casteilla L, Pénicaud L. Adipose tissues as an ancestral immune organ: site-specific change in obesity. FEBS Lett. 2005;579:3487–3492. doi: 10.1016/j.febslet.2005.05.031
- Schipper HS, Rakhshandehroo M, van de Graaf SF, Venken K, Koppen A, Stienstra R, Prop S, Meerding J, Hamers N, Besra G, et al. Natural killer T cells in adipose tissue prevent insulin resistance. *J Clin Invest* 2012;122:3343–3354. doi: 10.1172/JCI62739
- Feuerer M, Herrero L, Cipolletta D, Naaz A, Wong J, Nayer A, Lee J, Goldfine AB, Benoist C, Shoelson S, et al. Lean, but not obese, fat is enriched for a unique population of regulatory T cells that affect metabolic parameters. *Nat Med*, 2009;15:930–939, doi: 10.1038/nm.2002
- Nishimura S, Manabe I, Nagasaki M, Eto K, Yamashita H, Ohsugi M, Otsu M, Hara K, Ueki K, Sugiura S, et al. CD8+ effector T cells contribute to macrophage recruitment and adipose tissue inflammation in obesity. *Nat Med.* 2009;15:914–920. doi: 10.1038/nm.1964
- 54. Wu D, Molofsky AB, Liang HE, Ricardo-Gonzalez RR, Jouihan HA, Bando JK, Chawla A, Locksley RM. Eosinophils sustain adipose alternatively activated macrophages associated with glucose homeostasis. *Science*. 2011;332:243–247. doi: 10.1126/science.1201475
- Brigger D, Riether C, van Brummelen R, Mosher KI, Shiu A, Ding Z, Zbären N, Gasser P, Guntern P, Yousef H, et al. Eosinophils regulate adipose tissue inflammation and sustain physical and immunological fitness in old age. *Nat Metab.* 2020;2:688–702. doi: 10.1038/s42255-020-0228-3
- Zuk PA, Zhu M, Ashjian P, De Ugarte DA, Huang JI, Mizuno H, Alfonso ZC, Fraser JK, Benhaim P, Hedrick MH. Human adipose tissue is a source of multipotent stem cells. *Mol Biol Cell*. 2002;13:4279–4295. doi: 10.1091/mbc.e02-02-0105
- Romijn JA, Fliers E. Sympathetic and parasympathetic innervation of adipose tissue: metabolic implications. *Curr Opin Clin Nutr Metab Care*. 2005;8:440–444. doi: 10.1097/01.mco.0000172586.09762.55
- Chi J, Wu Z, Choi CHJ, Nguyen L, Tegegne S, Ackerman SE, Crane A, Marchildon F, Tessier-Lavigne M, Cohen P. Three-dimensional adipose tissue imaging reveals regional variation in beige fat biogenesis and PRDM16dependent sympathetic neurite density. *Cell Metab.* 2018;27:226–236.e3. doi: 10.1016/j.cmet.2017.12.011
- Xue Y, Petrovic N, Cao R, Larsson O, Lim S, Chen S, Feldmann HM, Liang Z, Zhu Z, Nedergaard J, et al. Hypoxia-independent angiogenesis in adipose tissues during cold acclimation. *Cell Metab.* 2009;9:99–109. doi: 10.1016/j.cmet.2008.11.009
- Villaret A, Galitzky J, Decaunes P, Estève D, Marques MA, Sengenès C, Chiotasso P, Tchkonia T, Lafontan M, Kirkland JL, et al. Adipose tissue endothelial cells from obese human subjects: differences among depots in angiogenic, metabolic, and inflammatory gene expression and cellular senescence. *Diabetes*. 2010;59:2755–2763. doi: 10.2337/db10-0398
- Elias I, Franckhauser S, Ferré T, Vilà L, Tafuro S, Muñoz S, Roca C, Ramos D, Pujol A, Riu E, et al. Adipose tissue overexpression of vascular endothelial growth factor protects against diet-induced obesity and insulin resistance. *Diabetes*. 2012;61:1801–1813. doi: 10.2337/db11-0832

- 62. Bagchi M, Kim LA, Boucher J, Walshe TE, Kahn CR, D'Amore PA. Vascular endothelial growth factor is important for brown adipose tissue development and maintenance. FASEB J. 2013;27:3257-3271. doi: 10.1096/fj.12-221812
- 63. Shimizu I, Aprahamian T, Kikuchi R, Shimizu A, Papanicolaou KN, MacLauchlan S, Maruyama S, Walsh K. Vascular rarefaction mediates whitening of brown fat in obesity. J Clin Invest. 2014;124:2099-2112. doi: 10.1172/JCI71643
- 64. Murano I, Barbatelli G, Giordano A, Cinti S. Noradrenergic parenchymal nerve fiber branching after cold acclimatisation correlates with brown adipocyte density in mouse adipose organ. J Anat. 2009;214:171-178. doi: 10.1111/j.1469-7580.2008.01001.x
- 65. Wang P, Loh KH, Wu M, Morgan DA, Schneeberger M, Yu X, Chi J, Kosse C, Kim D, Rahmouni K, et al. A leptin-BDNF pathway regulating sympathetic innervation of adipose tissue. Nature. 2020;583:839-844. doi: 10.1038/s41586-020-2527-y
- 66. Nguyen KD, Qiu Y, Cui X, Goh YP, Mwangi J, David T, Mukundan L, Brombacher F, Locksley RM, Chawla A. Alternatively activated macrophages produce catecholamines to sustain adaptive thermogenesis. Nature. 2011;480:104-108. doi: 10.1038/nature10653
- 67. Qiu Y, Nguyen KD, Odegaard JI, Cui X, Tian X, Locksley RM, Palmiter RD, Chawla A. Eosinophils and type 2 cytokine signaling in macrophages orchestrate development of functional beige fat. Cell. 2014;157:1292-1308. doi: 10.1016/j.cell.2014.03.066
- 68. Knights AJ, Vohralik EJ, Houweling PJ, Stout ES, Norton LJ, Alexopoulos SJ, Yik JJ, Mat Jusoh H, Olzomer EM, Bell-Anderson KS, et al. Eosinophil function in adipose tissue is regulated by Krüppel-like factor 3 (KLF3). Nat Commun. 2020;11:2922. doi: 10.1038/s41467-020-16758-9
- 69. Brestoff JR, Kim BS, Saenz SA, Stine RR, Monticelli LA, Sonnenberg GF, Thome JJ, Farber DL, Lutfy K, Seale P, et al. Group 2 innate lymphoid cells promote beiging of white adipose tissue and limit obesity. Nature. 2015;519:242-246. doi: 10.1038/nature14115
- 70. Saxton SN, Clark BJ, Withers SB, Eringa EC, Heagerty AM. Mechanistic links between obesity, diabetes, and blood pressure: role of perivascular adipose tissue. Physiol Rev. 2019;99:1701-1763. doi: 10.1152/ physrev.00034.2018
- 71. Padilla J, Vieira-Potter VJ, Jia G, Sowers JR. Role of perivascular adipose tissue on vascular reactive oxygen species in type 2 diabetes: a give-and-take relationship. Diabetes. 2015;64:1904-1906. doi: 10.2337/db15-0096
- 72. Jia G, Durante W, Sowers JR. Endothelium-derived hyperpolarizing factors: a potential therapeutic target for vascular dysfunction in obesity and insulin resistance. Diabetes. 2016;65:2118-2120. doi: 10.2337/dbi16-0026
- 73. Jia G, Aroor AR, Sowers JR. The role of mineralocorticoid receptor signaling in the cross-talk between adipose tissue and the vascular wall. Cardiovasc Res. 2017;113:1055-1063. doi: 10.1093/cvr/cvx097
- 74. Watts SW. Flood ED. Garver H. Fink GD. Roccabianca S. A new function for perivascular adipose tissue (PVAT): assistance of arterial stress relaxation. Sci Rep. 2020;10:1807. doi: 10.1038/s41598-020-58368-x
- 75. Lynch FM, Withers SB, Yao Z, Werner ME, Edwards G, Weston AH, Heagerty AM. Perivascular adipose tissue-derived adiponectin activates BKCa channels to induce anticontractile responses. Am J Physiol Heart Circ Physiol. 2013:304:H786-H795.
