Serum uric acid and cardiovascular disease

Adriana ILIESIU, MD, PhD, FESC; Alexandru CAMPEANU, MD, PhD; Dinu DUSCEAC, MD, PhD

Department of Internal Medicine, Caritas Hospital, “Carol Davila” University of Medicine and Pharmacy, Bucharest, Romania

The relationship between the serum uric acid (SUA) and gout, hypertension and obesity has been known since the late 19th century. After 1960s many epidemiological studies confirm the relationship between the SUA level and various cardiovascular diseases, such as arterial hypertension, atherosclerosis, stroke, acute and chronic heart failure. This relationship is observed not only in frank hyperuricemia (defined by SUA > 7 mg in men and > 6 mg in women), but also in the high – normal values of SUA (defined by SUA > 5.5 mg/dl) (1, 2). The significance of this association is still unknown and in literature there are controversies concerning the significance of hyperuricemia in cardiovascular disease: is it only a marker or is it also a risk factor?

Uric acid is the final oxidation product of purine catabolism in humans and in higher primates. The last metabolic step, the conversion of hypoxanthine to uric acid is regulated by the enzyme xanthine oxidoreductase (XO). As a part of this process reactive oxygen species (ROS) are produced (Figure 1). The major sources of XO are the liver and the small intestine, but there are evidences for local production of XO by the endothelium and myocardium. XO is associated with enhanced oxidative stress. XO activity is up-regulated in many cardiovascular diseases, such as myocardial ischemia, reperfusion injury, left ventricular remodeling after myocardial infarction and heart failure.

The SUA level reflects the net balance between its constant production and excretion. Dietary intake of urate provides a source of uric acid precursors. To maintain homeostasis, SUA is eliminated by kidney and the gastrointestinal tract. Two thirds of the daily turnover of the urate is excreted by the kidney, where it is completely filtered at the glomerulus, completely reabsorbed in the proximal tubule, then secreted (approxiimately 50% of the filtered load), and finally reabsorbed.

Hyperuricemia is a very common metabolic disorder. Elevated SUA levels occur in 2–18% of the population, varying in relation to age, sex, and many other factors (Table 1).

The SUA level is higher in postmenopausal women (because the uricosuric effect of estrogen) and in African Americans (3). The high SUA level can be due to an excessive production or to a decreased excretion. An increased
TABLE 1. Causes of the serum acid elevation, according to Johnson (56)

<table>
<thead>
<tr>
<th>Group</th>
<th>Mechanism</th>
</tr>
</thead>
<tbody>
<tr>
<td>Postmenopausal women</td>
<td>Estrogen is uricosuric</td>
</tr>
<tr>
<td>African Americans</td>
<td>Unknown</td>
</tr>
<tr>
<td>Renal disease</td>
<td>Decrease in GFR increases SUA levels</td>
</tr>
<tr>
<td>Diuretics</td>
<td>Volume contraction promotes SUA reabsorption</td>
</tr>
<tr>
<td>Hypertension</td>
<td>Urate ressertions increased in setting and is tightly linked to SUA reabsorption; microvascular disease predisposes to tissue ischemia that leads to increased urate generation (from adenosine breakdown) and reduced excretion (due to lactate competing with urate transporter in the proximal tubule); some hyperuricemic hypertension may be due to alcohol ingestion or lead intoxication</td>
</tr>
<tr>
<td>Alcohol use</td>
<td>Increases SUA generation, decreases SUA excretion</td>
</tr>
</tbody>
</table>

FIGURE 1. Uric acid synthesis: In humans uric acid is the terminal step of purine metabolism, catalyzed by xanthinoxidase, which also produces superoxide. Xanthinoxidase is inhibited by allopurinol.

dietary purine or fructose intake increases the SUA production. In malignancies, polycytemia vera or haemolytic anemias, the rapid cellular turnover determines excessive SUA production. Renal insufficiency is a common cause of increase in SUA. Hyperuricemia is highly prevalent in chronic kidney disease, reflecting the reducing renal excretion of SUA. The data of the recent experimental and clinical studies suggest that SUA is not only a marker of reduced kidney function, but it is also a causal factor in the development and progression of renal disease (4). The use of diuretics, by causing volume contraction, increases the SUA level by increasing urate absorption.

