The past 30 years have seen a renaissance of the interest of investigators and clinicians for the sympathetic nervous system, a development supported by multiple lines of evidence. First, sympathetic neural factors are involved in the genesis of life-threatening cardiac arrhythmias and sudden death, particularly in patients with advanced heart failure, severe obesity, or sleep apnea syndrome, ie, clinical conditions which display as a common pathophysiological hallmark a marked activation of the sympathetic cardiovascular drive. Second, abnormalities in sympathetic modulation of the cardiovascular function have been reported in metabolic disease, such as diabetes mellitus, obesity, and metabolic syndrome.

**Key Words:** hypertension essential ■ renal insufficiency ■ sympathetic nervous system

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promoting cardiovascular complications and favoring directly or indirectly the disease progression.3,5,6 Third, in several diseases, such as congestive heart failure, acute myocardial infarction, acute ischemic stroke, malignant cardiac arrhythmias, renal failure, and chronic obstructive pulmonary disease, the sympathetic overdrive has been reported to be an independent predictor of mortality.7–13 Finally, recent developments have suggested, although not univocally,14–17 that in patients with true resistant hypertension radiofrequency ablation of bilateral sympathetic afferent and efferent renal nerves might trigger a blood pressure reduction, thereby emphasizing the importance of the adrenergic overdrive as a target of therapeutic blood pressure–lowering interventions.18–21

The present article will provide an in-depth review of the knowledge available on the behavior of the sympathetic nervous system in uncomplicated and complicated human hypertension. After an introductory part critically addressing the pros and cons of the methodological approaches used in clinical research to assess human sympathetic neural function, the article will examine the evidence available in favor or against the hypothesis that sympathetic activation may represent a hallmark of the essential hypertensive state. This will be followed by a discussion of sympathetic abnormalities in the escalating and challenging problems of hypertension associated with obesity or renal failure. This will allow to discuss the metabolic-sympathetic as well as the renal-sympathetic crosstalks which may concur at the marked potentiation of the sympathetic overdrive detectable in these diseases. Finally, we review the background and the results of recent clinical studies aimed at assessing the effects of carotid baroreceptor stimulation and renal denervation on cardiovascular sympathetic drive and blood pressure. The future perspectives of the research in the field of the adrenergic nervous system in human hypertension will be finally highlighted.

Methods to Assess Human Sympathetic Cardiovascular Function

An understanding of methods for evaluating the sympathetic nervous system is facilitated by appreciation of 2 major features. The first is the profound regional differences in the characteristics, control, and function of the sympathetic nervous system. There are examples of qualitative, not simply quantitative, differences in sympathetic activity to various regional beds in both physiological and pathological states.22,23 Second, there are multiple levels of the sympathetic nervous system. These include (1) central neural regulation of sympathetic nerve activity to different regions, (2) ganglionic transmission, (3) prejunctional modulation of the release of norepinephrine (and other postulated neurotransmitters) at the postganglionic neuroeffector junction, (4) clearance and reuptake of the neurotransmitter, (5) adrenergic receptors, and (6) the responsiveness of the effector.22,23 As a result of these complexities, there is no such thing as overall or generalized sympathetic nerve activity or tone, and one cannot obtain an overall assessment of the sympathetic nervous system with any single method. Each method must be judged by its strengths and limitations and by the insight being sought.

Heart Rate

Throughout the years, several techniques have been developed systemic and regional sympathetic neuroadrenergic activity.22 One approach has been assessment of heart rate, based on the evidence that heart rate is regulated in part by the positive chronotropic effects of norepinephrine and epinephrine in addition to the negative chronotropic effects of vagal parasympathetic inhibitory influence on the sinus node.1,22 There are, however, several limitations to heart rate as a specific and sensitive marker of sympathetic adrenergic drive. Heart rate is not a specific marker of cardiac adrenergic activity because it is also regulated substantially by vagal cholinergic influences so that changes in heart rate cannot be taken as a measure of adrenergic drive without the benefit of the response to β-adrenergic blockade to assess the sympathetic contribution. In addition, there are both quantitative and qualitative differences between sympathetic activity to the heart as compared with the kidney or skeletal muscle. For example, in several conditions characterized by an increase in norepinephrine spillover rate or muscle sympathetic nerve traffic, heart rate may be within normal range.22 In a study of baseline heart rate, muscle sympathetic nerve traffic, and venous plasma norepinephrine in patients with essential hypertension, obesity, or heart failure, the relationship between heart rate and muscle sympathetic nerve traffic was of moderate degree (heart failure) or absent (essential hypertension and obesity) (Figure 1).23 This suggests that heart rate may not reflect sympathetic overdrive to other important regional circulations in heart failure, hypertension, obesity, and other conditions. These and other studies24,25 indicate that despite the ease of measurement, heart rate cannot be taken as either a marker of cardiac sympathetic adrenergic activity or a reliable indicator of sympathetic activity to other regional circulations.

Power Spectral Analysis of Heart Rate

Given the limitations of heart rate per se as a marker of cardiac sympathetic activity, power spectral analysis of heart rate has gained popularity for assessment of cardiac sympathetic in addition to parasympathetic cholinergic influences on the sinus node.22 The approach has had appeal because it is noninvasive and relatively easy and inexpensive to perform. Despite this appeal, the method has significant limitations, particularly as a quantitative and specific indicator of cardiac sympathetic activity as opposed to parasympathetic activity, and its insight does not extend beyond sympathetic control of heart rate.26 The same reservations extend to power spectral analysis of arterial pressure as an indicator of generalized sympathetic activity.26

Plasma Norepinephrine

A common and traditional approach for assessing adrenergic drive is measurement of plasma norepinephrine. The advantages of this approach are its relatively easy performance and wide applicability even in large scale studies. However, the sensitivity and reproducibility of the measurements are far from optimal.22 The limited reproducibility can be circumvented by repeating the sampling several times and averaging the values, in which case, the results of the assay approach values of reproducibility displayed by microneurographic recording of efferent postganglionic muscle sympathetic nerve traffic.22
The sensitivity of the method is more challenging as a result of several factors. First, circulating norepinephrine levels are only a minor fraction of the amount secreted from sympathetic nerve terminals. As a result, plasma norepinephrine values may not reflect sympathetic neural drive and secretion of the neurotransmitter. For example, Meredith et al demonstrated that in patients with autonomic failure, elevated plasma norepinephrine responses to head-up tilt do not reflect an increase in the release of norepinephrine from the adrenergic nerve terminal, but rather a reduced tissue clearance process of norepinephrine dependent on a reduction in tissue perfusion. Finally, there is a conceptual limitation to measurement of plasma norepinephrine as an indicator of sympathetic activity. Specifically, it treats the sympathetic nervous system as a humoral system that can be assayed with a single circulating marker. As mentioned earlier, the sympathetic nervous system has multiple levels and components. There can be profound regional differences in sympathetic nerve activity in physiological and pathological states. Thus, there is no such thing as overall or generalized sympathetic activity. Consequently, measurement of plasma norepinephrine (or any other single measurement) cannot be taken as a sensitive or specific marker of overall sympathetic activity and may miss alterations in sympathetic activity in various regions or levels.