- 76. Saxton SN, Ryding KE, Aldous RG, Withers SB, Ohanian J, Heagerty AM. Role of sympathetic nerves and adipocyte catecholamine uptake in the vasorelaxant function of perivascular adipose tissue. Arterioscler Thromb Vasc Biol. 2018;38:880-891.
- 77. Abu Bakar H, Robert Dunn W, Daly C, Ralevic V. Sensory innervation of perivascular adipose tissue: a crucial role in artery vasodilatation and leptin release. Cardiovasc Res. 2017;113:962-972. doi: 10.1093/cvr/cvx062
- 78. Police SB, Thatcher SE, Charnigo R, Daugherty A, Cassis LA. Obesity promotes inflammation in periaortic adipose tissue and angiotensin II-induced abdominal aortic aneurysm formation. Arterioscler Thromb Vasc Biol. 2009:29:1458-1464
- 79. Qi XY, Qu SL, Xiong WH, Rom O, Chang L, Jiang ZS. Perivascular adipose tissue (PVAT) in atherosclerosis: a double-edged sword. Cardiovasc Diabetol. 2018;17:134. doi: 10.1186/s12933-018-0777-x
- 80. Padilla J, Jenkins NT, Vieira-Potter VJ, Laughlin MH. Divergent phenotype of rat thoracic and abdominal perivascular adipose tissues. Am J Physiol Regul Integr Comp Physiol. 2013;304:R543-R552. doi: 10.1152/ ajpregu.00567.2012
- 81. Eringa EC, Bakker W, Smulders YM, Serné EH, Yudkin JS, Stehouwer CD. Regulation of vascular function and insulin sensitivity by adipose tissue: focus on perivascular adipose tissue. Microcirculation. 2007;14:389-402. doi: 10.1080/10739680701303584

- 82. Kwok KH, Lam KS, Xu A. Heterogeneity of white adipose tissue: molecular basis and clinical implications. Exp Mol Med. 2016;48:e215. doi: 10.1038/emm.2016.5
- 83. Gálvez-Prieto B, Bolbrinker J, Stucchi P, de Las Heras Al, Merino B, Arribas S, Ruiz-Gayo M, Huber M, Wehland M, Kreutz R, et al. Comparative expression analysis of the renin-angiotensin system components between white and brown perivascular adipose tissue. J Endocrino. 2008;197:55-64.
- 84. Chang L, Villacorta L, Li R, Hamblin M, Xu W, Dou C, Zhang J, Wu J, Zeng R, Chen YE. Loss of perivascular adipose tissue on peroxisome proliferator-activated receptor- $\gamma$  deletion in smooth muscle cells impairs intravascular thermoregulation and enhances atherosclerosis. Circulation. 2012;126:1067-1078. doi: 10.1161/CIRCULATIONAHA.112.104489
- 85. Henrichot E, Juge-Aubry CE, Pernin A, Pache JC, Velebit V, Dayer JM, Meda P, Carlo Chizzolini C, Meier CA. Production of chemokines by perivascular adipose tissue. Arterioscler Thromb Vasc Biol. 2005;25:2594-2599.
- 86. Fitzgibbons TP, Kogan S, Aouadi M, Hendricks GM, Straubhaar J, Czech MP. Similarity of mouse perivascular and brown adipose tissues and their resistance to diet-induced inflammation. Am J Physiol Heart Circ Physiol. 2011;301:H1425-H1437. doi: 10.1152/ajpheart.00376.2011
- 87. Nedergaard J, Bengtsson T, Cannon B. Unexpected evidence for active brown adipose tissue in adult humans. Am J Physiol Endocrinol Metab. 2007;293:E444-E452. doi: 10.1152/ajpendo.00691.2006
- 88. Efremova A, Senzacqua M, Venema W, Isakov E, Di Vincenzo A, Zingaretti MC, Protasoni M, Thomski M, Giordano A, Cinti S. A large proportion of mediastinal and perirenal visceral fat of Siberian adult people is formed by UCP1 immunoreactive multilocular and paucilocular adipocytes. J Physiol Biochem. 2020;76:185-192. doi: 10.1007/s13105-019-00721-4
- 89. Kusminski CM, Park J, Scherer PE. MitoNEET-mediated effects on browning of white adipose tissue. Nat Commun. 2014;5:3962. doi: 10.1038/ ncomms4962
- 90. Xiong W, Zhao X, Garcia-Barrio MT, Zhang J, Lin J, Chen YE, Jiang Z, Chang L. MitoNEET in perivascular adipose tissue blunts atherosclerosis under mild cold condition in mice. Front Physiol. 2017;8:1032. doi: 10.3389/fphys.2017.01032
- 91. Gao YJ, Zeng ZH, Teoh K, Sharma AM, Abouzahr L, Cybulsky I, Lamy A, Semelhago L, Lee RM. Perivascular adipose tissue modulates vascular function in the human internal thoracic artery. J Thorac Cardiovasc Sura. 2005;130:1130-1136. doi: 10.1016/j.jtcvs.2005.05.028
- 92. Boucher JM, Ryzhova L, Harrington A, Davis-Knowlton J, Turner JE, Cooper E, Maridas D, Ryzhov S, Rosen CJ, Vary CPH, et al. Pathological conversion of mouse perivascular adipose tissue by notch activation. Arterioscler Thromb Vasc Biol. 2020;40:2227-2243.
- 93. Gu P, Xu A. Interplay between adipose tissue and blood vessels in obesity and vascular dysfunction. Rev Endocr Metab Disord. 2013;14:49-58. doi: 10.1007/s11154-012-9230-8
- 94. Greenstein AS, Khavandi K, Withers SB, Sonoyama K, Clancy O, Jeziorska M, Laing I, Yates AP, Pemberton PW, Malik RA, et al. Local inflammation and hypoxia abolish the protective anticontractile properties of perivascular fat in obese patients. Circulation. 2009;119:1661-1670.
- 95. Withers SB, Forman R, Meza-Perez S, Sorobetea D, Sitnik K, Hopwood T, Lawrence CB, Agace WW, Else KJ, Heagerty AM, et al. Eosinophils are key regulators of perivascular adipose tissue and vascular functionality. Sci Rep. 2017;7:44571. doi: 10.1038/srep44571
- 96. Brosnihan KB, Li P, Ferrario CM. Angiotensin-(1-7) dilates canine coronary arteries through kinins and nitric oxide. Hypertension. 1996;27:523-528. doi: 10.1161/01.hyp.27.3.523
- 97. Durand MJ, Zinkevich NS, Riedel M, Gutterman DD, Nasci VL, Salato VK, Hijjawi JB, Reuben CF, North PE, Beyer AM. Vascular actions of angiotensin 1-7 in the human microcirculation; novel role for telomerase. Arterioscler Thromb Vasc Biol. 2016;36:1254-1262. doi: 10.1161/ ATVBAHA.116.307518
- 98. Bujak-Gizycka B, Madej J, Wołkow PP, Olszanecki R, Drabik L, Rutowski J, Korbut R. Measurement of angiotensin metabolites in organ bath and cell culture experiments by liquid chromatography - electrospray ionization mass spectrometry (LC-ESI-MS). J Physiol Pharmacol. 2007;58:529-540.
