**ABSTRACT**

**Introduction**
Obesity is a social disease constituting a global pandemic. It is present in 90% of diabetic and 65% of hypertensive patients. It is associated with cardiometabolic syndrome and with damaging physiological mechanisms, particularly for the vascular system and the kidneys. On Cuba’s Isle of Youth, a community-based epidemiological study of chronic kidney disease (CKD), hypertension (HTN), diabetes mellitus (DM) and cardio-cerebral vascular disease was carried out in total population, including an examination of common risk factors.

**Objective**
Based on the Isle of Youth Study (ISYS) data, determine prevalence of obesity and overweight, and their association with CKD, HTN, DM, and shared risk factors for all these conditions in total population of the Isle of Youth, Cuba.

**Methods**
Phase 1: Population diagnosis (November 2004-April 2006): 96.6% of the Isle of Youth’s total population (80,117) was studied, including all ages and both sexes. Information was offered to the public, and written informed consent obtained. Screening was conducted by participant questionnaire including risk factors, physical measurements (weight, height, blood pressure and body mass index), and a single first-morning urine sample to determine the presence of vascular-renal damage markers – proteinuria and hematuria (Combur 10 Test, Roche), and microalbuminuria (Micral Test, Roche). When results were positive, serum creatinine was determined and glomerular filtration rate (GFR) estimated with Modification of Diet in Renal Disease (MDRD) formula for adults and Schwartz formula for children <15 years. Data obtained were analyzed to determine association of participants’ nutritional status with prevalence of chronic kidney disease, hypertension and diabetes mellitus, as well as a set of common risk factors.

**Results**
Population <20 years: Obesity prevalence 3.2%; positive urine markers in 56.9% of obese and 8.9% of non-obese participants; positive albuminuria in 38% of obese and 3% of non-obese. Obese participants were found to be hyperfiltrating. DM prevalence was 9.5% in obese and 1.1% in non-obese participants. Population ≥20 years: Overweight, 31.3%; obese, 13.4%. Positive markers in normal-weight, overweight and obese individuals were 18.3%, 21.2% and 32.7%, respectively; microalbuminuria values increased with weight. Obese individuals were found to be hyperfiltrating. HTN prevalence in normal-weight, overweight and obese individuals was 18.3%, 31.5%, and 51.0% respectively; DM rates were 2.8%, 5.2% y 11.3%, respectively.

**Conclusion**
In Cuba, obesity poses significant risk for vascular and renal damage and should be the focus of increased prevention efforts.

**Keywords:** Vascular diseases, chronic kidney disease, CKD, chronic kidney insufficiency, hypertension, diabetes mellitus, risk factors, obesity, overweight, body mass index, glomerular filtration rate

**INTRODUCTION**

**Background:** After World War II, industrial-scale production was developed to manufacture relatively cheap, easy-to-prepare and highly caloric food products. These became accessible to a share of the global population, among whom obesity has since become a social disease of pandemic proportions.[1]

Global estimates put the number of overweight people at 1.7 billion and obese at 312 million, weight gain directly proportional to the aging of populations. Simultaneously, the new phenomenon of childhood and adolescent obesity already affects 155 million persons worldwide. The largest increase in the obesity pandemic is foreseen in middle-income developing countries.[2-5]

In North America, 49.6% of Canadians are overweight including 15.7% obese.[6] Some 66% of people in the United States are overweight, including 32% obese.[7] In Mexico, 69% are overweight, including 30% obese.[8]

On a social scale: excessive consumption of hypercaloric foods has been associated with genetic predisposition for weight gain, and obese mothers tend to deliver infants of above-average height and weight. The number of overweight and obese children and adolescents is increasing to alarming levels. A family’s eating habits can lead to obesity for its members. Among couples with one obese partner, it has been reported that 37% of spouses follow suit; the same applies to 40% of twins and 57% of very close friends.[9]

Obesity often coexists with other risk factors such as smoking and alcohol consumption, and is associated with vascular-damage diseases such as diabetes mellitus (DM), hypertension (HTN) and chronic kidney disease (CKD). Globally, obesity is present in 90% of type 2 diabetic patients and in 65-75% of patients with HTN. By 2000, there were 171 million diabetics globally, and their number is expected to reach 366 million by 2030, the largest increase predicted for developing countries. In 2000, 1 billion HTN
patients were reported, their numbers expected to reach 1.56 billion by 2025.[2]

Diabetes and hypertension are the two main causes of CKD. In the United States, for 72.4% of patients who started renal replacement therapy in 2003, diabetes or hypertensive vascular disease was the underlying cause of renal failure.[10]

