Morphological kidney and heart analysis of autopsied hypertensive patients

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This study aims to analyze the correlation between renal and cardiac fibrosis in autopsied hypertensive individuals, associating it with clinical and morphological data. This is a quantitative research, using clinical data from autopsy reports of hypertensive patients from the Federal University of Triângulo Mineiro, Uberaba/MG/Brazil, as well as fragments from these patients’ heart and kidney. A comparison of the urea and creatinine levels, considering the analyzed morphological changes (cardiac and renal fibrosis), was conducted through a laboratory analysis. After collecting the materials, the microscope plates were prepared and colored, and microscope and morphometric analysis was conducted, followed by statistical ones. In this study, no proof was found of a correlation between cardiac and renal fibrosis. The changes in the organs targeted by systemic arterial hypertension may have happened in moments and proportions that varied among individuals.

Descriptors: Hypertension; Heart; Kidney; Fibrosis; Stroke.

Este estudio tiene como objetivo analizar la correlación entre la fibrosis renal y cardíaca en individuos hipertensos autopsiados, asociándola a datos clínicos e morfológicos. Se trata de una pesquisa cuantitativa, en que fueron utilizados datos clínicos de los laudos de autópsias, provinientes de la Universidad Federal do Triângulo Mineiro (UFTM), de pacientes hipertensos, además de fragmentos de corazón y riñón de estos pacientes. Laboratorialmente, fueron verificados los niveles de urea y creatinina de estos pacientes, comparados a las alteraciones morfológicas analizadas (fibrosis cardíaca e renal). Después de la recolección de los materiales, se procedió a la confección de láminas y coloración, análisis microscópico, morfométrico, y después estadístico. En este estudio no fue comprobada la correlación entre fibrosis cardíaca e renal. No se puede decir que posiblemente las alteraciones en los órganos se hayan producido a falta de una hipertensión arterial sistémica que ocurrieron en momentos y proporciones diferentes entre los individuos.

Descritores: Hipertensão; Coração; Rim; Fibrose; Acidente vascular cerebral.
INTRODUCTION

According to the VII Brazilian Directive for Arterial Hypertension from the Brazilian Society of Cardiology, Systemic Arterial Hypertension (SAH) happens when the levels of arterial pressure (AP) are above the reference levels for the general population. It can be classified as prehypertension (121x81mmHg a 139x99mmHg) or actual hypertension, which has the stages: 1 (light - 140x90mmHg a 159x99mmHg), 2 (moderate - 160x100mmHg a 179x109mmHg) and 3 (severe - equal or above 180x110mmHg).1

The systemic arterial hypertension exposes the patient to the development of a series of functional and structural organic changes, and the hypertensive crisis is a clinical urgency that can damage many organs, leading to cerebral vascular accident, pulmonary edema, congestive cardiac insufficiency, aortic dissection, acute myocardial infarction, unstable angina, acute kidney failure and hypertensive encephalopathy.2

The SAH affects 25% of the adult world population, and the cases of the disease are predicted to grow 60% in 2025. Cerebrovascular and coronariopathy diseases have increasingly been associated to arterial pressure disturbances, which shows their high socioeconomic impact and their influence in the morbimortality of the population.3

In Brazil, the prevalence of SAH varies from 24.8 to 44.4%, making it the most prevalent of all cardiovascular diseases and affecting more than 36 million Brazilian adults. It is the greatest risk factor for cardiac and cerebrovascular lesions and the third greatest cause of disabilities4,5.

The SAH leads to a number of changes in the composition of the cardiac tissue, leading to a structural remodeling of the myocardium. Therefore, hypertensive patients are exposed to the development of the hypertensive hearth disease syndrome, which is the hypertrophy of the left ventricle in the absence of aortic stenosis and hypertrophic myocardial disease. In hypertensive cardiopathy, the hypertrophy and the apoptosis of cardiomyocytes, interstitial fibrosis and the hypertrophy of the cardiac microvasculature, through the exacerbated retention of collagen fibers types I and III in the interstice and around the intramyocardial arteries and arterioles, constitute the structural elements that define myocardial remodeling. The functional consequences of such remodelling are diverse, and the most representative is the development of congestive cardiac insufficiency6,7.