SUA AND CARDIOVASCULAR EVENTS

The relationship between SUA and the cardiovascular risk was demonstrated in many epidemiological studies (5). In the MONICA Ausburg study the increase in the SUA level was an independent factor for all causes of death and possible for the cardiovascular death (6). In the First National Health and Nutrition Study (NHANES I) study, for every 1.01 mg/dl
increase in the SUA level, the hazard ratio for total mortality and for cardiovascular mortality were 1.09 and 1.19 for men and 1.26 and 1.3 for women, respectively (7). The result of the LIFE Study pointed out an association between the baseline SUA level and the risk of cardiovascular events in a high risk population with coronary artery disease (8). In the Multiple Risk Factors Intervention Trial (MRFIT), both hyperuricemia and gout were independent risk factors for myocardial infarction in 12866 men followed for 6.5 years (9).

In contrast, in the Atherosclerotic Risk in Communities (ARIC) Study and in the Framingham Heart Study there was no association between SUA and incident cardiovascular disease (10, 11).

The difficulties in the assessment of the role of SUA independently from other traditional risk factors and the different methodologies used in the epidemiological studies may be responsible for the conflicting data regarding the relationship between the SUA level and cardiovascular disease.

SUA AND HYPERTENSION

The association between arterial hypertension (HT) and hyperuricemia is very common. It has been reported that 25-40% of patients with untreated HT and more than 80% of patients with malignant HT have high SUA levels (12). Hyperuricemia is more common in primary HT, especially in patients with HT of recent onset and in preHT associated with microalbuminuria (13). Many mechanisms are involved in high SUA level in HT. The reabsorption of the urate in the proximal tubule is increased as a consequence to the reduced renal blood flow. The microvascular renal disease leads to tissue ischemia and to the up-regulation of XO with increased the SUA production. The reduction of the SUA secretion in the proximal tubule and the use of diuretics may increase the SUA level.

In the recent years, several experimental studies have indicated that hyperuricemia per se can induce HT. In rats the high SUA level induced HT after several weeks (14). The HT was reversed after the normalization of SUA with allopurinol or with an uricozuric drug. Two main mechanisms are involved in the hyperuricemia-induced HT (FIGURE 2). In early stage the high SUA level induces renal vasoconstriction by the activation of the renal RAAS and by the endothelial dysfunction with decreased nitric oxide level at the macula densa. In this stage, HT is salt-resistant and it is reversed by lowering the SUA level. In later stage chronic hyperuricemia induces vascular muscle cell proliferation and local activation of RAAS system with the activation of the mediators of inflammation. Progressive microvascular renal disease is associated with afferent arteriolsclerosis and with interstitial fibrosis (15, 16). The renal histopathologic changes in chronic hyperuricemia are similar to those induced by HT. HT becomes salt-driven and renal-dependent and it is not normalized by lowering SUA.

Several clinical studies demonstrated that hyperuricemia precedes and it is associated with the development of HT. In the Framingham Heart Study, each increase in SUA by 1.3 mg/dl was associated to the development of HT with an odd ratio of 1.17 (17). In the Multiple Risk Factor Intervention (MRFIT) study, in normotensive men with the SUA level greater than 7 mg/dl there was an 80% increased risk for the development of HT (18). The association between hyperuricemia and HT was more common in young people. The high SUA was observed in nearly 90% of adolescents with primary HT and the SUA level correlates with both systolic and diastolic HT (19, 20). In a study including adolescents with HT of recent onset and hyperuricemia, the reduction in SUA to less than 5 mg/dl with allopurinol was associated to the reversal of HT in 86% of the patients (21).
SUA AND METABOLIC SYNDROME, INSULIN RESISTANCE AND DIABETES

Epidemiological and clinical studies have established a close link between the high SUA level and the increasing prevalence of the metabolic syndrome and all its individual components (glucose intolerance, insulin resistance, abdominal obesity, atherogenic dyslipidemia and HT) (22, 23). In the Third National Health and Nutrition Examination Survey, the prevalence of the metabolic syndrome was determined in persons with normal body mass index at different SUA levels (24). The prevalences of the metabolic syndrome was 18.9% for the SUA levels less than 6 mg/dL; in contrast, the prevalence of metabolic syndrome increased at 70.7% for the SUA levels of 10 mg/dL or greater. Moreover, hyperuricemia might independently predict the development of different components of the metabolic syndrome – obesity, hyperinsulinemia and diabetes (25-27).