Globally, prevalence of patients in dialysis increases 4-10% annually, their total number expected to double in this decade.[11] Extrapolating from a study of CKD prevalence carried out in the United States, over 600 million people worldwide may already suffer from some stage of CKD.[12]

A 17-year longitudinal study begun in 1983 of 100,863 subjects aged ≥20 years (Iseki in Okinawa) showed that CKD cumulative incidence increased with body mass index (BMI).[13] Hsu obtained similar results in the United States.[14]

The question then arises: Is obesity just excess fat? Excess fat is only the visible manifestation of obesity, the tip of the iceberg. Inside the body, major metabolic, hormonal, hemodynamic, biochemical, immunological and molecular changes are taking place – the basis for obesity’s systemic damage.[15] Thus, obesity is associated with a number of pathological conditions such as:

Cardiometabolic syndrome: Clinical observation has demonstrated frequent association between the anthropometric parameters of obesity (abdominal circumference ≥102 cm in men or ≥88 cm in women) and metabolic parameters such as lower glucose tolerance (≥110 mg/dl) and dyslipidemia (triglycerides ≥150 mg/dl, and HDL <40 mg in men or <50 mg in women), as well as hemodynamic parameters such as blood pressure ≥130/80 mm Hg. Based on this, the US National Cholesterol Education Program has concluded that if three of the above elements are present, a person suffers from cardiometabolic syndrome[16] and thus is at high risk for cardiovascular disease and systemic vascular damage.[17-19] The World Health Organization (WHO) has confirmed this assessment by including albuminuria (a vascular and renal damage marker) as another element to diagnose and describe this syndrome.[20]

Obesity-related glomerulopathy (ORG): In 1974, Weisinger reported an association between nephrotic syndrome and severe obesity.[21] In 1975, Cohen described obesity-related glomerulopathy as characterized by an increase in glomerular volume, slight hypercellularity, variable mesangial enlargement, glomerular basal membrane thickening and, occasionally, segmental and focal sclerosis.[22]

There are two histopathological presentations of ORG: glomerulomegaly and obesity-related focal segmental glomerulosclerosis (O-FSGS). Recent studies indicate (also comparing O-FSGS with primary focal segmental glomerulosclerosis, P-FSGS) that O-FSGS is clinically characterized by moderate proteinuria, little edema, high serum albumin, low cholesterol, lower incidence of nephrotic syndrome and slower progression towards end-stage renal disease (ESRD).[23]

A 15-year study conducted by Kambian of 6,818 original renal biopsies reported a tenfold increase in the incidence of kidney disease during that period and described the renal histopathology.[24] Optic microscope revealed presence of glomerulomegaly in 100% of cases; sclerosis was moderate and was not found frequently at the corticomedullary union, as was the case with P-FSGS. Moreover, there were perihilar and peripheral segmental lesions, podocytes were hypertrophied, loose and cap-shaped, and pedicels were fused. Particular emphasis was placed on the presence of focal “diabetoid” changes in the form of slight focal mesangial sclerosis and moderate focal thickening of glomerular and tubular basal membranes. Immunofluorescence revealed the presence of IgM and C3 glomerular deposits.

Nephrolithiasis: In a 46-year retrospective study of 4,827 kidney stone patients, Taylor observed that men weighing >100 kg had a 1.44 relative nephrolithiasis risk when compared to those weighing <68 kg. Comparing individuals with a ≥30 kg/m2 BMI to those with a 21-22.9 kg/m2 BMI, he found that the relative risk was 1.33 for men and 1.90 for women.[25] Taylor concluded that hyperinsulinism reduced ammoniogenesis and thus urine pH, and caused hypocitraturia and increased calcium excretion. Moreover, obese individuals also excreted more uric acid and oxalate. The combination of these factors facilitated development of kidney stones.

METHODS

The first phase of the “Community-Based Epidemiological Study of Chronic Kidney Disease, Cardiovascular Disease, Diabetes Mellitus and Hypertension (ISYS)” was carried out in total population on the Isle of Youth, Cuba, November 2004-April 2006. Phase 1 consisted of active screening for renal-vascular damage markers and associated risk factors for these conditions in total population of the Isle of Youth, Cuba.

ISYS’ general methodology has been published previously.[26] Particularly relevant is the fact that the Isle of Youth’s population is composed of a representation of all Cuban provinces, and its age and gender demographics closely resemble those of the rest of the country, except for persons aged >60 years, whose percentage on the Isle of Youth is slightly lower.[27] Of the 80,117 persons physically resident in the Isle of Youth, distributed in 21,115 family-household units, 77,423 (96.6%) were screened; their demographics the same as the Isle’s total.