The essential SAH has devastating effects on the brain, associated to high rights of morbimortality. It is the main risk factor for cerebral vascular accidents and the main cause for cognitive deficits and dementia.8 Hypertensive encephalopathy is a neurological syndrome brought forth by a sudden and sustained increase in systemic blood arterial pressure. It can be asymptomatic or lead to silent cerebral infarction and symptoms such as headaches, mental confusion, vision alterations, severe hypertension, vomit, sensory alterations and convulsions.9

Chronic hypertension leads to a structural adaptation of cerebral vascular resistance, and consequently, to an elevation of the cerebral self-regulation threshold. Thus, hypertensive encephalopathy is believed to result from a sustained increase in blood pressure that exceeds the upper limit of self-regulation, dilating the cerebral arterioles and dissolving the blood-brain barrier, generating vasogenic edemas.2

Although reversible if readily recognized and treated, hypertensive encephalopathy can evolve into coma and even death, if treatment is late. Even if it occurs in people with primary hypertension, it is more common in the mutual presence of hypertension and kidney disease, and it may be a complication of kidney transplants.10

The mechanism through which SAH could cause kidney lesions can be divided into three categories: glomerular ischemia secondary to vasoconstriction, glomerulosclerosis due to intracapillary hypertension, and interstitial fibrosis. Goldblatt et al., in the decade of 1930,
demonstrated that the reduction of kidney perfusion can generate a sustained elevation of blood pressure, subsequently associated to the renin-angiotensin-aldosterone system\textsuperscript{11,12}. Aldosterone plays an important role in the pathogenesis of hypertension, vascular remodeling, left ventricular hypertrophy and kidney diseases, especially those which cause with proteinuria and glomerulosclerosis in hypertensive individuals\textsuperscript{13}.

In the advanced phase of kidney failure, aldosterone values increase significantly with the decrease of glomerular filtration due to the activation of the renin-angiotensin-aldosterone system, secondary to the alteration of glomerular hemodynamics. This contributes to the lesion of target organs, affecting primarily kidneys, brain and heart\textsuperscript{13}.

Kidney interstitial fibrosis is the final pathway for almost all forms of renal disease. The study of fibrosis is of great value for several kidney diseases that become terminal. During SAH, such a study can be worthwhile, since interstitial fibrosis, in addition to being a bad prognosis, is an efficient marker to evaluate all pathways that injure the kidney in the SAH (ischemia, glomerular hypertension and inflammatory state)\textsuperscript{14}.

Chronic kidney disease has demonstrated to be an independent risk factor for cardiovascular disease, which in turn is the main cause of morbidity and mortality in chronic kidney disease patients. Similar to this, numerous studies indicate that chronic kidney diseases are associated with the high prevalence of cerebral vascular accidents\textsuperscript{15}.

Hypertension is a systemic disease with high morbidity and mortality, and thus, it becomes important to understand the relationship between the organs affected by the SAH, and compare the intensity of lesions such as cardiac fibrosis, kidney fibrosis and the presence of CVAs in autopsied hypertensive patients. The present study aims to analyze kidney and heart fibrosis in autopsied subjects, associating them to clinical and morphological data.

**METHOD**

Clinical data of the autopsy reports of twenty-three hypertensive patients were used, all from the General Hospital of the Federal University of the Triângulo Mineiro (HC-UFTM), Uberaba, Brazil, and who had been studied in General Pathology classes in the period from 1986 to 2007. Heart and kidney fragments from these patients were also used for morphometric analyses, laboratory data (urea and creatinine levels compared to morphological changes) and morphological aspects of autopsied patients (cardiac and kidney fibrosis).

This study included patients who presented hypertensive heart disease and/or kidney sclerosis (benign or malignant) as changes provoked by the SAH. Patients with primary kidney diseases (glomerulopathy, interstitial nephritis such as pyelonephritis, vasculopathy) or carriers of systemic diseases damaging to the kidney (diabetes, lupus erythematosus, hepatitis or acute or chronic inflammatory diseases) were excluded from the study.

To analyze the kidney material, fragments of the middle pole of the right kidney were used, and to analyze the cardiac material, fragments of the middle third of the left ventricle were removed from the autopsied patients. Após o processamento, o fragmento parafinizado foi submetido a cortes seriais de espessura adequada para confecção das lâminas. Then, the colorations of picro-sirius (PS) and hematoxylin and eosin (HE) were carried out.