This brings us to a discussion of the strengths and limitations of the 2 methods that have greatly advanced understanding of the sympathetic nervous system in humans in the past 35 years: (1) norepinephrine isotope dilution method for measurement of regional and total norepinephrine spillover and (2) direct intraneural microneurographic recordings of postganglionic efferent sympathetic nerve activity to skeletal muscle or cutaneous circulations. Although each of these methods has its limitations, the 2 methods have proven highly complementary and have been primarily responsible for enormous advances in understanding sympathetic nervous system alterations in human hypertension, often in ways not currently realized in experimental animals using available techniques.

Regional and Total Norepinephrine Spillover

The norepinephrine isotope dilution method involving infusion of small doses of radiolabeled norepinephrine represents an approach that overcomes several limitations related to plasma norepinephrine measurement. This method permits precise quantification of the net release of the adrenergic neurotransmitter from sympathetic nerves and the amount of norepinephrine undergoing clearance from the bloodstream. Given the regional differences in sympathetic nerve activity in physiological and pathological states, there is no such thing as overall or generalized sympathetic activity. Consequently, measurement of plasma norepinephrine (or any other single measurement) cannot be taken as a sensitive or specific marker of overall sympathetic activity and may miss alterations in sympathetic activity in various regions or levels.

Figure 1. Relationships between heart rate (HR) and muscle sympathetic nerve traffic (MSNA), as assessed by the microneurographic technique in the peroneal nerve (MSNA, expressed as bursts frequency over time), in healthy controls, essential hypertensive patients, obese subjects, and patients with heart failure. In each panel number of subjects (n), correlation coefficients (r), and P values are shown. Note that only in heart failure patients a significant relationship was found. Figure redrawn from data derived from Ref. 23.
requirements limit widespread use of this method. Fortunately, the investigators at the Baker Institute in Melbourne, Australia, who developed the method, have mastered its use. As a result of their work, this method has yielded valuable insights into the sympathetic nervous system in humans, including human hypertension.

**Microneurography**
The microneurographic method for direct intraneural recording of efferent postganglionic sympathetic nerve activity to skeletal muscle and skin has contributed greatly to study of sympathetic cardiovascular drive in humans. There are profound differences in the characteristics and control of sympathetic nerve activity to skeletal muscle and skin, and the focus in most microneurographic studies of cardiovascular control has been on sympathetic activity to skeletal muscle. The technique is minimally invasive, requiring the percutaneous insertion and positioning of a 200-μm tungsten recording electrode in superficial nerves (usually peroneal or radial nerve), allowing to directly record efferent postganglionic sympathetic neural bursts.22 The advantages of the approach include (1) direct measurement of central nervous system sympathetic neural outflow to either the skeletal muscle circulation or to skin; (2) a continuous moment-to-moment recording of sympathetic nerve activity, allowing a dynamic assessment of sympathetic drive in a given experimental session; (3) the ease of performing repeated recordings over time; and (4) the high reproducibility of the activity between recordings in 2 different nerves and over time.22 The limitations are 2-fold. First, postganglionic sympathetic nerve traffic may not always reflect the release of the neurotransmitter at the neuroeffector junction. Second, microneurographic recordings do not provide direct information on sympathetic activity to visceral tissues such as the kidney and heart. In healthy humans, there are good relationships between the microneurographic data collected in the peroneal nerve and the ones characterizing the cardiac or the renal neural network, but this may not pertain in pathological states.22

**Sympathetic Adrenergic Neuroimaging**
One further approach of investigating sympathetic activity in humans deserves to be briefly mentioned, namely the so-called neuroimaging technique, which uses radiolabeled sympathetic amines (specifically 123-metaiodobenzylguanidine) to provide imaging of innervation of a given organ, most frequently the heart.22 The cost of the method and its inability to provide dynamic, rather than static, assessment of adrenergic cardiovascular drive have limited its widespread use in clinical research.

In summary, measurements of regional norepinephrine spillover to kidney and heart and direct microneurographic recordings of central neural sympathetic traffic to the skeletal muscle circulation represent the gold standards for evaluating the sympathetic nervous system in humans today. They have complementary strengths and limitations and together have greatly advanced understanding of sympathetic cardiovascular regulation in human hypertension and beyond. The current review draws mainly on the results of studies using these 2 state-of-the-art methods.

**Evidence in Favor and Against Adrenergic Overdrive as Hallmark of Human Hypertension**
Studies examining indirect and direct markers of sympathetic function have provided compelling evidence in the early stages of hypertension or in young hypertensives, and the sympathetic nervous system is activated. In classic studies by Julius et al.,23 young subjects with hyperkinetic borderline hypertension had increased heart rate which was frequently associated with an elevation in plasma norepinephrine. Subsequent studies using microneurography or measurements of norepinephrine spillover in young adults with borderline or established hypertension have also demonstrated marked sympathetic activation.28,32,33 Although there are several exceptions,34,35 the weight of evidence suggests that in the early stages of hypertension, particularly in young patients, there is marked adrenergic overdrive, a finding that supports the hypothesis that sympathetic overactivity occurs in the clinical phases of essential hypertension. It is more challenging to know (1) if the adrenergic activation precedes and promotes the elevation of blood pressure and (2) if the sympathetic overdrive is also present in stable hypertensive state of middle age or elderly patients. As far as the first issue is concerned, old data from the Framingham study seem to support the possibility that adrenergic overdrive precedes the elevation of blood pressure, by showing that young adults who display a resting tachycardia are more prone to develop an elevation in blood pressure over years than are age-matched controls with normal heart rate.36 This finding is supported by the data collected in a longitudinal study in which arterial epinephrine (but not norepinephrine) predicted the development of hypertension during the 20-year follow-up.37 As far as the second issue is concerned, microneurographic studies almost universally shown that sympathetic nerve traffic is potentiated in established hypertension in middle age and in elderly as well as young patients.38–46 This finding suggests that neuroadrenergic factors may contribute to the maintenance and progression as well as the development of the hypertensive state. Interestingly, studies of elderly hypertensives have not always documented an increase in renal norepinephrine spillover in contrast to the finding of increased sympathetic nerve traffic to the skeletal muscle circulation.22,28,39,47 This highlights the regional differences in sympathetic nerve activity in some pathological states and emphasizes the value of direct measurements of sympathetic activity to specific regional circulations.