- 99. Gao YJ, Lu C, Su LY, Sharma AM, Lee RM. Modulation of vascular function by perivascular adipose tissue: the role of endothelium and hydrogen peroxide. Br J Pharmacol. 2007;151:323-331. doi: 10.1038/sj.bjp.0707228
- 100. Arita Y, Kihara S, Ouchi N, Takahashi M, Maeda K, Miyagawa J, Hotta K, Shimomura I, Nakamura T, Miyaoka K, et al. Paradoxical decrease of an adipose-specific protein, adiponectin, in obesity. Biochem Biophys Res Commun. 1999;257:79-83. doi: 10.1006/bbrc.1999.0255
- 101. Gu W, Nowak WN, Xie Y, Le Bras A, Hu Y, Deng J, Issa Bhaloo S, Lu Y, Yuan H, Fidanis E, et al. Single-cell RNA-sequencing and metabolomics

- analyses reveal the contribution of perivascular adipose tissue stem cells to vascular remodeling. *Arterioscler Thromb Vasc Biol.* 2019;39:2049–2066.
- 102. Jia G, Sowers JR. Endothelial dysfunction potentially interacts with impaired glucose metabolism to increase cardiovascular risk. *Hypertension*. 2014;64:1192–1193.
- 103. Villacorta L, Chang L. The role of perivascular adipose tissue in vasoconstriction, arterial stiffness, and aneurysm. Horm Mol Biol Clin Investig. 2015;21:137–147. doi: 10.1515/hmbci-2014-0048
- 104. Thanassoulis G, Massaro JM, Corsini E, Rogers I, Schlett CL, Meigs JB, Hoffmann U, O'Donnell CJ, Fox CS. Periaortic adipose tissue and aortic dimensions in the Framingham Heart Study. J Am Heart Assoc. 2012;1:e000885. doi: 10.1161/JAHA.112.000885
- 105. Li W, Jin D, Takai S, Hayakawa T, Ogata J, Yamanishi K, Yamanishi H, Okamura H. Impaired function of aorta and perivascular adipose tissue in IL-18-deficient mice. Am J Physiol Heart Circ Physiol. 2019;317:H1142–H1156. doi: 10.1152/ajpheart.00813.2018
- 106. Talman AH, Psaltis PJ, Cameron JD, Meredith IT, Seneviratne SK, Wong DT. Epicardial adipose tissue: far more than a fat depot. *Cardiovasc Diagn Ther*. 2014;4:416–429. doi: 10.3978/j.issn.2223-3652.2014.11.05
- 107. Marchington JM, Mattacks CA, Pond CM. Adipose tissue in the mammalian heart and pericardium: structure, foetal development and biochemical properties. Comp Biochem Physiol B. 1989;94:225–232. doi: 10.1016/0305-0491(89)90337-4
- 108. Sacks HS, Fain JN, Holman B, Cheema P, Chary A, Parks F, Karas J, Optican R, Bahouth SW, Garrett E, et al. Uncoupling protein-1 and related messenger ribonucleic acids in human epicardial and other adipose tissues: epicardial fat functioning as brown fat. J Clin Endocrinol Metab. 2009;94:3611–3615. doi: 10.1210/jc.2009-0571
- 109. Chatterjee TK, Stoll LL, Denning GM, Harrelson A, Blomkalns AL, Idelman G, Rothenberg FG, Neltner B, Romig-Martin SA, Dickson EW, et al. Proinflammatory phenotype of perivascular adipocytes. Circ Res. 2009;104:541–549.
- Aldiss P, Davies G, Woods R, Budge H, Sacks HS, Symonds ME. 'Browning' the cardiac and peri-vascular adipose tissues to modulate cardiovascular risk. Int J Cardiol. 2017;228:265–274. doi: 10.1016/j.ijcard.2016.11.074
- 111. lacobellis G, Pistilli D, Gucciardo M, Leonetti F, Miraldi F, Brancaccio G, Gallo P, di Gioia CR. Adiponectin expression in human epicardial adipose tissue in vivo is lower in patients with coronary artery disease. *Cytokine*. 2005;29:251–255. doi: 10.1016/j.cyto.2004.11.002
- 112. Silaghi A, Achard V, Paulmyer-Lacroix O, Scridon T, Tassistro V, Duncea I, Clément K, Dutour A, Grino M. Expression of adrenomedullin in human epicardial adipose tissue: role of coronary status. Am J Physiol Endocrinol Metab. 2007;293:E1443–E1450. doi: 10.1152/ajpendo.00273.2007
- 113. Teijeira-Fernandez E, Eiras S, Grigorian-Shamagian L, Fernandez A, Adrio B, Gonzalez-Juanatey JR. Epicardial adipose tissue expression of adiponectin is lower in patients with hypertension. *J Hum Hypertens*. 2008;22:856–863. doi: 10.1038/jhh.2008.75
- 114. Le Jemtel TH, Samson R, Ayinapudi K, Singh T, Oparil S. Epicardial adipose tissue and cardiovascular disease. Curr Hypertens Rep. 2019;21:36. doi: 10.1007/s11906-019-0939-6
- 115. Dicker D, Atar E, Kornowski R, Bachar GN. Increased epicardial adipose tissue thickness as a predictor for hypertension: a cross-sectional observational study. J Clin Hypertens (Greenwich). 2013;15:893–898. doi: 10.1111/jch.12201
- Eroğlu S, Sade LE, Yıldırır A, Demir O, Müderrisoğlu H. Association of epicardial adipose tissue thickness by echocardiography and hypertension. *Turk Kardiyol Dem Ars.* 2013;41:115–122. doi: 10.5543/tkda.2013.83479
- 117. Austys D, Dobrovolskij A, Jablonskienė V, Dobrovolskij V, Valevičienė N, Stukas R. Epicardial adipose tissue accumulation and essential hypertension in non-obese adults. *Medicina (Kaunas)*. 2019;55:456.
- 118. Tomasz M, LiFeng Z, Andrew Z, Mannion JD, Diehl JT, Arafat H, Sarov-Blat L, O'Brien S, Keiper EA, Johnson AG, et al. Human epicardial adipose tissue is a source of inflammatory mediators. *Circulation*. 2003;108:2460–2466.
- Iacobellis G, Ribaudo MC, Zappaterreno A, Iannucci CV, Leonetti F. Relation between epicardial adipose tissue and left ventricular mass. Am J Cardiol. 2004;94:1084–1087. doi: 10.1016/j.amjcard.2004.06.075
- Iacobellis G, Leonetti F, Singh N, M Sharma A. Relationship of epicardial adipose tissue with atrial dimensions and diastolic function in morbidly obese subjects. *Int J Cardiol.* 2007;115:272–273. doi: 10.1016/j. ijcard.2006.04.016
- 121. Thanassoulis G, Massaro JM, O'Donnell CJ, Hoffmann U, Levy D, Ellinor PT, Wang TJ, Schnabel RB, Vasan RS, Fox CS, et al. Pericardial fat is associated with prevalent atrial fibrillation: the Framingham Heart Study. Circ Arrhythm Electrophysio. 2010;3:345–350.