The microscope plates that had been colored with picro-sirius were used to analyze cardiac fibrosis and quantify the collagen in the mesangial matrix. The morphometry was performed through the capturing of images subsequently sent and recorded for analysis in the ImageJ software (an Image ProPlus software). The morphometric analysis of the fibrous tissue was performed in the kidney and heart samples using digital morphometry. In the polarized image, the fibrous connective tissue showed itself to be birefringent and was marked by the observer, and through it, the percentage of fibrosis per area of the analyzed field was obtained, as shown in Image 1.
For the statistical analysis, a spreadsheet was created in Microsoft Excel. The clinical, laboratory and morphological variables were tested to verify that they presented normal distribution, through the test of Kolmogorov-Smirnov. The following tests were used: student’s t-test (to compare two groups with parametric variables) Mann-Whitney’s test (to compare two groups with non-parametric variables), Pearson’s (to correlate groups that presented normal distribution) and Spearman’s correlation (to correlate non-parametric samples between groups). The differences were considered statistically significant when p was less than 5% (P < 0.05).

**Image 1.** Computer system for the quantification of fibrosis. Autopsied patients from 1986 to 2007, HC-UFTM. Uberaba, MG, 2016.

Automatic microscopic computed morphometry. In (A) the light microscope can be seen, connected to a camera which in turn is connected to the computer. In (B) the steps followed by the KS 300 Carl Zeiss software can be seen; in (C), the capture of the image to be quantified with polarized light; and in (D), the result of the quantification of the captured image, in percentages.

**RESULTS**

The age average of the 23 autopsied hypertensive patients was 59 years of age (SD=16.14), 54.5% of which were elders (>65 years old) (Image 2A).16 patients (72.6%) were male (Image 2B). The most common cause of death was cardiovascular (45.5%), followed by infection (27.3%), and by digestive (22.7%) and neoplastic (4.5%) causes (Image 2C). The average Body Mass Index (BMI) was 22.27 (SD=4.96) (Image 2D). The average cardiac fibrosis, on the other hand, was 2.78% (SD=1.03) while renal fibrosis had an average of 12.91% (SD =2.62) (Image 3).


Percentage of cardiac fibrosis and kidney fibrosis. Horizontal lines represent the averages and vertical lines represent the standard deviation from the average. Statistical analysis calculated through Student's T-Test. A positive but not significant correlation was observed between the percentage of cardiac and kidney fibrosis (p = 0.15 and r = 0.31) (Image 4).


Note: correlation of the percentage of cardiac fibrosis (Y-axis) relative to the percentage of kidney fibrosis (X-axis). Statistical analysis conducted through Spearman's correlation test.
Among the hypertensive patients evaluated in this study, 16 (72.7%) had had previous episodes of cerebral vascular accidents. From these, 80% had hemorrhagic CVAs and 20%, ischemic CVAs.

When comparing the percentage of cardiac fibrosis to that of kidney fibrosis among the patients who had had CVAs and those who had not, a statistically significant difference was not observed for any of the two measures (P=0.52 and P=0.53, respectively) (Images 5A and B).

**Image 5.** Percentage of cardiac and kidney fibrosis according to CVA. Autopsied patients from 1986 to 2007. Uberaba, MG, 2016.

When comparing the percentage of cardiac fibrosis to that of kidney fibrosis among the hyperuricemic patients and those who had normal urea levels, a statistically significant difference was not observed for any of the two measures (P=0.59 and P=0.76, respectively). Similarly, there were no statistically significant differences in the comparison between cardiac fibrosis and kidney fibrosis, when comparing normal serum creatinine levels to those who had alterations in these levels.

**DISCUSSION**

Systemic arterial hypertension (SAH) is responsible for approximately 46% of all deaths in Brazil. This disease, frequent and disturbing, affects several organs, such as heart, kidney and brain, which are therefore said to be the target organs of SAH. The mechanisms through which SAH damages target organs and leads to vascular events are still not well-known.