Adrenergic neural factors may also contribute to the development and progression of target organ damage, as documented by the pronounced increase in sympathetic nerve traffic and cardiac norepinephrine spillover rate in patients in which hypertension is complicated by left ventricular hypertrophy or left ventricular diastolic dysfunction.40–44,45 A similar potentiation has been described when renal damage complicates the hypertensive state (see below).46–51 Finally, the arteriolar remodeling that characterizes hypertension and makes the elevation of total peripheral resistance partly dependent on structural factors, ie, an increased wall-to-lumen ratio, has been found to be in large part mediated by sympathetic neural
influences throughout direct effects on the different components of the arterial vascular wall.52,53

In discussing the evidence in favor and against adrenergic overdrive as hallmark of the hypertensive state, it is worthy to mentioning that sympathetic activation may not be confined to the essential or idiopathic hypertensive state. Some studies have reported that patients with hypertension secondary to renal artery stenosis or primary aldosteronism also display an increase in sympathetic nerve traffic to skeletal muscle circulation.54–56 In contrast, other microneurographic studies failed to show an increase in sympathetic nerve traffic in these forms of secondary hypertension.38,54

In summary, measurements of sympathetic nerve traffic to skeletal muscle and of norepinephrine spillover from kidney and heart in the past several decades provide direct evidence for sympathetic activation to skeletal muscle, kidney, and heart in the early as well as established stages of human hypertension. This evidence supports the concept that the sympathetic nervous system contributes to the development and maintenance of hypertension and to end organ damage. The sympathetic overactivity is not, however, uniform or universal. It does not, for example, extend to skin sympathetic activity.57 Because of limited methodology, there is much less information about sympathetic activity to the splanchnic circulation in humans. Given the recent interest in the role of the splanchnic circulation in animal models of hypertension, this highlights the need for advances in methodology for direct measurement of sympathetic activity to splanchnic circulation in humans.

**Sympathetic Nervous System in Obesity-Related Hypertension and the Metabolic Syndrome: Insights From Studies in Humans**

During the past 20 years, there has been burgeoning interest in the role of the sympathetic nervous system in obesity, hypertension, and the metabolic syndrome. Drawing frequently on the power of microneurographic recordings58 and measurements of regional norepinephrine spillover,59 studies of humans have provided unique insights not feasible in studies of experimental animals. These studies have demonstrated that while obesity and essential hypertension are both characterized by sympathetic overactivity, there are features of the altered sympathetic nervous system in human obesity and obesity-associated hypertension that are distinct from those in essential hypertension. Most notably, elevated sympathetic activity in obesity often occurs in the absence of hypertension.3,59,60

**Sympathetic Nervous System in Obese Normotensive and Hypertensive Humans**

There have been 2 major competing concepts regarding the role of the sympathetic nervous system in obesity. The first, introduced by Landsberg,83 postulated that obesity produces insulin resistance and hyperinsulinemia which stimulates the sympathetic nervous system and promotes thermogenic metabolism. The second, advanced by Bray,84 proposed that most obese states are characterized by underactivity of the sympathetic nervous system that decreases thermogenesis and predisposes to obesity. Sophisticated studies using microneurography and radiolabeled norepinephrine spillover have unequivocally supported Landsberg’s concept of sympathetic overactivity in most human obesity.1,59,60,63 In an exception, however, there is evidence that lean Pima Indians have lower muscle sympathetic nerve traffic than lean Caucasians.64 According to Bray’s hypothesis,82 this could contribute to the known predisposition of Pima Indians to obesity. It has also been speculated that the lower sympathetic activity may explain why Pima Indians have a low prevalence of obesity-associated hypertension.85

A remarkable feature of sympathetic overactivity in human obesity is that it often occurs in the absence of hypertension.1,59,60 Furthermore, the neurophysiologic alterations in the sympathetic nervous system in obese normotensive and hypertensive humans differ from those in lean hypertensives (Figure 2). In obese hypertensive humans, there is enhanced muscle sympathetic nerve activity, but the neurophysiologic basis of this increase is different in obese versus lean hypertensives.66 In lean patients with hypertension, the increased sympathetic activity is related to increased firing frequency of single nerve fibers. In contrast, in obese normotensive and hypertensive subjects, the increased activity is because of recruitment of additional fibers and not to increasing firing frequency of single nerve fibers.66 The precise mechanistic and hemodynamic significance of increased single fiber versus multifiber sympathetic activity is not clear, but the contrast between the neurophysiologic pattern of increased muscle sympathetic nerve traffic in obese versus lean hypertensives suggests that the mechanisms of adrenergic activation differ in lean and obese human hypertensives.

There are 2 other notable features of sympathetic activity in obese normotensive and hypertensive humans. First, cardiac norepinephrine spillover is decreased in obese normotensives and is only marginally increased in obese hypertensives.59,60 This contrasts with marked elevation in cardiac norepinephrine spillover in lean hypertensives.59 Second, in obese humans, renal norepinephrine spillover is not significantly higher in hypertensive than it is in normotensive subjects.59,60 In contrast, in lean humans renal norepinephrine spillover is significantly higher in hypertensive versus normotensive subjects.27,28 Given the view that sympathetic activity to the kidney is critical to the pathogenesis of hypertension, the finding that renal norepinephrine spillover is not different in obese normotensives and hypertensives suggests that increased renal...
sympathetic activity is not a sufficient explanation for the development of obesity-associated hypertension in humans. The contribution of renal sympathetic activity to the development of hypertension seems to be greater in lean versus obese individuals.