- 122. Baker AR, Silva NF, Quinn DW, Harte AL, Pagano D, Bonser RS, Kumar S, McTernan PG. Human epicardial adipose tissue expresses a pathogenic profile of adipocytokines in patients with cardiovascular disease. Cardiovasc Diabetol. 2006;5:1. doi: 10.1186/1475-2840-5-1
- 123. Nguyen Dinh Cat A, Briones AM, Callera GE, Yogi A, He Y, Montezano AC, Touyz RM. Adipocyte-derived factors regulate vascular smooth muscle cells through mineralocorticoid and glucocorticoid receptors. *Hypertension*. 2011;58:479–488. doi: 10.1161/HYPERTENSIONAHA.110.168872
- 124. Nguyen Dinh Cat A, Touyz RM. A new look at the renin-angiotensin system–focusing on the vascular system. *Peptides*. 2011;32:2141–2150. doi: 10.1016/j.peptides.2011.09.010
- 125. Frederich RC Jr, Kahn BB, Peach MJ, Flier JS. Tissue-specific nutritional regulation of angiotensinogen in adipose tissue. *Hypertension*. 1992;19:339–344. doi: 10.1161/01.hyp.19.4.339
- 126. Massiéra F, Bloch-Faure M, Ceiler D, Murakami K, Fukamizu A, Gasc JM, Quignard-Boulange A, Negrel R, Ailhaud G, Seydoux J, et al. Adipose angiotensinogen is involved in adipose tissue growth and blood pressure regulation. FASEB J. 2001;15:2727–2729. doi: 10.1096/fj.01-0457fje
- 127. Engeli S, Gorzelniak K, Kreutz R, Runkel N, Distler A, Sharma AM. Co-expression of renin-angiotensin system genes in human adipose tissue. J Hypertens. 1999;17:555–560. doi: 10.1097/00004872-199917040-00014
- 128. Giacchetti G, Faloia E, Sardu C, Camilloni MA, Mariniello B, Gatti C, Garrapa GG, Guerrieri M, Mantero F. Gene expression of angiotensinogen in adipose tissue of obese patients. *Int J Obes Relat Metab Disord*. 2000;24(suppl 2):S142–S143. doi: 10.1038/sj.ijo.0801305
- 129. Zhang Y, Somers KR, Becari C, Polonis K, Pfeifer MA, Allen AM, Kellogg TA, Covassin N, Singh P. Comparative expression of renin-angiotensin pathway proteins in visceral versus subcutaneous fat. Front Physiol. 2018;9:1370. doi: 10.3389/fphys.2018.01370
- 130. Jones BH, Standridge MK, Taylor JW, Moustaïd N. Angiotensinogen gene expression in adipose tissue: analysis of obese models and hormonal and nutritional control. *Am J Physiol.* 1997;273:R236-R242. doi: 10.1152/ajpregu.1997.273.1.R236
- 131. Liu C, Kraja AT, Smith JA, Brody JA, Franceschini N, Bis JC, Rice K, Morrison AC, Lu Y, Weiss S, et al; CHD Exome+ Consortium; ExomeBP Consortium; GoT2DGenes Consortium; T2D-GENES Consortium; Myocardial Infarction Genetics and CARDIoGRAM Exome Consortia; CKDGen Consortium. Meta-analysis identifies common and rare variants influencing blood pressure and overlapping with metabolic trait loci. Nat Genet. 2016;48:1162–1170. doi: 10.1038/ng.3660
- 132. Morimoto H, Mori J, Nakajima H, Kawabe Y, Tsuma Y, Fukuhara S, Kodo K, Ikoma K, Matoba S, Oudit GY, et al. Angiotensin 1-7 stimulates brown adipose tissue and reduces diet-induced obesity. Am J Physiol Endocrinol Metab. 2018;314:E131–E138. doi: 10.1152/ajpendo.00192.2017
- 133. Kawabe Y, Mori J, Morimoto H, Yamaguchi M, Miyagaki S, Ota T, Tsuma Y, Fukuhara S, Nakajima H, Oudit GY, et al. ACE2 exerts anti-obesity effect via stimulating brown adipose tissue and induction of browning in white adipose tissue. Am J Physiol Endocrinol Metab. 2019;317:E1140–E1149. doi: 10.1152/ajpendo.00311.2019
- 134. Than A, Xu S, Li R, Leow MK, Sun L, Chen P. Angiotensin type 2 receptor activation promotes browning of white adipose tissue and brown adipogenesis. Signal Transduct Target Ther. 2017;2:17022. doi: 10.1038/sigtrans.2017.22
- 135. de Kloet AD, Krause EG, Scott KA, Foster MT, Herman JP, Sakai RR, Seeley RJ, Woods SC. Central angiotensin II has catabolic action at white and brown adipose tissue. Am J Physiol Endocrinol Metab. 2011;301:E1081– E1091. doi: 10.1152/ajpendo.00307.2011
- 136. Tsukuda K, Mogi M, Iwanami J, Kanno H, Nakaoka H, Wang XL, Bai HY, Shan BS, Kukida M, Higaki A, et al. Enhancement of adipocyte browning by angiotensin II type 1 receptor blockade. *PLoS One.* 2016;11:e0167704. doi: 10.1371/journal.pone.0167704
- 137. Hotamisligil GS, Shargill NS, Spiegelman BM. Adipose expression of tumor necrosis factor-alpha: direct role in obesity-linked insulin resistance. Science. 1993;259:87–91. doi: 10.1126/science.7678183
- Cildir G, Akıncılar SC, Tergaonkar V. Chronic adipose tissue inflammation: all immune cells on the stage. *Trends Mol Med.* 2013;19:487–500. doi: 10.1016/j.molmed.2013.05.001
- 139. Spalding KL, Arner E, Westermark PO, Bernard S, Buchholz BA, Bergmann O, Blomqvist L, Hoffstedt J, Näslund E, Britton T, et al. Dynamics of fat cell turnover in humans. *Nature*. 2008;453:783–787. doi: 10.1038/nature06902
- 140. Goossens GH. The metabolic phenotype in obesity: fat mass, body fat distribution, and adipose tissue function. *Obes Facts*. 2017;10:207–215. doi: 10.1159/000471488

- 141. Moreno-Indias I, Tinahones FJ. Impaired adipose tissue expandability and lipogenic capacities as ones of the main causes of metabolic disorders. J Diabetes Res. 2015;2015:970375. doi: 10.1155/2015/970375
- 142. Guzik TJ, Skiba DS, Touyz RM, Harrison DG. The role of infiltrating immune cells in dysfunctional adipose tissue. *Cardiovasc Res.* 2017;113:1009– 1023. doi: 10.1093/cvr/cvx108
- Fried SK, Lee MJ, Karastergiou K. Shaping fat distribution: new insights into the molecular determinants of depot- and sex-dependent adipose biology. Obesity (Silver Spring). 2015;23:1345–1352. doi: 10.1002/oby.21133
- 144. Palmer BF, Clegg DJ. The sexual dimorphism of obesity. *Mol Cell Endocrinol*. 2015;402:113–119. doi: 10.1016/j.mce.2014.11.029
- 145. Lee MJ, Fried SK. Sex-dependent depot differences in adipose tissue development and function; role of sex steroids. J Obes Metab Syndr. 2017;26:172–180. doi: 10.7570/jomes.2017.26.3.172
- 146. Haarbo J, Marslew U, Gotfredsen A, Christiansen C. Postmenopausal hormone replacement therapy prevents central distribution of body fat after menopause. *Metabolism*. 1991;40:1323–1326. doi: 10.1016/0026-0495(91)90037-w
- 147. Chen X, McClusky R, Chen J, Beaven SW, Tontonoz P, Arnold AP, Reue K. The number of x chromosomes causes sex differences in adiposity in mice. *PLoS Genet*. 2012;8:e1002709. doi: 10.1371/journal.pgen.1002709
- 148. Link JC, Wiese CB, Chen X, Avetisyan R, Ronquillo E, Ma F, Guo X, Yao J, Allison M, Chen YI, et al. X chromosome dosage of histone demeth-ylase KDM5C determines sex differences in adiposity. J Clin Invest. 2020;130:5688-5702. doi: 10.1172/JCI140223
- Liu R, Nikolajczyk BS. Tissue immune cells fuel obesity-associated inflammation in adipose tissue and beyond. Front Immunol. 2019;10:1587. doi: 10.3389/fimmu.2019.01587
- 150. Kanda H, Tateya S, Tamori Y, Kotani K, Hiasa K, Kitazawa R, Kitazawa S, Miyachi H, Maeda S, Egashira K, et al. MCP-1 contributes to macrophage infiltration into adipose tissue, insulin resistance, and hepatic steatosis in obesity. J Clin Invest. 2006;116:1494–1505. doi: 10.1172/JCl26498
- 151. Patsouris D, Li PP, Thapar D, Chapman J, Olefsky JM, Neels JG. Ablation of CD11c-positive cells normalizes insulin sensitivity in obese insulin resistant animals. *Cell Metab.* 2008;8:301–309. doi: 10.1016/j.cmet.2008.08.015
- 152. Wentworth JM, Naselli G, Brown WA, Doyle L, Phipson B, Smyth GK, Wabitsch M, O'Brien PE, Harrison LC. Pro-inflammatory CD11c+CD206+ adipose tissue macrophages are associated with insulin resistance in human obesity. *Diabetes*. 2010;59:1648–1656. doi: 10.2337/db09-0287
- 153. Amano SU, Cohen JL, Vangala P, Tencerova M, Nicoloro SM, Yawe JC, Shen Y, Czech MP, Aouadi M. Local proliferation of macrophages contributes to obesity-associated adipose tissue inflammation. *Cell Metab.* 2014;19:162–171. doi: 10.1016/j.cmet.2013.11.017
- 154. Cancello R, Tordjman J, Poitou C, Guilhem G, Bouillot JL, Hugol D, Coussieu C, Basdevant A, Bar Hen A, Bedossa P, et al. Increased infiltration of macrophages in omental adipose tissue is associated with marked hepatic lesions in morbid human obesity. *Diabetes*. 2006;55:1554–1561. doi: 10.2337/db06-0133
- 155. Murano I, Barbatelli G, Parisani V, Latini C, Muzzonigro G, Castellucci M, Cinti S. Dead adipocytes, detected as crown-like structures, are prevalent in visceral fat depots of genetically obese mice. *J Lipid Res.* 2008;49:1562–1568. doi: 10.1194/jlr.M800019-JLR200
- 156. Winer S, Chan Y, Paltser G, Truong D, Tsui H, Bahrami J, Dorfman R, Wang Y, Zielenski J, Mastronardi F, et al. Normalization of obesity-associated insulin resistance through immunotherapy: CD4+ T cells control glucose homeostasis. Nat Med. 2009;15:921–929.