The SAH is closely linked to the kidney, because lesions in that organ can be both the cause and the consequence of high blood pressure. The brain is another target organ of the SAH, since it leads to the deterioration of the walls of cerebral arteries, causing cerebral vascular accidents (CVAs), as the final results of predisposing conditions. Some studies suggest an intimate relationship between blood pressure and CVA mortality, being that among patients treated with antihypertensive medication, an increase of one mmHg in the systolic arterial BP increases mortality by CVA in 2%.

In this work, most individuals were male, corroborating previous studies. Cardiovascular complications were the main causes of death in this study. The BMI of the patients was within a normal range (< 25) suggesting that the SAH can occur in non-obese patients, although literature findings show that increased body fat is associated with an increase in the risk of SAH.

Within the morphological alterations of hypertensive heart disease, fibrosis is one of the most common changes among hypertensive patients. It is characterized by
excessive collagen fibers or their diffuse accumulation, increasing myocardial rigidity, which can lead to a dysfunction of the left ventricle, finally culminating in cardiac insufficiency. In this study, not all patients presented fibrosis of the heart tissue in large quantities. These results may be due to the intensity of the SAH and to the age group of the individuals analyzed in the study, since some individuals were non-elderly adults, with an average age of 59 years, perhaps not yet presenting an increase in the accumulation of collagen fibers, corroborating works that show that the elderly are more susceptible to fibrosis, since in this age group there is an increase in the synthesis and a decrease in collagen degradation.

One of the main morphological findings in the kidney of hypertensive patients is that glomerulosclerosis may occur through 3 main pathways: glomerular ischemia due to vascular lesion; glomerular hypertension, through the loss of self-regulation; and through the activation of the renin-angiotensin system (RAS). All these pathways can compromise not only kidney vessels (arterial hyalinosis, fibrinoid necrosis) or the glomerulus (glomerulosclerosis), but also the kidney interstitium, leading to fibrosis.

In this present study, the percentage of fibrosis in the medium portion of the right kidney was evaluated, showing areas with the presence of collagen in all evaluated individuals, corroborating a work in which the increase in the production and an accumulation of collagen fibers type I and III was proportionate to the intensity of the lesions caused by the SAH.

In this study, more than 70% of individuals presented episodes of CVA, and 80% of these were hemorrhagic. This can be justified by the fact that hemorrhagic CVAs seem to be directly related to the elevation of blood pressure, while the ischemic CVA is explained by atherosclerotic lesions, although these are also related to the SAH.

In this study no correlation was found between cardiac and kidney fibrosis; what explains this result is the fact that some patients presented cardiac fibrosis in small quantities, even presenting high kidney fibrosis. Literature, however, reports that the increase of arterial BP is a determining factor that acts on the structure of the heart. Similarly, there was no relationship was observed between the intensity of cardiac and kidney fibrosis and CVA episodes. No previous reports were found in the literature highlighting this relationship. Considering this, it is suggested that the involvement of the target organs can occur at different times and that they are very changeable, from one individual to another. However, studies show that there is a higher prevalence of CVA in hypertensive patients with glomerulosclerosis than in those without it.

The laboratory analysis of the patients who presented episodes of CVA and those who did not found a similarity between their levels of urea and creatinine. However, concerning creatinine concentration, studies show that high concentration levels are significantly related to CVA cases in patients with or without hypertension. In another study, however, urea and creatinine levels showed no significant difference between patients with CVA and patients from a control group.

CONCLUSION

The most common causes of death had cardiovascular origins, and all patients presented cardiac and kidney fibrosis. Although the correlation test between cardiac and kidney fibrosis was positive and regular, it was also not significant.

72% of the surveyed patients had had previous instances of cerebral vascular accident, 80% of the hemorrhagic type. Considering all patients, the percentages of cardiac and kidney fibrosis were not significant.

The percentage of cardiac and kidney fibrosis did not vary between patients who presented or not changes in urea (p=0.59 and p=0.76, respectively) and creatinine levels (p=0.74 and p=0.67, respectively).

It can be concluded that, in the group of patients surveyed in the period considered, among the individuals with systemic arterial
hypertension, there is kidney fibrosis and cardiac fibrosis, as well as other changes in the target organs of systemic arterial hypertension, which probably took place at different times and in varying proportions from one individual to the other.

REFERENCES

CONTRIBUTIONS
All authors contributed equally in the various stages of the research and in the writing of the article.