Two studies using pharmacological antagonists to evaluate the contribution of the sympathetic nervous system to obesity-associated hypertension merit discussion. Wofford et al compared the antihypertensive effect of combined α- and β-adrenergic receptor blockade for 1 month in obese versus lean hypertensive patients with comparable baseline blood pressure. Based on the previous discussion, one might predict that adrenergic blockade would produce a greater fall in blood pressure in lean patients with hypertension. Surprisingly, this was not the result. Adrenergic blockade produced a significantly greater decrease in blood pressure in obese versus lean patients with hypertension. The effect of adrenergic blockade on blood pressure in obese normotensive subjects was not studied.

Shibao et al evaluated the sympathetic contribution to blood pressure in obesity using a different strategy. These investigators used acute ganglionic blockade to reversibly disrupt sympathetic activity in lean and obese normotensive subjects and in obese hypertensive subjects. There were 3 key findings. First, ganglionic blockade produced an exaggerated fall in vascular resistance and blood pressure in obese compared with lean normotensive subjects. This indicates that even in the absence of overt obesity-associated hypertension, obesity is associated with an exaggerated sympathetic contribution to blood pressure in humans. Among other implications, this may help explain why weight loss tends to decrease blood pressure in obese individuals even without a history of hypertension. Second, the fall in blood pressure with ganglionic blockade in the obese hypertensives was double that in the obese normotensives, consistent with an important role for the sympathetic nervous system in maintaining blood pressure in obese hypertensives. Third, after ganglionic blockade, blood pressure was normalized in the obese normotensives. Blood pressure also decreased in the obese hypertensives, but it did not completely indicating that factors other than the sympathetic nervous system contribute to obesity-associated hypertension. This study supports an exaggerated sympathetic contribution to blood pressure in both normotensive and hypertensive obese humans.

So there are 2 seemingly conflicting observations. First, renal norepinephrine spillover, although elevated, is not higher in hypertensive versus normotensive obese humans. Second, pharmacological studies indicate that the sympathetic contribution to blood pressure is greater in hypertensive versus normotensive obese humans. What is the explanation(s) for these seemingly conflicting observations. The most likely is a prohypertensive contribution of increased muscle sympathetic nerve activity which is higher in hypertensive versus normotensive obese humans. This explanation is consistent with the finding that ganglionic blockade produced an exaggerated fall in vascular resistance in obese hypertensives. An additional explanation may be that heightened sympathetic activity is a necessary but not sufficient cause for the development of hypertension in obesity. The residual elevation in blood pressure in obese hypertensives after ganglionic blockade in the study by Shibao et al indicates, not surprisingly, that factors other than the sympathetic nervous system contribute to the hypertension in obesity. The interaction with these factors may enhance the prohypertensive influence of the sympathetic nervous system on blood pressure and the development of obesity-associated hypertension.

**Sympathetic Nervous System in the Metabolic Syndrome**

Obese humans with the metabolic syndrome have elevated muscle sympathetic nerve traffic values even in the absence of hypertension. During diet-induced weight loss, the decreases in sympathetic activity are accompanied by reduction in all components of the metabolic syndrome. Muscle sympathetic nerve activity correlates directly with waist circumference in subjects with the metabolic syndrome. This is consistent with evidence that muscle sympathetic nerve traffic correlates with visceral but not subcutaneous fat in obesity. Visceral adiposity is a key driver of the metabolic syndrome.

Huggett et al demonstrated that patients with metabolic syndrome without hypertension had increased muscle sympathetic nerve activity (specifically increased single fiber firing) compared with a control group matched for body weight and lacking the metabolic syndrome. The presence of hypertension in addition to other components of the metabolic syndrome was associated with further increases in single fiber firing and multifiber muscle sympathetic nerve activity. This suggests that the metabolic syndrome is accompanied by greater sympathetic drive than is obesity in the absence of metabolic syndrome. We speculate that this may reflect the influence of visceral as opposed to subcutaneous fat.

**Mechanisms of Sympathetic Overactivity and Hypertension in Human Obesity**

Since the discovery of leptin in 1994, there has been an explosion of research on the genetic–neurobiological mechanisms of obesity. This in turn has fueled mounting interest and insights, particularly in experimental animals, on the mechanisms of obesity-associated hypertension. Many of these mechanisms such as the role of the renin–angiotensin system, the kidney, reflex mechanisms, obstructive sleep apnea, and brain oxidative and endoplasmic stress relate to mechanisms in experimental models of essential hypertension and are not discussed here, but several are distinct to obesity-associated hypertension. In 1999, it was suggested that the emerging biology of obesity prompted 2 new concepts regarding obesity-associated hypertension. First, the effect of obesity on blood pressure may depend critically on the genetic–neurobiological mechanisms underlying the obesity. Second, obesity is not consistently associated with increased blood pressure. We focus here on evidence from humans relating to 3 possible mechanisms that are of particular relevance to the role of sympathetic nervous system in to obesity-associated hypertension and to the role of the genetic–neurobiological mechanisms of obesity in the regulation of the sympathetic nervous system and blood pressure.

**Interaction of Insulin, the Sympathetic Nervous System, and Insulin Sensitivity**

Landsberg proposed that the sympathoexcitatory response to obesity is triggered by hyperinsulinemia and that the
sympathetic activation in turn increases vascular resistance. The concept that insulin-induced sympathetic activation produces vasoconstriction is a linchpin of the hypothesis that hyperinsulinemia is a major contributor to obesity-associated hypertension. Two decades ago, studies from several laboratories demonstrated that hyperinsulinemic, euglycemic clamp produced striking increases in muscle sympathetic nerve traffic and plasma norepinephrine in healthy lean subjects.\textsuperscript{25,78,79} The surprising finding was that despite insulin-induced sympathetic activation, insulin failed to increase blood pressure because the sympathetic vasoconstriction was opposed and overridden by a vasodilator action of insulin.\textsuperscript{25} These findings were consistent with studies in dogs demonstrating that physiological increases in plasma insulin failed to increase blood pressure because increases in cardiac output were offset by substantial decreases in vascular resistance.\textsuperscript{90} Thus, in normal humans, hyperinsulinemia increases sympathetic nerve activity but does not increase blood pressure because the sympathetic activation is offset by a direct vasodilator action of insulin\textsuperscript{25} (Figure 3, left). There is, however, evidence that the balance between the vasodilator and sympathetic actions of insulin is altered in obesity and hypertension in favor of the sympathetic pressor effect. Laakso et al\textsuperscript{82} reported that insulin-induced vasodilation in skeletal muscle is reduced in obese, insulin resistant humans. In contrast, Lembo et al\textsuperscript{83} demonstrated that insulin-induced increases in sympathetic activity and norepinephrine release in skeletal muscle are substantially exaggerated in patients with hypertension (Figure 3, right). These observations raise the possibility, not yet established experimentally, that in obese, hypertension-prone individuals, chronic hyperinsulinemia might contribute to increases in blood pressure as well as sympathetic nerve activity.

