Review

Genetic and Non-Genetic Basis of Essential Hypertension: Maladaptation of Human Civilization to High Salt Intake

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The relation between salt intake and hypertension has long been controversial. Available evidence provides conflicting results. However, biological and evolutionary insights into the structure and principle function of the kidney and the blood pressure sustaining machinery, with the extracellular fluid volume and the renin-angiotensin system acting as a central player, clearly indicate that salt has been a precious commodity in the terrestrial animal kingdom and that essential hypertension may be a consequence of the kidney's maladaptation to excess salt intake, which is specific to recent human civilization. This review provides a hypothesis to explain how and why hypertension develops and attempts to define the roles of genetic, non-genetic, and environmental factors in its pathogenesis. (Hypertens Res 1998; 21: 67-71)

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The human kidneys, consisting of approximately 2 million nephrons, receive 1 l/min of blood flow, the highest blood flow per organ weight in the body. When the blood flows through the glomeruli, a highly efficient filtration system, each kidney generates 100 ml/min of urine. Approximately 99% of this glomerular filtrate is reabsorbed along the tubules, a highly organized and regulated epithelial transporting system. Why is such a process necessary? The principal function of the kidney is to maintain the homeostasis of our milieu interieur, or the extracellular fluid (ECF), despite wide variations in and ranges of daily fluid and electrolyte intake. The range of daily fluid and electrolyte intake that causes no significant disturbances in the milieu interieur is quite large: water intake could range from 0 to 30 l/d; salt, 0 to 1,000 mEq/d; potassium, 0 to 700 mEq/d, and so on. This wide allowance in the intake of fluids and electrolytes is due to a high glomerular filtration rate (GFR) of 140 d, with approximately 99% reabsorption of the glomerular filtrate. A patient with a GFR of 10% of normal would have to have a much lower intake of fluid and electrolytes, because a higher intake would cause pathological changes in the milieu interieur. It is this remarkable kidney function of high filtration and high reabsorption that allows us great flexibility in our daily fluid and electrolyte intake. A detailed discussion of some of the issues outlined in this overview has appeared elsewhere (1, 2).

Autoregulation of Renal Blood Flow

To maintain a high GFR despite possible fluctuations in blood pressure in a terrestrial environment with 1.0 G gravity, the kidney must maintain a constant blood supply and GFR. Table 1 summarizes the principal functions of the kidney (2). Constancy of renal blood flow is achieved with a two-component system: a myogenic response of the afferent artery (AA), and tubuloglomerular feedback (TGF) by the juxtaglomerular apparatus (JGA). In response to changes in renal perfusion pressure, vascular smooth muscles of the AA respond to maintain the down-stream perfusion pressure so that the renal blood flow remains constant, i.e., myogenic response. The very last portion of the AA at the entrance to the glomerulus is regulated principally by TGF. Table 1 summarizes the principal functions of the kidney (2). Constancy of renal blood flow is achieved with a two-component system: a myogenic response of the afferent artery (AA), and tubuloglomerular feedback (TGF) by the juxtaglomerular apparatus (JGA). In response to changes in renal perfusion pressure, vascular smooth muscles of the AA respond to maintain the down-stream perfusion pressure so that the renal blood flow remains constant, i.e., myogenic response. The very last portion of the AA at the entrance to the glomerulus is regulated principally by TGF; this portion contracts or relaxes in response to an increase or decrease, respectively, in macula densa chloride (Cl\textsuperscript{−}) delivery (3). Moreover, Holstein-Rathlou and colleagues (4, 5) have reported oscillations with a frequency of approximately 20 s of the distal tubular fluid [Cl\textsuperscript{−}] just beyond the macula densa and of the proximal tubule pressure, reflecting single nephron glomerular filtration rates (SNGFR) of the same tubule. This oscillation is driven by TGF at the entrance of the AA to the glomerulus. These observations clearly indicate that the apparent constancy of the SNGFR is maintained by the TGF.

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Autoregulation and Maintenance of Single Nephron Glomerular Filtration Rates

What is the possible physiological importance of autoregulation of renal glomerular plasma flow? Consider a situation where salt intake is very low, perhaps less than 2 g/d. The systemic blood pressure and renal perfusion pressure will drop. A decrease in AA resistance ($R_A$) due to the myogenic response of AA and also due to TGF at the entrance to the glomerulus maintains glomerular plasma flow (GPF). A drop in ECF volume and systemic blood pressure maximally activates the renin-angiotensin system (RAS). Maximal activation of the RAS leads to a preferential increase in efferent arteriolar resistance ($R_E$) by angiotensin II (AII) to maintain glomerular capillary pressure, thus maintaining the SNGFR. This process allows a wide range of the daily intake of fluids and minerals, despite a low salt intake (6).