CKD was used as a tracer disease in the ISYS study since both renal damage and vascular damage markers can be detected in urine, thus revealing the presence of other chronic vascular diseases.

During Phase 1 of ISYS, public information about the study was provided, and written informed consent obtained from participants. A survey was conducted on family and individual history of vascular disease and risk factors, using the Framingham score.[28] Blood pressure, weight and height were measured. In
a single first-morning urine sample, vascular and renal damage markers were studied: hematuria and proteinuria (Combur-10 Test, Roche) in total population screened, and microalbuminuria (Micral Test, Roche) in risk groups. Risk groups for vascular-renal damage were defined as individuals with obesity, HTN, DM, a history of CKD or chronic vascular disease, children aged <5 years and adults >60 years.

A cohort of 14,322 individuals (18.5%) was obtained, who tested positive for these markers. Blood samples were taken to determine serum creatinine and glomerular filtration rate (GFR), estimated with MDRD (Modification of Diet in Renal Disease) formula for persons aged ≥15 years and with Schwartz’s formula for children aged 2-15 years. CKD stages were then classified for those aged ≥2 years: stage 1, ≥90 ml/min; stage 2, 89-60 ml/min; stage 3, 59-30 ml/min; stage 4, 29-15 ml/min; stage 5, <15 ml/min. Stages were correlated with population demographics, two risk factors (obesity or overweight and age) and other chronic vascular diseases (DM and HTN in population aged ≥20 years and DM in those <20 years).

The screened population was divided into age groups, and nutritional status was determined by body mass index (BMI = kg/m²). Population aged ≥20 years were classified as: underweight, <18.5 kg/m²; normal weight, 18.5-24.9 kg/m²; overweight, 25.0-29.9 kg/m²; and obese, ≥30.0 kg/m². For population <20 years, the same BMI formula was used; nutritional status assessment was based on age, sex and percentiles, and persons were classified as obese or non-obese. For this portion of the study, relation between BMI and established risk groups was established for 20,649 children and adolescents aged <20 years, and 52,430 adults aged ≥20 years.

All data was introduced into a Microsoft Access database, and a qualitative and quantitative data validation system was employed with minimum and maximum value intervals to minimize error margin.

Frequency distributions of kidney damage markers – proteinuria, proteinuria plus hematuria, microalbuminuria and isolated hematuria – were calculated for cases testing positive within the total population studied. Frequency distributions were constructed to classify renal function, based on GFR estimates for cases aged ≥2 years with positive kidney damage markers whose serum creatinine levels were measured.

Likewise, double-entry tables were created to relate this classification to the following variables for participants aged ≥20 years: sex, age, kidney damage markers, diabetes and hypertension. For participants aged 2-19.9 years, all the above variables were considered except hypertension.

Frequency distributions were calculated for various combinations of four main risk factors: diabetes, hypertension, obesity and overweight. For all combinations, a cross-classification table was also created to reflect status of renal function.

RESULTS

Population aged <20 years

Obesity prevalence was 3.2% in this age group. Urine vascular-renal damage markers were positive in 56.9% of obese and 8.9% of non-obese individuals. These parameters were similar for all ages within this group. Frequency of each positive marker taken alone was higher in obese than in non-obese individuals.

Positive microalbuminuria within risk groups was 38% for obese and 3% for non-obese individuals.

In all ages <20 years, obese individuals were hyperfiltrating in comparison to non-obese subjects (Figure 1), an expression of glomerular capillary hypertension, which induces kidney damage.

DM prevalence for this age group was 1.3%-9.5% for obese and 1.1% for non-obese individuals. This distribution was the same for all ages within this group (Figure 2).

Population aged ≥20 years

In the adult population, nutritional status analysis revealed 6.0% underweight, and 44.6% with BMI ≥25 kg/m²; 31.3% were overweight and 13.4% obese. Overweight and obesity increased with age and risk was higher after 40 years. Percentage of underweight persons increased from age 60 years forward.

Figure 1: Obesity and Glomerular Filtration Rate (GFR) in Population <20 Years by Age Group; ISYS, Cuba

Figure 2: Obesity and Prevalence of Diabetes Mellitus (DM) in Population <20 Years by Age Group; ISYS, Cuba
Correlating nutritional status with positive urine markers revealed higher risk among persons underweight (46.5%) or obese (32.7%) than among persons of normal weight (18.3%) or who were overweight (21.2%), although the risk was higher in the latter than in those whose weight was normal. Higher risk for individuals at both extremes of nutritional status was present in all age groups, but older age brought added risk of obesity and, after 70, risk distribution took the form of a U-shaped curve (Figure 3).