An iconoclastic and major development in the insulin-sympathetic story was the demonstration that reflex sympathetic activation promotes insulin resistance in skeletal muscle. In studies of insulin-induced vasodilation in humans, Laakso et al\textsuperscript{82} introduced the concept that insulin-induced increases in skeletal muscle blood flow play an important role in glucose delivery and uptake, ie, impaired insulin-induced vasodilation in skeletal muscle contributes importantly to reduced glucose uptake in skeletal muscle and to insulin resistance. Shortly thereafter, 2 key studies demonstrated that reflex sympathetic vasoconstriction in the forearm of normal subjects decreased glucose uptake during hyperinsulinemia.\textsuperscript{84,85} Recently, Gamboa et al\textsuperscript{86} expanded on this discovery by demonstrating that ganglionic blockade improved insulin-glucose metabolism in obese insulin resistant subjects with elevated baseline muscle sympathetic nerve activity but not in obese insulin sensitive subjects. These studies advanced the concept that the sympathetic nervous system promotes insulin resistance in skeletal muscle through hemodynamic mechanisms and that the interaction of insulin, the sympathetic nervous system and insulin sensitivity are 2-way street.\textsuperscript{87} Insulin increases sympathetic activity. In turn, increased sympathetic vasoconstriction decreases glucose uptake in skeletal muscle and thereby promotes insulin resistance and compensatory hyperinsulinemia.

Brain Melanocortin Pathway, Sympathetic Activity, and Obesity-Associated Hypertension

The brain melanocortin pathway has emerged from studies in both experimental animals\textsuperscript{73,74} and humans\textsuperscript{88,89} as a crucial pathway in the regulation of sympathetic activity and blood pressure in obesity. In addition to inhibiting appetite and increasing metabolism, stimulation of hypothalamic melanocortin-4 receptors increases sympathetic activity and blood pressure.\textsuperscript{73,74} Rodents with loss-of-function mutations in the hypothalamic melanocortin-4 receptors are obese but lack sympathetic activation and obesity-associated hypertension.\textsuperscript{73,74} There are rare patients with obesity caused by loss-of-function mutations in hypothalamic melanocortin-4 receptors. These individuals have lower sympathetic activity and blood pressure than control, weight-matched patients with common human obesity.\textsuperscript{88,89} In addition, administration of an hypothalamic melanocortin-4 receptors agonist in healthy subjects increases blood pressure.\textsuperscript{88} These data support the concept, discussed above, that the genetic–neurobiological basis of obesity can critically influence the sympathetic and blood pressure response to obesity\textsuperscript{77} and demonstrate that hypothalamic melanocortin-4 receptors contribute to regulation of sympathetic activity and blood pressure in obese humans.

Leptin, the Sympathetic Nervous System and Obesity-Associated Hypertension

In addition to its effects on appetite and metabolism, leptin acts in the brain to produce receptor-mediated increases in regional sympathetic activity.\textsuperscript{90} A large body of evidence has demonstrated that leptin contributes to sympathetic overactivity and hypertension in several animal models of monogenic and diet-induced obesity.\textsuperscript{73,74} Surprisingly then, evidence that leptin contributes to obesity-associated hypertension in humans is inconclusive.\textsuperscript{76} Ozata et al\textsuperscript{91} reported that 3 adults and 1 child with severe obesity and complete leptin deficiency had sympathetic hypofunction manifest by orthostatic hypotension and an attenuated cold pressor test. Baseline blood pressures were not presented, but the patients reportedly did not have hypertension.\textsuperscript{92} These observations suggest that loss of physiological leptin action in humans is associated with decreases in sympathetic activity and blood pressure. Machleidt

![Figure 3](image-url)
et al. demonstrated that a bolus injection of leptin in healthy lean men produced an acute increase in muscle sympathetic nerve activity, but blood pressure did not increase. The vexing question is whether hyperleptinemia contributes to obesity-associated hypertension in humans. The lack of a safe, effective, reversible leptin antagonist for studies in humans has greatly impeded evaluation of the contribution of leptin to obesity-associated hypertension in humans. There are, however, data from several studies in relatively large numbers of lean and obese subjects that have failed to show an increase in blood pressure with chronic or acute administration of leptin. This is in contrast with increases in blood pressure with administration of a hypothalamic melanocortin-4 receptors agonist in humans. We conclude that studies in humans have not yet convincingly demonstrated that hyperleptinemia contributes significantly to obesity-associated human hypertension.

Summary: the Sympathetic Nervous System in Obesity, Hypertension, and the Metabolic Syndrome

Obesity-associated hypertension in humans is characterized by sympathetic overactivity, but there are features of the sympathetic nervous system in obesity that are distinct from those in lean hypertensives. Cardiac norepinephrine spillover is lower in obese normotensive and hypertensive patients than in their lean counterparts. Elevated renal norepinephrine spillover and muscle sympathetic nerve traffic are observed in normotensive as well as hypertensive obese subjects. Muscle sympathetic nerve activity is higher in hypertensive versus normotensive obese subjects, but renal norepinephrine spillover is not different in hypertensive versus normotensive obese subjects. This contrasts with essential hypertension where renal norepinephrine spillover is significantly higher in lean hypertensive versus normotensive humans. These observations suggest that renal sympathetic activity is more important in the development of hypertension in lean versus obese individuals. In juxtaposition to the neurophysiologic studies, experiments with pharmacological antagonists indicate that the sympathetic contribution to vascular resistance and blood pressure is exaggerated in obesity-associated hypertension in humans. These observations suggest an important contribution of elevated muscle sympathetic nerve traffic to obesity-associated hypertension. Muscle sympathetic nerve activity is higher in hypertensive versus normotensive obese subjects. In addition, other factors mediating obesity-associated hypertension may interact to enhance the prohypertensive contribution of elevated sympathetic activity.