- 157. Cipolletta D, Feuerer M, Li A, Kamei N, Lee J, Shoelson SE, Benoist C, Mathis D. PPAR-γ is a major driver of the accumulation and phenotype of adipose tissue Treg cells. *Nature*. 2012;486:549–553. doi: 10.1038/nature11132
- 158. Fuster JJ, Ouchi N, Gokce N, Walsh K. Obesity-induced changes in adipose tissue microenvironment and their impact on cardiovascular disease. Circ Res. 2016;118:1786–1807. doi: 10.1161/CIRCRESAHA.115.306885
- Osborn O, Olefsky JM. The cellular and signaling networks linking the immune system and metabolism in disease. *Nat Med.* 2012;18:363–374. doi: 10.1038/nm.2627
- 160. Saito M, Okamatsu-Ogura Y, Matsushita M, Watanabe K, Yoneshiro T, Nio-Kobayashi J, Iwanaga T, Miyagawa M, Kameya T, Nakada K, et al. High incidence of metabolically active brown adipose tissue in healthy adult humans: effects of cold exposure and adiposity. *Diabetes*. 2009;58:1526–1531. doi: 10.2337/db09-0530
- 161. Yoneshiro T, Aita S, Matsushita M, Kayahara T, Kameya T, Kawai Y, Iwanaga T, Saito M. Recruited brown adipose tissue as an antiobesity agent in humans. J Clin Invest. 2013;123:3404–3408. doi: 10.1172/JCI67803

- 162. Lowell BB, S-Susulic V, Hamann A, Lawitts JA, Himms-Hagen J, Boyer BB, Kozak LP, Flier JS. Development of obesity in transgenic mice after genetic ablation of brown adipose tissue. *Nature*. 1993;366:740–742. doi: 10.1038/366740a0
- 163. Feldmann HM, Golozoubova V, Cannon B, Nedergaard J. UCP1 ablation induces obesity and abolishes diet-induced thermogenesis in mice exempt from thermal stress by living at thermoneutrality. *Cell Metab.* 2009;9:203– 209. doi: 10.1016/j.cmet.2008.12.014
- 164. Sakamoto T, Nitta T, Maruno K, Yeh YS, Kuwata H, Tomita K, Goto T, Takahashi N, Kawada T. Macrophage infiltration into obese adipose tissues suppresses the induction of UCP1 level in mice. Am J Physiol Endocrinol Metab. 2016;310:E676–E687. doi: 10.1152/ajpendo.00028.2015
- 165. Peterson KR, Flaherty DK, Hasty AH. Obesity alters B cell and macrophage populations in brown adipose tissue. *Obesity (Silver Spring)*. 2017;25:1881–1884. doi: 10.1002/oby.21982
- 166. Bi P, Shan T, Liu W, Yue F, Yang X, Liang XR, Wang J, Li J, Carlesso N, Liu X, et al. Inhibition of Notch signaling promotes browning of white adipose tissue and ameliorates obesity. *Nat Med.* 2014;20:911–918. doi: 10.1038/nm.3615
- 167. Lawes CM, Vander Hoorn S, Rodgers A; International Society of Hypertension. Global burden of blood-pressure-related disease, 2001. Lancet. 2008;371:1513–1518. doi: 10.1016/S0140-6736(08)60655-8
- 168. Zhou D, Xi B, Zhao M, Wang L, Veeranki SP. Uncontrolled hypertension increases risk of all-cause and cardiovascular disease mortality in US adults: the NHANES III Linked Mortality Study. Sci Rep. 2018;8:9418.
- 169. Burke GL, Bertoni AG, Shea S, Tracy R, Watson KE, Blumenthal RS, Chung H, Carnethon MR. The impact of obesity on cardiovascular disease risk factors and subclinical vascular disease: the multi-ethnic study of atherosclerosis. *Arch Intern Med.* 2008;168:928–935. doi: 10.1001/archinte.168.9.928
- 170. Stevens VJ, Obarzanek E, Cook NR, Lee IM, Appel LJ, Smith West D, Milas NC, Mattfeldt-Beman M, Belden L, Bragg C, et al; Trials for the Hypertension Prevention Research Group. Long-term weight loss and changes in blood pressure: results of the Trials of Hypertension Prevention, phase II. Ann Intern Med. 2001;134:1–11. doi: 10.7326/0003-4819-134-1-200101020-00007
- 171. Lee DC, Sui X, Church TS, Lavie CJ, Jackson AS, Blair SN. Changes in fitness and fatness on the development of cardiovascular disease risk factors hypertension, metabolic syndrome, and hypercholesterolemia. *J Am Coll Cardiol.* 2012;59:665–672. doi: 10.1016/j.jacc.2011.11.013
- 172. Matsuzawa Y, Funahashi T, Nakamura T. The concept of metabolic syndrome: contribution of visceral fat accumulation and its molecular mechanism. J Atheroscler Thromb. 2011;18:629–639. doi: 10.5551/jat.7922
- 173. Huang PL. A comprehensive definition for metabolic syndrome. Dis Model Mech. 2009;2:231–237. doi: 10.1242/dmm.001180
- 174. van Harmelen V, Elizalde M, Ariapart P, Bergstedt-Lindqvist S, Reynisdottir S, Hoffstedt J, Lundkvist I, Bringman S, Arner P. The association of human adipose angiotensinogen gene expression with abdominal fat distribution in obesity. Int J Obes Relat Metab Disord. 2000;24:673–678. doi: 10.1038/sj.ijo.0801217
- 175. Rahmouni K, Mark AL, Haynes WG, Sigmund CD. Adipose depot-specific modulation of angiotensinogen gene expression in diet-induced obesity. Am J Physiol Endocrinol Metab. 2004;286:E891–E895. doi: 10.1152/ajpendo.00551.2003
- 176. Yiannikouris F, Gupte M, Putnam K, Thatcher S, Charnigo R, Rateri DL, Daugherty A, Cassis LA. Adipocyte deficiency of angiotensinogen prevents obesity-induced hypertension in male mice. *Hypertension*. 2012;60:1524–1530.