Among risk groups, positivity for microalbuminuria was high in underweight persons in all age groups; it increased proportionately with weight above normal ranges (overweight, obese) and with age (Figure 4).

In the adult population, underweight, normal-weight, overweight and obese subjects had mean GFR values of 107 ml/min, 91 ml/min, 101 ml/min and 127 ml/min, respectively, for a J-shaped curve with the highest point at obesity.

HTN prevalence correlated directly with weight gain, reaching its peak in obese individuals, among whom 51% were hypertensive, while mean prevalence in adults was 26.5%. HTN prevalence was 13.3% in underweight individuals, 18.3% in normal-weight, and 31.5% in overweight. Stratifying correlation between hypertension and nutritional status by age group revealed exactly the same tendency, where age was compounded by obesity (Figure 5).

DM also increased with greater weight, and DM prevalence nearly tripled in obese persons (11.3%) when compared to DM in the adult population (4.7%). Prevalence in underweight, normal-weight and overweight was 3.6%, 2.8% and 5.2%, respectively. This tendency was observed in all adult age groups, and prevalence increased with age (Figure 6).

Classifying CKD by stages and correlating to nutritional status revealed a higher percentage of persons who were overweight or obese in stages 2 and 3 of renal failure, in contrast with those whose weight was normal or low (Table 1). Age-adjusted risk analysis for adults, calculated without considering HTN and DM, showed a linear gradient of greater risk with increased age.

Risk quantification adjusted for nutritional status, without HTN and DM, clearly revealed the same inverted-J risk distribution ob-
The decline observed in obesity prevalence coincided with reduced food consumption by the Cuban population during the country’s severe economic crisis of the 1990s, known as the “Special Period.”

The 2nd National Survey on Risk Factors and Chronic Diseases in the Cuban Population[31] carried out in 2000-2001 showed 42.5% of persons aged >15 years with BMI>24.9 kg/m², of whom 11.8% were obese (BMI>30). ISYS data obtained among adults coincide with these earlier results, and constitute a call for prevention of continued increase in obesity prevalence.

Observing the last 15 years in Cuba, prevalence of obesity decreased in the mid 1990s and has gradually increased during subsequent years. Several studies bear this out. In Havana’s 10 de Octubre municipality, adults were studied between 1988 and 1994, obtaining an obesity prevalence of 18.8% and 6.3%, respectively.[29] Other research has revealed an obesity prevalence in Havana Province of 11.9% (1982), 5.4% (1994) and 9.3% (1998); in Cienfuegos Province, of 14.3% (1990), 7.2% (1995) and 12.1% (2001).[30]

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ISYS results corroborate that obesity in Cuba is a risk factor for atherosclerosis-linked chronic vascular disease, a risk that increases with age. These vascular diseases clinically manifest in different forms depending on the main organ affected, usually presenting co-morbidity in which each condition enhances the damage of the other, constituting a systemic multiplier of damage. As shown in previous studies, overweight and obesity in Cuba is gradually increasing, a situation that merits particular and comprehensive attention by all sectors that influence the population’s health.

Table 1: Stage of Chronic Kidney Disease (CKD) and Nutritional Status in Population ≥20 Years (MDRD Formula); ISYS, Cuba

<table>
<thead>
<tr>
<th>CKD Stage</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Underweight</td>
<td>63.0%</td>
<td>30.8%</td>
<td>5.0%</td>
<td>0.2%</td>
<td>1.0%</td>
</tr>
<tr>
<td>Normal</td>
<td>59.0%</td>
<td>34.3%</td>
<td>5.6%</td>
<td>0.6%</td>
<td>0.5%</td>
</tr>
<tr>
<td>Overweight</td>
<td>54.3%</td>
<td>39.1%</td>
<td>6.0%</td>
<td>0.2%</td>
<td>0.4%</td>
</tr>
<tr>
<td>Obese</td>
<td>52.5%</td>
<td>40.6%</td>
<td>6.3%</td>
<td>0.5%</td>
<td>0.2%</td>
</tr>
</tbody>
</table>

**DISCUSSION**

ISYS results show that risk of vascular-renal damage is as great for underweight as for obese persons. One hypothesis to explain this finding is that among the underweight cohort are the most elderly people, who may have been nutritionally compromised due to kidney damage, as well as those who suffer from type 1 diabetes from an early age. This group should be studied separately and more thoroughly.