In the past 20 years, extensive research in experimental animals has implicated multiple and mounting mechanisms in the pathogenesis of sympathetic activation and hypertension in obesity. In contrast, experimental studies in humans have not yet conclusively identified the major mediators of obesity-associated sympathetic activation and hypertension in humans. This difference between the abundant and compelling data on the mechanisms of obesity-associated sympathetic activation and hypertension in experimental animals and the paucity of conclusive evidence in humans highlights the need for mechanistic, state-of-the-art patient-oriented research on this topic.

Renal Sympathetic Nervous System and Kidney-Related Hypertension

The nerves of the kidney occupy a special place in the panoply of hypertension. The renal sympathetic nerves, through their multiple influences on tubular processing of sodium, on renin secretion and on renal vascular resistance, with activation of the renal sympathetic outflow in essential hypertension as the driver, are pivotal in hypertension pathogenesis. Beyond that, nociceptive sensory nerves of the kidneys which detect renal injury, through projection to the central nervous system, contribute to the systemic sympathetic nervous system activation of severe essential hypertension, renal hypertension and end-stage renal disease.

Renal Efferent Sympathetic Nerves

Renal sympathetic activation is thought to be central to the pathogenesis of essential hypertension. In untreated essential patients with hypertension, the application of regional norepinephrine isotope dilution methodology demonstrates that a high level of activation of the renal sympathetic outflow is present (Figure 4). The sympathetic neural outflow is commonly activated also to the heart, shown with selective cardiac norepinephrine spillover measurements, and to the skeletal muscle vasculature, demonstrated with microneurographic nerve traffic recording, but it is the renal sympathetic activation which is central to hypertension pathogenesis.

The renal tubules receive a dense sympathetic innervation, at all tubular levels. A specific and important relation of the renal sympathetic nerves to renal tubular sodium reabsorption, key to hypertension pathogenesis, concerns pressure natriuresis, the normal capacity of the kidneys to excrete sodium at higher arterial perfusion pressures. Impairment of pressure

Figure 4. Measurements of renal norepinephrine spillover to plasma, used to assess sympathetic activity in the kidneys in healthy volunteers and patients with arterial hypertension. Renal sympathetic activation was commonly evident in patients with hypertension. In untreated patients with mild-moderately severe essential hypertension (middle column), renal norepinephrine spillover was increased overall, and elevated in ≈50%. In drug-resistant hypertension, with patients administered on average 5 antihypertensive drug classes, renal norepinephrine spillover was higher again, attributable to their hypertension, and perhaps its treatment. From unpublished data by Murray Esler, Markus Schlaich, Gavin Lambert, and Dagmara Hering.
natriuresis is thought to be a central element in the development of hypertension. Renal sympathetic denervation shifts the renal pressure-natriuresis curve to the left, promoting urinary sodium excretion and lowering of blood pressure. Surgical sympathectomy in experimental hypertension abolishes the hypertension or prevents its development.

**Renal Afferent Sympathetic Nerves**

Renal sensory nerves are of 2 principal types. The first, of the renal parenchyma and bearing nociceptive receptors, respond to a renal injury signal by projecting to the hypothalamus to increase central sympathetic outflow. The second are pressure-sensitive receptors in the renal pelvis; these are sympathoinhibitory and important in mediating reno-renal reflexes. The nociceptive renal afferents have been implicated in generating increased systemic sympathetic activity in drug-resistant hypertension and chronic kidney disease.

What the renal injury signal is in these 2 contexts is not known. Renal ischemia has been invoked, but this proposition is unproven. Chemical irritation, from intrarenal phenol injection, can act as a potent stimulus in experimental studies in rats. In patients with drug-resistant hypertension, there are few if any clinical markers of injury. Renal ischemia has been invoked, but this proposition is unproven. Chemical irritation, from intrarenal phenol injection, can act as a potent stimulus in experimental studies in rats. In patients with drug-resistant hypertension, there are few if any clinical markers of injury. Glomerular filtration rate is commonly normal and proteinuria is often absent. But these clinical indices are insensitive and belie the fact that an injury signal must be operating. Ablation of the renal afferent nerves with endovascular delivery of radiofrequency energy in patients with drug-resistant hypertension causes inhibition of central sympathetic outflow.

**Chronic Kidney Disease**

Sympathetic nervous system activation is present in patients with chronic kidney disease, intimated first with the finding of elevated plasma norepinephrine concentrations, then more definitively with demonstration of whole-body norepinephrine spillover measurements and increased sympathetic nerve firing recorded with clinical microneurography. Consistent with this, pronounced blood pressure lowering is observed with pharmacological adrenergic inhibition by clonidine and debrisoquine. Sympathetic activation is present in the early phases of renal disease, in the absence of uremia, being seen in patients with nephrotic syndrome and with autosomal dominant polycystic kidney disease despite normal renal function, and increases with disease progression. The mechanism of sympathetic activation seems to be complex, no doubt involving afferent renal nerve signaling, but also perhaps other mechanisms.

**End-Stage Renal Disease**

The highest level of systemic sympathetic nervous activation is seen in patients with end-stage renal disease maintained on hemodialysis, equal to or exceeding that seen in New York Heart Association class IV cardiac failure. This has been conclusively demonstrated, in elegant clinical studies, to be generated by renal afferent nerve signaling, not systemic uremic toxins. The sympathetic activation of end-stage renal disease, and commonly also the hypertension, is not reversed by successful renal transplantation but is reversed by bilateral removal of the diseased native kidneys, the surgery removing the influence of the afferent renal nerves. These clinical observations have a parallel in experimental studies in rats with 5/6 nephrectomy, where the hypertension and sympathetic activation in this renal failure model are abolished by dorsal rhizotomy, which abolishes renal afferent nerve input to the central nervous system.