The elevated SUA level observed in the metabolic syndrome has been attributed to hyperinsulinemia, since insulin reduces renal excretion of uric acid. In animal studies, hyperuricemia might induce metabolic syndrome by two mechanisms. Firstly, high level of uric acid have been shown to inhibit endothelial NO bioavailability. Because insulin requires endothelial NO to stimulate skeletal muscle glucose uptake, hyperuricemia may have a causal role in the pathogenesis of insulin-resistance. Secondly, in animal models, hyperuricemia induces oxidative and inflammatory changes in adipocytes, inducing metabolic syndrome in obese mice (28).

SUA AND ATHEROSCLEROSIS

The pathophysiological link between the elevated SUA and atherosclerosis are endothelial dysfunction and inflammation. ROS production by XO can induce endothelial dysfunction by reducing bioavailability of nitric oxide (33). SUA, by its antioxidant properties, could counteract ROS generation. There are also evidences in animal experiments that the high SUA impairs endothelial dependent vaso-dilatation (34). An independent association between the SUA level and C-reactive protein and other inflammatory markers (blood neutrophils, interleukin, TNF-alfa) has also been described (35, 36). So far there is evidence that the increased SUA level is associated with subclinical atherosclerosis.

The relationship between SUA and the development of coronary artery disease and cerebrovascular disease was investigated in many studies. In NHANES I, ARIC and Rotterdam studies the high SUA level was associated with an increased risk of stroke (7, 10, 29). In NHANES I study there was a 48% increase in the risk of ischemic stroke in women for every 1.01 mg/dl increase in SUA. In ARIC study there was an independent and positive relationship between the incidence of the ischemic stroke and SUA (10).

SUA as a risk factor for the developing CAD remains controversial. In MRFIT study, the hyperuricemia and gout had an independent relationship with the risk of myocardial infarction, after adjustments for other risk factors (9). In AMORIS study a moderate increase in the SUA level was associated with increased incidence of myocardial infarction, stroke and heart failure in middle-aged subjects without prior cardiovascular disease (31). Other studies (ARIC study, Framingham study or an Austrian study) did not found an independent association between the SUA levels and the increased risk of CAD (10, 11, 32).

The aforementioned studies demonstrated the strong association of SUA with CAD, particularly in patients at high risk for heart disease and in women. The role of SUA as a causal factor for cardiovascular events in these conditions remains to be determined.

SUA AND HEART FAILURE

Hyperuricemia is a common condition in chronic heart failure (CHF). Its prevalence increases as the disease progresses (37). In a cross sectional study, 51% of patients hospitalized from chronic heart failure had hyperuricemia (30). The SUA level is higher in patients with end-stage CHF and in cachectic patients (38). It is inversely associated with functional NYHA class and maximal oxygen consumption and it is significant correlated with the severity of diastolic dysfunction (38-42).

Hyperuricemia is also an independent prognostic marker in chronic and in acute heart failure (AHF) (43, 44). In a validation study, SUA was the most powerful predictor of survival for patients with severe CHF (NYHA class III or IV): in patients with high levels of SUA (> 9.5 mg/
dl), the relative risk of death was 7.4 (44). In a study with AHF and systolic dysfunction the high SUA level was associated with higher risk of death and new heart failure readmission (45). Hyperuricemia was also an independent predictor of all-cause mortality in an unselected consecutive patients admitted with AHF (46).

Recently, hyperuricemia was associated to incident heart failure in community adults (47, 48). In the Cardiovascular Health Study the incident heart failure occurred in 21 % participants with hyperuricemia and in 18% participants without hyperuricemia. Each 1 mg/dl increase in SUA was associated to 12 % increase in incident heart failure (47). In the Framingham Offspring cohort, the incidence rates of heart failure were 6-fold higher among those at the highest quartile of SUA (>6.3 mg/dL) compared to those at the lowest quartile (<3.4 mg/dL) (48). Hyperuricemia appears as a novel, independent risk factor for heart failure in a group of young general community dwellers.