Phylogenic Considerations of Juxtaglomerular Apparatus

It is thought that the JGA and TGF appear in the nephrons of amphibians, reptiles, birds, and mammals (7, 8). We could therefore envision the functional roles of JGA or TGF from an evolutionary perspective. Simply compare the milieu interieur of the sea fish, teleost, to that of humans. The basic constituents of intracellular and extracellular fluids are comparable: thus, the ECF, or milieu interieur, is isotonic saline, and potassium is the major cation of the intracellular fluids for both teleosts and humans. When a fish ingests its meal, the salt it absorbs has a higher concentration than that of the milieu interieur, i.e., 0.9% NaCl in sea water. Thus, the teleost does not need salt, but rather needs salt-free water. The milieu interieur of the teleost is maintained primarily by the gills and the kidney has only an ancillary role.

In contrast, there is no guarantee of salt intake for terrestrial creatures. In fact, the availability of salt is, in general, very limited for terrestrial life, since the main sources of food stuff in the natural environment are plants and animal meats. Nonetheless, it is necessary to maintain a high GFR to maintain the milieu interieur in response to a wide range of fluid and electrolyte intake. This is probably why the JGA first appeared in amphibians, and it is likely that the JGA was an essential component of the nephron in order for organisms to live on lands, where a high GFR must be maintained with a minimal salt intake. It should be noted that blood pressure must rise in terrestrial life to ensure that the blood supplies oxygen throughout the body system in 1.0 G gravity (9). The blood pressure rise helps to develop a high GFR.

Renal Cross Transplantation in Rat Models of Hypertension

Results of renal cross transplantation studies in an experimental model of hypertension in the rat strongly indicate that the kidney has a primary role in the genesis of hypertension (10). Thus, available data clearly indicate that the kidney determines whether or not hypertension eventually develops. Similar results have also been obtained from studies in humans. In chronic dialysis patients with end stage renal failure due to severe hypertension, successful kidney transplantation from a normotensive, healthy donor often leads to the disappearance of hypertension (11). These observations are consistent with the notion that it is the kidney that dictates the development of hypertension.

Aberrant Tubuloglomerular Feedback as an Intrarenal Mechanism Responsible for Abnormal Salt Excretion

There are numerous reviews and monographs on the role of the kidney in the genesis of hypertension. In particular, two recent reviews address the critical role of the kidney. One role proposed by
Laragh and colleagues emphasizes the presence of a subset of nephrons that secrete inappropriately increased amounts of renin for any given salt intake (12). Another hypothesis, proposed by Brenner and colleagues, indicates that a decreased number of nephrons, and thus a less effective glomerular filtration surface, is associated with the development of hypertension (13). This hypothesis is also persuasive.

A third hypothesis that we advance is that an abnormal JGA or resetting of the TGF might be involved in the development of hypertension (2). Available data indicate similarities in the TGF response of spontaneously hypertensive rats (SHR) and Wistar Kyoto rats (WKY) and that of normal rats with and without All. Thus, with a given salt intake, TGF is more activated in SHR than in WKY. This difference is quite similar to that between normal rats with and without All (14, 15). The data suggest that the TGF of SHR is behaving as if TGF is more sensitive to All for any given salt intake or ECF volume. Thus, in response to a saline load, the RAS will be inhibited because of volume expansion, and the TGF curve will become either less activated or inhibited; at the same time, distal Cl⁻ delivery is increased by the saline load, but the inhibition of TGF increases the GFR, and saline is effectively excreted in the urine. In hypertensive subjects, however, this TGF inhibition is incomplete or aberrant; thus, in response to increased distal Cl⁻ delivery, the GFR might go unchanged or even drop, rather than increase, leading to volume expansion with a slower urinary excretion of loaded saline. This is a characteristic response in hypertensive patients or even some normotensive subjects with a family history of essential hypertension (16, 17). This series of changes in response to salt loading eventually leads to hypertension with increased peripheral resistance if excess salt intake continues. Thus, in hypertensive patients, inability of the kidneys to excrete salt must be responsible for the development and maintenance of hypertension (18). The hypothesis of Laragh and colleagues is consistent with the abnormal TGF governed by the RAS, particularly in superficial nephrons.

If this hypothesis is correct, then in persons with a family history of essential hypertension, genetic defects must be somehow involved in the growth and development of the kidney during fetal life, since the number of nephrons would be determined by the time of birth and would not increase significantly thereafter. Such genetic abnormalities have not yet been identified. Nonetheless, it is important to note that there are other conditions that might affect the nephron number in individuals born after a full-term pregnancy. Available evidence suggests a good correlation between birth weight and later development of hypertension; indeed, birth weight is inversely correlated with blood pressure in childhood, adolescence, and adulthood, and this correlation is amplified with increasing age (19-21).

According to Brenner, this correlation might be due to a lower number of nephrons, which is determined by low birth weight and due principally to intrauterine protein malnutrition. Such conditions might also cause hypertension with the high salt intake of our culture in those who were in utero during poor nutritional conditions, such as those associated with poverty, low socioeconomic conditions, and war. Hypertension might develop in siblings because of similar intrauterine nutrition provided by their mother. Nonetheless, it is clear that hypertension is not necessarily a genetically determined event. It is thus important to note that such conditions should be regarded as a non-genetic cause of hypertension, even in the presence of a “family history.”