- 177. Boustany CM, Brown DR, Randall DC, Cassis LA. AT1-receptor antagonism reverses the blood pressure elevation associated with diet-induced obesity. Am J Physiol Regul Integr Comp Physiol. 2005;289:R181–R186. doi: 10.1152/ajpregu.00507.2004
- 178. Cohen P, Levy JD, Zhang Y, Frontini A, Kolodin DP, Svensson KJ, Lo JC, Zeng X, Ye L, Khandekar MJ, et al. Ablation of PRDM16 and beige adipose causes metabolic dysfunction and a subcutaneous to visceral fat switch. Cell. 2014;156:304–316. doi: 10.1016/j.cell.2013.12.021
- 179. Harms MJ, Lim HW, Ho Y, Shapira SN, Ishibashi J, Rajakumari S, Steger DJ, Lazar MA, Won KJ, Seale P. PRDM16 binds MED1 and controls chromatin architecture to determine a brown fat transcriptional program. *Genes Dev.* 2015;29:298–307. doi: 10.1101/gad.252734.114
- 180. Cittadini A, Mantzoros CS, Hampton TG, Travers KE, Katz SE, Morgan JP, Flier JS, Douglas PS. Cardiovascular abnormalities in transgenic mice with reduced brown fat: an animal model of human obesity. *Circulation*. 1999;100:2177–2183.

- 181. Dall'Asta C, Vedani P, Manunta P, Pizzocri P, Marchi M, Paganelli M, Folli F, Pontiroli AE. Effect of weight loss through laparoscopic gastric banding on blood pressure, plasma renin activity and aldosterone levels in morbid obesity. Nutr Metab Cardiovasc Dis. 2009;19:110-114. doi: 10.1016/i.numecd.2008.06.001
- 182. Maffei M, Halaas J, Ravussin E, Pratley RE, Lee GH, Zhang Y, Fei H, Kim S, Lallone R, Ranganathan S. Leptin levels in human and rodent: measurement of plasma leptin and ob RNA in obese and weight-reduced subjects. Nat Med. 1995;1:1155–1161. doi: 10.1038/nm1195-1155
- 183. Considine RV, Sinha MK, Heiman ML, Kriauciunas A, Stephens TW, Nyce MR, Ohannesian JP, Marco CC, McKee LJ, Bauer TL. Serum immunoreactive-leptin concentrations in normal-weight and obese humans. N Engl J Med. 1996;334:292–295. doi: 10.1056/NEJM199602013340503
- 184. Russell CD, Petersen RN, Rao SP, Ricci MR, Prasad A, Zhang Y, Brolin RE, Fried SK. Leptin expression in adipose tissue from obese humans: depot-specific regulation by insulin and dexamethasone. Am J Physiol. 1998;275:E507–E515. doi: 10.1152/ajpendo.1998.275.3.E507
- 185. Van Harmelen V, Reynisdottir S, Eriksson P, Thörne A, Hoffstedt J, Lönnqvist F, Arner P. Leptin secretion from subcutaneous and visceral adipose tissue in women. *Diabetes*. 1998;47:913–917. doi: 10.2337/ diabetes.47.6.913
- 186. Schoof E, Stuppy A, Harig F, Carbon R, Horbach T, Stöhr W, Rascher W, Dötsch J. Comparison of leptin gene expression in different adipose tissues in children and adults. Eur J Endocrinol. 2004;150:579–584. doi: 10.1530/eje.0.1500579
- 187. Simonds SE, Pryor JT, Ravussin E, Greenway FL, Dileone R, Allen AM, Bassi J, Elmquist JK, Keogh JM, Henning E, et al. Leptin mediates the increase in blood pressure associated with obesity. *Cell.* 2014;159:1404–1416. doi: 10.1016/j.cell.2014.10.058
- Rahmouni K, Morgan DA, Morgan GM, Mark AL, Haynes WG. Role of selective leptin resistance in diet-induced obesity hypertension. *Diabetes*. 2005;54:2012–2018. doi: 10.2337/diabetes.54.7.2012
- 189. Mark AL, Correia M, Morgan DA, Shaffer RA, Haynes WG. State-of-the-art-lecture: obesity-induced hypertension: new concepts from the emerging biology of obesity. *Hypertension*. 1999;33:537–541. doi: 10.1161/01.hyp.33.1.537
- 190. Haynes WG. Interaction between leptin and sympathetic nervous system in hypertension. Curr Hypertens Rep. 2000;2:311–318. doi: 10.1007/s11906-000-0015-1
- 191. Mark AL, Shaffer RA, Correia ML, Morgan DA, Sigmund CD, Haynes WG. Contrasting blood pressure effects of obesity in leptin-deficient ob/ob mice and agouti yellow obese mice. *J Hypertens*. 1999;17:1949–1953. doi: 10.1097/00004872-199917121-00026
- 192. Halaas JL, Boozer C, Blair-West J, Fidahusein N, Denton DA, Friedman JM. Physiological response to long-term peripheral and central leptin infusion in lean and obese mice. *Proc Natl Acad Sci USA*. 1997;94:8878–8883. doi: 10.1073/pnas.94.16.8878
- Lembo G, Vecchione C, Fratta L, Marino G, Trimarco V, d'Amati G, Trimarco B. Leptin induces direct vasodilation through distinct endothelial mechanisms. *Diabetes*. 2000;49:293–297. doi: 10.2337/diabetes.49.2.293
- 194. Nakagawa K, Higashi Y, Sasaki S, Oshima T, Matsuura H, Chayama K. Leptin causes vasodilation in humans. Hypertens Res. 2002;25:161–165.
- 195. Bravo PE, Morse S, Borne DM, Aguilar EA, Reisin E. Leptin and hypertension in obesity. Vasc Health Risk Manag. 2006;2:163–169. doi: 10.2147/vhrm.2006.2.2.163
- 196. Mark AL. Selective leptin resistance revisited. Am J Physiol Regul Integr Comp Physiol. 2013;305:R566–R581. doi: 10.1152/ajpregu.00180.2013
- 197. Correia ML, Haynes WG, Rahmouni K, Morgan DA, Sivitz WI, Mark AL. The concept of selective leptin resistance: evidence from agouti yellow obese mice. *Diabetes*. 2002;51:439–442. doi: 10.2337/diabetes.51.2.439
- 198. Gabriely I, Ma XH, Yang XM, Atzmon G, Rajala MW, Berg AH, Scherer P, Rossetti L, Barzilai N. Removal of visceral fat prevents insulin resistance and glucose intolerance of aging: an adipokine-mediated process? *Diabetes*. 2002;51:2951–2958. doi: 10.2337/diabetes.51.10.2951
- 199. Langheim S, Dreas L, Veschini L, Maisano F, Foglieni C, Ferrarello S, Sinagra G, Zingone B, Alfieri O, Ferrero E, et al. Increased expression and secretion of resistin in epicardial adipose tissue of patients with acute coronary syndrome. Am J Physiol Heart Circ Physiol. 2010;298:H746–H753. doi: 10.1152/ajpheart.00617.2009
- 200. Park SY, Kim KH, Seo KW, Bae JU, Kim YH, Lee SJ, Lee WS, Kim CD. Resistin derived from diabetic perivascular adipose tissue up-regulates vascular expression of osteopontin via the AP-1 signalling pathway. J Pathol. 2014;232:87–97. doi: 10.1002/path.4286

- Steppan CM, Bailey ST, Bhat S, Brown EJ, Banerjee RR, Wright CM, Patel HR, Ahima RS, Lazar MA. The hormone resistin links obesity to diabetes. Nature. 2001;409:307–312. doi: 10.1038/35053000
- 202. Takata Y, Osawa H, Kurata M, Kurokawa M, Yamauchi J, Ochi M, Nishida W, Okura T, Higaki J, Makino H. Hyperresistinemia is associated with coexistence of hypertension and type 2 diabetes. *Hypertension*. 2008;51:534–539.