There are several mechanisms involved in hyperuricemia- induced heart failure. The increased SUA production may be due to increased XO substrate (ATP breakdown to adenosine and hypoxanthine) and to the up-regulation and increase in XO activity. When released from necrotic tissue, SUA can produce additional adverse effects on cardiovascular system and can mediate the immune response. (49). In heart failure hyperuricemia is a marker of XO activation (44).

Several studies have shown that the reduction in the SUA levels may be associated with the reduction in cardiovascular morbidity and mortality. In the Losartan Intervention for Endpoint Reduction in Hypertension (LIFE) study, the attenuation of the SUA levels by losartan was associated with 29% reduction in the composite outcome of cardiovascular death, fatal or nonfatal myocardial infarction and fatal or nonfatal stroke (50). Some of the cardiovascular benefits of atorvastatin reported in the Greek Atorvastatin and Coronary-Heart-Disease Evaluation (GREACE) study have also been attributed to the ability of statins to lower the SUA levels (51). Allopurinol and oxypurinol are XO inhibitors which has been used to treat hyperuricemia. The reducing the SUA level in HT with XO inhibitors lowers blood pressure in young with HT of recent onset (52). Other studies outline the potential benefits of XO inhibition in heart failure. In CHF allopurinol improves endothelial dysfunction, peripheral vasodilatator capacity and myocardial energy by reducing markers of oxidative stress (52). In OPT-CHF Study oxypurinol increased left ventricular ejection fraction and improved clinical outcome in CHF patients presenting with high SUA levels (53).

**URIC ACID PARADOX**

The uric acid has several biological properties which can be either beneficial or detrimental. SUA is a powerful antioxidant and it protects against free radical damage. Along with ascorbate, SUA accounts for up to 60% of the serum free radical scavenging capacity. SUA reacts with a variety of oxidants and it prevents the formation of peroxynitrite and the inactivation of the nitric oxid by superoxide anions. In individuals with hyperuricemia, the plasma total antioxidant capacity is elevated, which suggests that hyperuricemia may be a compensatory mechanism to counteract the oxidative stress damage related to atherosclerosis (54).

The SUA paradox consists in the fact that high SUA, which has antioxidant properties, is associated with an increased cardiovascular risk. It has been proposed the theory of the antioxidant, pro-oxidant redox shuttle: SUA, which under normal circumstances is an antioxidant, becomes pro-oxidant in the atherosclerotic medium with ROS generation (55). The excess of SUA has deleterious effects: endothelial dysfunction, proliferation of vascular smooth muscle cells, increases platelet adhesiveness, oxidation of LDL-cholesterol and lipid peroxidation. All these pathological processes might contribute to the pathogenesis of atherosclerosis and cardiovascular disease.

**CONCLUSION**

Clinical and epidemiological evidences have shown that the SUA level is associated to metabolic syndrome, cardiovascular disease and chronic kidney disease. Elevated SUA level has been recently recognized as a risk factor for the development of the arterial hypertension, subclinical atherosclerosis, stroke and heart failure. The role of the uric acid as an independent risk factor for the cardiovascular disease is controversial, since hyperuricemia is associated to other traditional risk factors. Elevated SUA
level also represents a strong prognostic marker for cardiovascular events, particularly in patients at high cardiovascular risk or with established cardiovascular disease. Many factors contribute to high SUA level and the mechanisms linking the urate and the cardiovascular disease are not completely understood yet. When associated with increase oxidative stress, hyperuricemia may be a marker of the increased XO activity. If SUA has a protective role as an antioxidant or a causative and deleterious role is still debatable. More prospective randomized trials lowering SUA are needed in order to clarify the role of the uric acid in the development and progression of cardiovascular disease and to establish if reducing SUA level will translate into a better cardiovascular outcome. Hyperuricemia will become then a meaningful target for the prevention and treatment of cardiovascular disease.

REFERENCES


36. Ruggiero C, Cherubini A, Miller E et al – Usefulness of uric acid to predict changes in C-reactive protein and interleukin-6 in 3-year period in Italians aged 21 to 98 years. Am J Cardiol 2007; 100:115-121
42. Cicoira M, Zanolli L, Rossi A et al – Elevated serum uric acid levels are associated with diastolic dysfunction in patients with dilated cardiomyopathy, American Heart Journal 2002; 143:1077-1111
43. Stefan D Anker, Andrew JS Coats – Metabolic, functional, and hemodynamic staging for CHF? The Lancet 1996; 348:1530