**Influence of Sympathetic Activity and Its Inhibition on Kidney Disease Progression and Clinical Outcome**

Sympathetic nervous system activation in renal hypertension and end-stage renal disease contributes to the blood pressure elevation, to kidney disease progression, and to cardiovascular complications and clinical outcomes. Sympathetic activation aggravates existing hypertension, proteinuria, interstitial fibrosis, and glomerulosclerosis. A striking observation is that of Zoccali et al., who demonstrated the plasma concentration of norepinephrine to predict the incidence of cardiovascular events, and survival, in patients with end-stage renal disease. High sympathetic activity in the heart is arrhythmogenic, providing a partial explanation for the high rate of sudden death in chronic renal failure. Some of these effects can be blocked by pharmacological sympathetic inhibition. The central sympatholytic drug, moxonidine, has been demonstrated to reduce urinary protein excretion in patients with type I diabetes mellitus, at a dose not influencing blood pressure, and in another study in patients with chronic renal failure, to minimize progression of renal disease compared with a comparator drug, nitrindipine, an effect not explained by the small falls in blood pressure.

**Catheter-Based Renal Denervation in Renal Hypertension**

Given the importance of the renal afferent nerves in generating high sympathetic nervous activity in renal hypertension and end-stage renal disease, thereby contributing to the blood pressure elevation, on theoretical grounds there might be a special place for catheter-based renal denervation in the treatment of the hypertension of renal disease. To this point, only 2 small pilot studies have been conducted, both uncontrolled but successful, in renal hypertension, and in end-stage renal disease. In any definitive trials, yet to be conducted, there are challenging technical matters which will need to be overcome. The first is to preserve renal function during the necessary angiographic imaging, avoiding radiocontrast nephropathy by, for example, the use of carbon dioxide angiographic imaging. The second technical difficulty, in end-stage renal disease, is that the denervation procedure will often need to be performed on small diameter renal arteries with low blood flow, which increases the risk of damaging the artery. Because bilateral nephrectomy is sometimes performed for uncontrollable hypertension in end-stage renal disease, exploring the possible option of renal denervation through a well-designed clinical trial is warranted.

**Sympathetic Deactivation as a Goal of Antihypertensive Treatment: the Long and Circular Path From Surgical Sympathectomy on the Pressor Nerves to Catheter-Based Renal Denervation**

Thomas Willis and the 17th century London neuroanatomical school he led provided the first accurate depictions of the sympathetic nervous system. Stimulation of the sympathetic
nerves, by Claude Bernard and Charles Brown-Sequard demonstrated them to be vasoconstrictor, and to elevate blood pressure, leading to their designation as the pressor nerves.\textsuperscript{109} By the first years of the 20th century, this information, and his own clinical observations, led Geisbock\textsuperscript{110} to propose that human hypertension was caused by the influence of the brain on the sympathetic nervous system.

**Surgical Sympathectomy**
In this era, no treatment of hypertension was available until the introduction of surgical sympathectomy.\textsuperscript{111} The aim was to surgically sever sections of the sympathetic chain and all accessible sympathetic nerves of the thorax and abdomen, cutting as many pressor nerves as possible to remove their systemic vasoconstrictor influence. Around this time, the first measurements of cardiac output demonstrated that blood pressure elevation in severe hypertension was directly attributable to increased total peripheral resistance, plausibly thought to derive from the pressor nerves, which were now targeted. Selective renal sympathectomy was not performed, as no theory existed suggesting specific importance of the sympathetic nerves of the kidneys in hypertension pathogenesis, although surgical sympathectomy no doubt often interrupted the sympathetic outflow to the kidneys. Surgical sympathectomy for the treatment of hypertension, applied in the years 1935 to 1960,\textsuperscript{111} took many forms, with the various surgeries being demonstrably of value in lowering blood pressure and prolonging life in patients with severe and malignant hypertension, but at the cost of disabling side effects, most notably postural and postprandial hypotension and syncope, and sexual dysfunction.

**Antiadrenergic Drugs**
Ganglionic blocking drugs, discovered by Paton and Zaimis,\textsuperscript{112} ended the period of surgical sympathectomy for hypertension and ushered in the era of antiadrenergic drugs. Ganglion blockers constituted the first antiadrenergic pharmacotherapy for hypertension and could achieve what surgery sympathectomy achieved minus surgical risk, but regrettably not minus complications, which as expected were almost identical with those of sympathectomy. But the new concept of antiadrenergic antihypertensive pharmacotherapy had been established. Based on identification of the sympathetic neurotransmitter as norepinephrine, documentation of central neural mechanisms controlling sympathetic outflow, and categorization of adrenergic receptors, centrally-acting sympathetic nervous inhibitors including methyldopa and clonidine, neurone-blocking drugs such as guanethidine and debrisoquine, \(\beta\)-adrenergic receptor blocking drugs including propranolol, and \(\alpha\)-adrenergic receptor blockers was developed in quick succession.\textsuperscript{111} Ganglion blockers rapidly became a footnote to history. Antiadrenergic drugs, coupled with diuretics and direct-acting vasodilators such as hydralazine, became the preferred antihypertensive therapy for 30 years.\textsuperscript{114}

**Drug-Resistant Hypertension**
From the 1990s, drugs antagonizing the renin–angiotensin system have become the dominant antihypertensive therapy. Angiotensin-converting enzyme inhibitor drugs and angiotensin receptor blocking drugs gradually replaced antiadrenergic drugs as the preferred antihypertensive agents because they were at least equally efficacious, and substantially better tolerated. Subsequently joined by dihydropyridine calcium channel blocking drugs, the \(\alpha\)-renin drugs, calcium channel blockers, and diuretics came to occupy the top rung of hypertension treatment international guidelines lists,\textsuperscript{114} with antiadrenergic antihypertensive drugs drifting toward the bottom. But there was a problem. Despite the widespread availability and prescribing of angiotensin-converting enzyme inhibitors, angiotensin receptor blockers, diuretics, and calcium channel blockers, in a substantial minority of patients with essential hypertension, perhaps 10%,\textsuperscript{115} goal blood pressure was not achieved. In these drug-resistant hypertensives, a new strategy was needed, and in fact, devised. This was the development of 2 device-based therapies targeting the sympathetic nervous system, the surgically implanted barostimulator device\textsuperscript{116} and catheter-based renal denervation.\textsuperscript{20,117} These new therapies were developed on the well-established premise that activation of the sympathetic nervous system commonly initiates and sustains the blood pressure elevation in essential hypertension.