- 203. Jiang Y, Lu L, Hu Y, Li Q, An C, Yu X, Shu L, Chen A, Niu C, Zhou L, et al. Resistin induces hypertension and insulin resistance in mice via a TLR4-dependent pathway. Sci Rep. 2016;6:22193. doi: 10.1038/srep22193
- 204. Calabro P, Samudio I, Willerson JT, Yeh ET. Resistin promotes smooth muscle cell proliferation through activation of extracellular signal-regulated kinase ½ and phosphatidylinositol 3-kinase pathways. *Circulation*. 2004;110:3335–3340. doi: 10.1161/01.CIR.0000147825.97879.E7
- Segawa K, Fukuhara A, Hosogai N, Morita K, Okuno Y, Tanaka M, Nakagawa Y, Kihara S, Funahashi T, Komuro R, et al. Visfatin in adipocytes is upregulated by hypoxia through HIF1alpha-dependent mechanism. *Biochem Biophys Res Commun.* 2006;349:875–882. doi: 10.1016/j.bbrc.2006.07.083
- 206. Gunes F, Akbal E, Cakir E, Akyurek O, Altunbas M, Ozbek M. Visfatin may be a novel marker for identifying stages of essential hypertension in advanced age patients. *Intern Med.* 2012;51:553–557. doi: 10.2169/ internalmedicine.51.6609
- 207. Dogru T, Sonmez A, Tasci I, Yilmaz MI, Erdem G, Erturk H, Bingol N, Kilic S, Ozgurtas T. Plasma visfatin levels in young male patients with uncomplicated and newly diagnosed hypertension. *J Hum Hypertens*. 2007;21:173–175. doi: 10.1038/sj.jhh.1002114
- 208. Curat CA, Wegner V, Sengenès C, Miranville A, Tonus C, Busse R, Bouloumié A. Macrophages in human visceral adipose tissue: increased accumulation in obesity and a source of resistin and visfatin. *Diabetologia*. 2006;49:744–747. doi: 10.1007/s00125-006-0173-z
- Reneau J, Goldblatt M, Gould J, Kindel T, Kastenmeier A, Higgins R, Rengel LR, Schoyer K, James R, Obi B, et al. Effect of adiposity on tissue-specific adiponectin secretion. *PLoS One*. 2018;13:e0198889. doi: 10.1371/journal.pone.0198889
- 210. Ohashi K, Kihara S, Ouchi N, Kumada M, Fujita K, Hiuge A, Hibuse T, Ryo M, Nishizawa H, Maeda N, et al. Adiponectin replenishment ameliorates obesity-related hypertension. *Hypertension*. 2006;47:1108–1116. doi: 10.1161/01.HYP.0000222368.43759.a1
- 211. Yilmaz MI, Sonmez A, Caglar K, Celik T, Yenicesu M, Eyileten T, Acikel C, Oguz Y, Yavuz I, Vural A. Effect of antihypertensive agents on plasma adiponectin levels in hypertensive patients with metabolic syndrome. *Nephrology (Carlton)*. 2007;12:147–153. doi: 10.1111/j.1440-1797.2007.00764.x
- 212. Yin C, Chu H, Li H, Xiao Y. Plasma Sfrp5 and adiponectin levels in relation to blood pressure among obese children. *J Hum Hypertens*. 2017;31:284–291. doi: 10.1038/jhh.2016.76
- 213. Imatoh T, Miyazaki M, Momose Y, Tanihara S, Une H. Adiponectin levels associated with the development of hypertension: a prospective study. *Hypertens Res.* 2008;31:229–233. doi: 10.1291/hypres.31.229
- 214. Shatat IF, Freeman KD, Vuguin PM, Dimartino-Nardi JR, Flynn JT. Relationship between adiponectin and ambulatory blood pressure in obese adolescents. *Pediatr Res.* 2009;65:691–695. doi: 10.1203/ PDR.0b013e31819ea776
- 215. Peri-Okonny PA, Ayers C, Maalouf N, Das SR, de Lemos JA, Berry JD, Turer AT, Neeland IJ, Scherer PE, Vongpatanasin W. Adiponectin protects against incident hypertension independent of body fat distribution: observations from the Dallas Heart Study. *Diabetes Metab Res Rev.* 2017;33.
- 216. Ouchi N, Ohishi M, Kihara S, Funahashi T, Nakamura T, Nagaretani H, Kumada M, Ohashi K, Okamoto Y, Nishizawa H, et al. Association of hypoadiponectinemia with impaired vasoreactivity. *Hypertension*. 2003;42:231–234.
- 217. Almabrouk TAM, Ugusman AB, Katwan OJ, Salt IP, Kennedy S. Deletion of AMPKα1 attenuates the anticontractile effect of perivascular adipose tissue (PVAT) and reduces adiponectin release. *Br J Pharmacol*. 2017;174:3398–3410. doi: 10.1111/bph.13633
- 218. Yang RZ, Lee MJ, Hu H, Pray J, Wu HB, Hansen BC, Shuldiner AR, Fried SK, McLenithan JC, Gong DW. Identification of omentin as a novel depot-specific adipokine in human adipose tissue: possible role in modulating insulin action. *Am J Physiol Endocrinol Metab.* 2006;290:E1253–E1261. doi: 10.1152/ajpendo.00572.2004
- 219. de Souza Batista CM, Yang RZ, Lee MJ, Glynn NM, Yu DZ, Pray J, Ndubuizu K, Patil S, Schwartz A, Kligman M, et al. Omentin plasma levels and gene expression are decreased in obesity. *Diabetes*. 2007;56:1655–1661. doi: 10.2337/db06-1506

- 220. Moreno-Navarrete JM, Catalán V, Ortega F, Gómez-Ambrosi J, Ricart W, Frühbeck G, Fernández-Real JM. Circulating omentin concentration increases after weight loss. Nutr Metab (Lond). 2010;7:27. doi: 10.1186/1743-7075-7-27
- 221. Kazama K, Okada M, Hara Y, Yamawaki H. A novel adipocytokine, omentin, inhibits agonists-induced increases of blood pressure in rats. J Vet Med Sci. 2013;75:1029-1034. doi: 10.1292/jvms.12-0537
- 222. Brunetti L, Leone S, Orlando G, Ferrante C, Recinella L, Chiavaroli A, Di Nisio C, Shohreh R, Manippa F, Ricciuti A, et al. Hypotensive effects of omentin-1 related to increased adiponectin and decreased interleukin-6 in intra-thoracic pericardial adipose tissue. Pharmacol Rep. 2014;66:991-995. doi: 10.1016/j.pharep.2014.06.014
- 223. Ohashi K, Shibata R, Murohara T, Ouchi N. Role of anti-inflammatory adipokines in obesity-related diseases. Trends Endocrinol Metab. 2014;25:348-355. doi: 10.1016/i.tem.2014.03.009
- 224. Zabetian-Targhi F, Mirzaei K, Keshavarz SA, Hossein-Nezhad A. Modulatory role of Omentin-1 in inflammation: cytokines and dietary intake. J Am Coll Nutr. 2016;35:670-678. doi: 10.1080/07315724.2015.1126207
- 225. Sargolzaei J, Chamani E, Kazemi T, Fallah S, Soori H. The role of adiponectin and adipolin as anti-inflammatory adipokines in the formation of macrophage foam cells and their association with cardiovascular diseases. Clin Biochem. 2018;54:1-10. doi: 10.1016/j.clinbiochem.2018.02.008
- 226. Enomoto T, Ohashi K, Shibata R, Higuchi A, Maruyama S, Izumiya I, Walsh K, Murohara T, Ouchi N. Adipolin/C1qdc2/CTRP12 protein functions as an adipokine that improves glucose metabolism\*. J Biol Chem. 2011:286:34552-34558
- 227. Ogawa H, Ohashi K, Ito M, Shibata R, Kanemura N, Yuasa D, Kambara T, Matsuo K, Havakawa S, Hiramatsu-Ito M, et al. Adipolin/CTRP12 protects against pathological vascular remodelling through suppression of smooth muscle cell growth and macrophage inflammatory response. Cardiovasc Res. 2020;116:237-249. doi: 10.1093/cvr/cvz074
- 228. Saxton SN, Whitley AS, Potter RJ, Withers SB, Grencis R, Heagerty AM. Interleukin-33 rescues perivascular adipose tissue anticontractile function in obesity. Am J Physiol Heart Circ Physiol. 2020;319:H1387-H1397. doi: 10.1152/ajpheart.00491.2020
- 229. Titone D, Aroonsakool N, Li J, et al. Increased serum interleukin-33 in patients with pulmonary arterial hypertension: a role for IL-33/ST2 in disease pathogenesis. In: B95. NOVEL THERAPEUTIC TARGETS IN PULMONARY HYPERTENSION: INSIGHTS FROM TRANSLATIONAL AND PRECLINICAL STUDIES. American Thoracic Society International Conference Abstracts. American Thoracic Society; 2014:A3636-A3636.