Structural Alterations in the Kidney in Essential Hypertension

Whatever pathogenic factors underlie essential hypertension, the dominant and most characteristic renal abnormality in essential hypertension has been documented to be afferent arteriolar hyalinosis, which is directly related to both the severity and duration of hypertension (22). The glomeruli and postglomerular structures might be subject to ischemic changes. Such pathologic changes in the kidney are consistent with aberrant TGF, since TGF is associated with increased afferent arteriolar resistance. Nonetheless, as hypertension progresses in severity and duration, renal arterioles and the glomeruli begin to show direct influences of the high perfusion pressure as well as ischemic changes.

In contrast, if the lower number of nephrons is the prevailing cause of essential hypertension, one might expect to see the presence of hypertrophic glomeruli with focal sclerosis in the glomerular tufts in early hypertension, early renal lesions typically seen in remnant kidney, a model of oligonephronia. This must be carefully examined, particularly in hypertensive subjects with good documentation of uterine malnutrition and low birth weight (20, 21). In any event, a detailed examination of early renal lesions in subjects with essential hypertension might reveal a relative prevalence of distinct pathogenic factors, which might vary depending upon region, race, historical events, and other factors.
Salt Intake in Human Civilization

According to the hypothesis outlined here, hypertension will not develop where salt intake is very low. Epidemiological studies indicate that this is indeed the case. People with very low salt intake, e.g., Papua New Guineans and Yanomamo Indians in the Amazon, do not develop hypertension, and blood pressure does not rise as they get older (23).

We must realize that only humans have acquired the habit of consuming excess salt (23). Due to this habit of high salt intake, some will become hypertensive because they cannot adapt appropriately to a chronic excess salt intake because of a disorder in the TGF. In a natural environment, mammals do not constantly consume excess salt. The salt intake of carnivores is 20 to 40 mEq/d/60 kg body weight, but that of herbivores is less than 10 mEq/d/60 kg body weight. Thus, terrestrial mammals in a natural environment are in a state of chronic volume depletion with a highly activated RAS.

Analyses of Paleolithic nutrition suggest that salt intake by human beings, Homo sapiens, was 30 mmol/d at most, with a potassium intake of 500 to 700 mmol/d (24). The development of agriculture approximately 10,000 years ago did not change this salt intake when vegetable foods came to make up 90% of the diet. Thus, throughout our civilization, salt has been a valuable commodity and not an item in abundance. Abundant evidence suggests the importance of salt in our culture and civilization. Interested readers are referred to excellent monographs by Denton (23) and Astrap (25) and a recent brief review by Ritz (26).

Genetic, Non-Genetic, and Environmental Factors of Hypertension

Our body system has evolved through years of evolution accompanied by genetic mutations. The kidneys and the blood pressure maintenance system are no exception. Thus, our body system is genetically adapted to the salt depletion of the terrestrial environment. “Industrialized civilizations” have introduced an excessive salt intake, however, leading to a condition called essential hypertension or a condition of maladaptation to chronic high salt intake in a subset of people. Obviously, the last few centuries have not provided enough time for our body to genetically adapt to such excessive salt intake. In fact, some genetic abnormalities responsible for hypertension, such as a mutation in the distal Na+ channels leading to increased Na+ reabsorption in Liddle syndrome, would have been advantageous for survival in a natural environment with a low salt intake. This abnormality has been shown to be a cause of hypertension only in recent years because of our high salt intake as a culture. Indeed, mutations that have an opposite effect on the same Na+ channel, i.e., decreased distal Na+ reabsorption, would have caused severe dehydration in children who inherited the trait, a condition quite disadvantageous for survival in a natural ‘primitive’ environment with low salt availability. Thus, such mutations of the Na+ channel could not be conferred to subsequent generations. This is the reason why we do not see such mutations. These are the bases for genetic and environmental factors in the genesis of hypertension.

Moreover, as observed in the case of oligonephronia, maternal protein malnutrition can cause hypertension in offsprings in a culture with high salt intake. This mechanism is not genetically determined, but might appear in the siblings, and thus can be viewed as “hypertension with a positive family history.” In any event, the differences among the three hypotheses presented here on the intrarenal mechanisms of essential hypertension could be viewed from slightly different perspectives. Two of the hypotheses, one by Laragh proposing the presence of a subset of nephrons with unsuppressed renin secretion for a given volume, and the other by Brenner, proposing a lesser number of nephrons, both explain the mechanism of how the kidney is unable to appropriately excrete extra salt, leading to development of hypertension. Abnormal JGA function or an abnormal TGF setting as a cause of hypertension likewise explains as well as the other two hypotheses how the kidney is unable to excrete salt intake, but it also explains why such abnormalities could have been present in humans. As proposed here, the kidney must have acquired, during the evolution of life and the transition from the sea to the land, the ability to maintain a high GFR in the face of a very low salt intake. Thus, high salt intake was never expected at the genetic level during
human evolution. Stated differently, genetic abnormalities associated with essential hypertension would have been beneficial for survival in a natural environment with a low salt intake and were therefore preserved for many generations, only to become manifest in recent centuries because of the high salt intake of our culture (Table 2).

References