**Arterial Baroreceptor Stimulation**
Devices stimulating the human carotid baroreflex were first developed 5 decades ago,\textsuperscript{118} and after a prolonged absence have had a resurgence. In experimental models of hypertension, activation of central baroreflex pathways by continuous electric stimulation of the nerves of the carotid sinus baroreceptors reduces sympathetic outflow from the central nervous system and lowers blood pressure. The Rheos implantable carotid sinus stimulator (CVRx, Minneapolis, MN) has been studied in patients with severe drug-resistant hypertension.\textsuperscript{116} In a large-scale, double-blind study, the Rheos device was implanted in 265 patients who were subsequently randomized (2:1) to immediate baroreceptor stimulation for the first 6 months (n=181), or deferred stimulation after 6 months (n=84).\textsuperscript{116} At 6 months, blood pressure fell in the group receiving immediate treatment, but the primary efficacy end point was not reached, partly because blood pressure likewise decreased in many control subjects.\textsuperscript{118} The future of this procedure is therefore uncertain, to be determined by an ongoing clinical trial utilizing a less cumbersome electrode than that used previously, and with unilateral rather than bilateral stimulation of the carotid sinus.

**Catheter-Based Renal Denervation**
Three facts provided the knowledge base for the development of catheter-based renal denervation for treatment of essential hypertension: (1) the presence of an activated renal sympathetic outflow in patients with hypertension, (2) the blood-pressure-lowering effect of surgical renal denevration in experimental models of hypertension, and (3) the anatomy of the postganglionic renal sympathetic nerves in their passage to the kidneys.\textsuperscript{91-95} In humans, the nerves pass from the sympathetic chain and ganglia to the kidneys via the outer adventitia of the renal arteries, or just beyond in perirenal adipose tissue and connective tissue, potentially within reach of radiofrequency energy delivered by a catheter in the artery lumen.\textsuperscript{119-123} Drawing on these concepts, the first to suggest that essential hypertension might be treated with
a renal nerve ablation catheter were Howard Levin and Mark Gelfand in US provisional patents 60/370,190 (April 2002), 60/415,575 (October 2002), and 60/442,970 (January 2003). The California start-up company, Ardian, acquired the Levin and Gelfand patent rights and commenced a developmental program to design a radiofrequency ablation catheter suitable for human use, testing this purpose-designed catheter for safety and renal denervation capacity in pigs. The first-in-man studies were conducted in Melbourne, in patients with drug-resistant hypertension. This patient class, of resistant hypertension, was selected because of the evident clinical need, and because the potential benefit-risk balance made the study defensible ethically. Only recently has it been shown that among patients with hypertension, activation of the renal sympathetic outflow is at its highest in drug-resistant hypertension (Figure 4). This trial, commenced in June 2007, is now known as SYMPLICITY HTN-1.20

Renal Denervation Antihypertensive Trials

The SYMPLICITY trials in endovascular renal denervation,20,117 their subsequent continuation to specified end points, accompanying renal denervation registry files, trials with other, newly engineered radiofrequency renal denervation devices,121 and application of denervating ultrasonic energy122 have established these therapeutic principles:

1. Efferent sympathetic renal denervation can be achieved with luminal delivery of radiofrequency and ultrasonic energy.
2. Blood pressure reduction across the trials shows consistency, office systolic blood pressure falling on average by 20 to 30 mm Hg.
3. The blood pressure reduction is durable, persisting beyond 3 years, with no evidence that renal nerve regeneration occurs, to cancel out blood pressure lowering.
4. The development of renal artery stenosis in the region of radiofrequency energy delivery is uncommon.

Testing for Achieved Renal Denervation

The belief that renal denervation has been achieved in clinical trials of catheter-based renal denervation is almost invariably based on trust rather than testing. This contrasts with the studies of surgical renal denervation in experimental hypertension, where good experimental design demands that the effectiveness of denervation is always confirmed, typically by documenting 90% to 95% reduction in the kidney content of norepinephrine.120

Among clinical studies of endovascular renal denervation only in the SYMPLICITY HTN-1 trial20 was renal denervation confirmed, by measurements of norepinephrine spillover (release of the transmitter from the renal nerves to plasma). Measurement of regional norepinephrine spillover is well established as a valid test for sympathetic denervation, having been applied for 2 decades in the diagnosis of pure autonomic failure,121 a disorder characterized by spontaneous degeneration of postganglionic sympathetic nerve fibers, and denervation of the heart, kidneys, and other organs. The degree of renal denervation achieved in the SYMPLICITY HTN-1 trial was less than expected (on average 47%),20 but did seem to be sufficient, in that the antihypertensive response was adequate. Subsequent analyses in Melbourne confirm that denervation is often incomplete and surprisingly nonuniform from patient to patient (Figure 5). Achieved denervation can be <25%, no doubt inadequate for a full therapeutic effect. Although the denervation catheter technique might look easy, compared with many other interventional cardiological and radiological procedures, achieving denervation is difficult. Illustrations showing the ready proximity of renal sympathetic nerves to the renal artery lumen were misleading. Older surgical anatomies of the renal nerves120 demonstrated that at the origin of the renal artery from the aorta the sympathetic nerves were more remote from the artery, but they converged on the renal arteries distally, near the renal artery branch point. Contemporary anatomic studies confirm this.159 Clearly, ablative energy should be preferentially focused on the distal renal artery.

Renal Denervation Defrocked? The SYMPLICITY HTN-3 Trial

A challenge to the percutaneous renal denervation treatment of resistant hypertension came with the 9 January, 2014, press release concerning the SYMPLICITY HTN-3 trial in drug-resistant hypertension, the pivotal study for US Food and Drug Administration licensure, and in the subsequent New England Journal of Medicine publication on 29 March,124 indicating that the primary efficacy end point had not been reached in the trial. A lot was expected of the SYMPLICITY HTN-3 study. Five times larger than the first 2 SYMPLICITY renal denervation
Conclusions: the Future of Research on the Sympathetic Nervous System in Human Hypertension

Because this review has focused on the sympathetic nervous system in human hypertension, we reflect in closing on the future of research on the sympathetic neural mechanisms in human hypertension. The past 30 years witnessed enormous progress in understanding the role and importance of the sympathetic nervous system in the pathogenesis of human hypertension and its consequences. These advances resulted substantially from hypothesis based, mechanistic, patient-oriented research drawing on the sophistication and power of molecular biology and genetics and clinical trials. What is needed to renew that vitality in the coming years? Ingenious new methods. A coupling of patient-oriented research to basic research and advances in human genetics and neuroscience. Most of all, a serious commitment, not just lip service introduced with the word translational, to the training and careers of physician-scientists pursuing mechanistic patient-oriented research.

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