- 230. Gutierrez AK. The Role of Interleukin-33 in the Progression of Pulmonary Arterial Hypertension Through an ST2/MyD88 Pathway. 2018. Accessed January 11, 2021. https://escholarship.org/uc/item/52d491rq
- 231. Selcuk A, Bulucu F, Kalafat F, Cakar M, Demirbas S, Karaman M, Ay SA, Saglam K, Balta S, Demirkol S, et al. Skinfold thickness as a predictor of arterial stiffness: obesity and fatness linked to higher stiffness measurements in hypertensive patients. Clin Exp Hypertens. 2013;35:459-464. doi: 10.3109/10641963.2012.746357
- 232. Strasser B, Arvandi M, Pasha EP, Haley AP, Stanforth P, Tanaka H. Abdominal obesity is associated with arterial stiffness in middle-aged adults. Nutr Metab Cardiovasc Dis. 2015;25:495-502. doi: 10.1016/j. numecd.2015.01.002
- 233. Sutton-Tyrrell K, Newman A, Simonsick EM, Havlik R, Pahor M, Lakatta E, Spurgeon H, Vaitkevicius P. Aortic stiffness is associated with visceral adiposity in older adults enrolled in the study of health, aging, and body composition. Hypertension. 2001;38:429-433. doi: 10.1161/01.hyp.38.3.429
- 234. Tounian P, Aggoun Y, Dubern B, Varille V, Guy-Grand B, Sidi D, Girardet JP, Bonnet D. Presence of increased stiffness of the common carotid artery and endothelial dysfunction in severely obese children: a prospective study. Lancet. 2001;358:1400-1404. doi: 10.1016/S0140-6736(01)06525-4
- 235. Cote AT, Phillips AA, Harris KC, Sandor GG, Panagiotopoulos C, Devlin AM. Obesity and arterial stiffness in children: systematic review and

- meta-analysis. Arterioscler Thromb Vasc Biol. 2015;35:1038-1044. doi: 10.1161/ATVBAHA.114.305062
- 236. Mitchell GF, Hwang SJ, Vasan RS, Larson MG, Pencina MJ, Hamburg NM, Vita JA, Levy D, Benjamin EJ. Arterial stiffness and cardiovascular events: the Framingham Heart Study. Circulation. 2010;121:505-511. doi: 10.1161/CIRCULATIONAHA.109.886655
- 237. Kim HL, Lee JM, Seo JB, Chung WY, Kim SH, Zo JH, Kim MA. The effects of metabolic syndrome and its components on arterial stiffness in relation to gender. J Cardiol. 2015;65:243-249. doi: 10.1016/j.jjcc.2014.05.009
- 238. Cooper LL, Palmisano JN, Benjamin EJ, Larson MG, Vasan RS, Mitchell GF, Hamburg NM, Microvascular function contributes to the relation between aortic stiffness and cardiovascular events: the Framingham Heart Study. Circ Cardiovasc Imaging. 2016;9:e004979.
- 239. Bostick B, Habibi J, DeMarco VG, Jia G, Domeier TL, Lambert MD, Aroor AR, Nistala R, Bender SB, Garro M, et al. Mineralocorticoid receptor blockade prevents Western diet-induced diastolic dysfunction in female mice. Am J Physiol Heart Circ Physiol. 2015;308:H1126-H1135.
- 240. Jia G, Habibi J, Aroor AR, Martinez-Lemus LA, DeMarco VG, Ramirez-Perez FI, Sun Z, Hayden MR, Meininger GA, Mueller KB, et al. Endothelial mineralocorticoid receptor mediates diet-induced aortic stiffness in females. Circ Res. 2016;118:935-943. doi: 10.1161/CIRCRESAHA.115.308269
- 241. Sowers JR, Habibi J, Aroor AR, Yang Y, Lastra G, Hill MA, Whaley-Connell A, Jaisser F, Jia G. Epithelial sodium channels in endothelial cells mediate diet-induced endothelium stiffness and impaired vascular relaxation in obese female mice. Metabolism. 2019;99:57-66. doi: 10.1016/j. metabol.2019.153946
- 242. Sehgel NL, Vatner SF, Meininger GA. "Smooth muscle cell stiffness syndrome"-revisiting the structural basis of arterial stiffness. Front Physiol. 2015;6:335. doi: 10.3389/fphys.2015.00335
- 243. Sowers JR, Habibi J, Jia G, Bostick B, Manrique-Acevedo C, Lastra G, Yang Y, Chen D, Sun Z, Domeier TL, et al. Endothelial sodium channel activation promotes cardiac stiffness and diastolic dysfunction in Western diet fed female mice. Metabolism. 2020;109:154223. doi: 10.1016/j.
- 244. Hill MA, Jaisser F, Sowers JR. Role of the vascular endothelial sodium channel activation in the genesis of pathologically increased cardiovascular stiffness, Cardiovascular Research. 2020;cvaa326. doi: 10.1093/cvr/ cvaa326
- 245. Leopold Jane A. Cellular and molecular mechanisms of arterial stiffness associated with obesity. Hypertension. 2013;62:1003-1004.
- 246. Wang J, Barbry P, Maiyar AC, Rozansky DJ, Bhargava A, Leong M, Firestone GL, Pearce D. SGK integrates insulin and mineralocorticoid regulation of epithelial sodium transport. Am J Physiol Renal Physiol. 2001;280:F303-F313. doi: 10.1152/ajprenal.2001.280.2.F303
- 247. von Wowern F, Berglund G, Carlson J, Månsson H, Hedblad B, Melander O. Genetic variance of SGK-1 is associated with blood pressure, blood pressure change over time and strength of the insulin-diastolic blood pressure relationship. Kidney Int. 2005;68:2164-2172. doi: 10.1111/j.1523-1755.2005.00672.x
- 248. Huang DY, Boini KM, Osswald H, Friedrich B, Artunc F, Ullrich S, Rajamanickam J, Palmada M, Wulff P, Kuhl D, et al. Resistance of mice lacking the serum- and glucocorticoid-inducible kinase SGK1 against salt-sensitive hypertension induced by a high-fat diet. Am J Physiol Renal Physiol. 2006;291:F1264-F1273. doi: 10.1152/ajprenal.00299.2005
- 249. Li P, Pan F, Hao Y, Feng W, Song H, Zhu D. SGK1 is regulated by metabolic-related factors in 3T3-L1 adipocytes and overexpressed in the adipose tissue of subjects with obesity and diabetes. Diabetes Res Clin Pract. 2013;102:35-42. doi: 10.1016/j.diabres.2013.08.009
- 250. Norlander AE, Saleh MA, Pandey AK, Itani HA, Wu J, Xiao L, Kang J, Dale BL, Goleva SB, Laroumanie F, et al. A salt-sensing kinase in T lymphocytes, SGK1, drives hypertension and hypertensive end-organ damage. JCI Insight. 2018;2:e92801.