**Physiopathological Mechanisms of Obesity Responsible for Vascular and Renal Damage**

Adipose tissue is an active endocrine organ that secretes several adipokines with auto-, para- and endocrine effects that have inflammatory, mitogenic, hemodynamic and other functions. Adipokines include: leptin, adiponectin, resistin, angiotensin II (AgII), TNF-α, PAI-1, interleukin (IL)-1-6-8-10, prostaglandins, HGF, VEGF, IGF-1, estrogens, and others. Visceral adipose tissue has more endocrine activity and increases structurally and functionally with obesity; thus the importance of abdominal obesity.[32,33]

An excess of adipokines reduces the amount of glucose-transporting intracellular proteins (Glut 1-Glut 4). Concurrently, obesity increases cell endoplasmic reticulum stress. The combination of these two factors causes an increase in tissue resistance to insulin, which results in hyperinsulinism.[20,34,35] Hyperinsulinism acts on various biomolecules and their receptors causing:

- Increased sympathetic activity, salt sensitivity and AgII and ET levels and decreased nitric oxide and natriuresis, leading to systemic hemodynamic changes, hypertension and systemic vascular damage;
- Increased glomerular capillary pressure, hyperfiltration, podocyte damage and albuminuria; and
- Release of mitogenic factors (IGF-1, TGF-β, PAI-1) that induce mesangial cell proliferation and greater glomerular matrix deposition.

HTN and efferent arteriole vasoconstriction generated by adipokines and hyperinsulinism increase glomerular capillary pressure. At the same time, the metabolic demand resulting from increased body fat increases cardiac output and renal plasma flow, resulting in greater glomerular capillary hypertension, hyperfiltration and albuminuria, leading to tubulointerstitial damage.[36]

Mononuclear cells in the adipose tissue matrix produce oxidizing radicals, with well-known deleterious systemic effects which, added to those caused by AgII and TGF-β induction, causes an increased glomerular matrix deposition.[37] Obesity increases the reactivity of medullary baroreceptors, thus increasing the adrenosympathetic activity—another mechanism that contributes to HTN.[15]

Seventy-five percent of people with sleep apnea are obese. Apnea causes hypoxia, which releases so-called “hypoxia-induced factors (HIF)”; these allow cells to adapt to low oxygen levels. The persistence and intensity of this physiological mechanism causes inflammatory cell infiltration and higher levels of TGF-β, CTGF, metalloproteinase inhibitors and PAI-1, which cause inflammation, apoptosis, podocyte damage, and renal fibrosis.[38]

Physiopathological changes promoted by obesity include: increased production of adipokines, hyperinsulinism, dyslipidemia, glomerular capillary hypertension, oxidative stress, stimulation of the renin-angiotensin-aldosterone system, hypoxia, inflammation and others. These integrate agonistically into a perpetual system of kidney cell damage. Kidney cells do not remain a passive target: not only are they damaged but they also react and change phenotypically. They are transformed into immunocompetent cells which become hypertrophied, proliferate, present antigens, and liberate vasoactive factors, inflammatory cytokines, growth factors, matrix proteins, etc. This results in a reverberating damage-producing system leading to glomerulosclerosis and accelerating progression of any underlying renal disease, also causing primary kidney disease.[39-42]
At the same time, ISYS provides further evidence that obesity should be understood as a chronic disease that is genetically based, social in origin, and pandemic in behavior. It is expressed externally by excess fatty accumulation and internally by profound metabolic disturbances, and poses increased risk for development of chronic vascular conditions.

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THE AUTHORS

Raúl Herrera Valdés (Corresponding Author: raul.herrera@infomed.sld.cu) specialist in nephrology, full professor, Higher Institute of Medical Sciences of Havana, senior researcher and academician, Institute of Nephrology, Havana, Cuba.

Miguel Almaguer López, specialist in nephrology, associate professor, Higher Institute of Medical Sciences of Havana, senior researcher, Chief of Preventive Nephrology Services, Institute of Nephrology, Havana, Cuba.

José Chipi Cabrera, specialist in nephrology, instructor, Isle of Youth Faculty of Medicine, Head of Nephrology Service, Héroes de Baire Hospital, Isle of Youth, Cuba.

Orquidea Martínez Soto, specialist in nephrology, Héroes de Baire Hospital, Isle of Youth, Cuba.

Osvaldo Castellanos Rabanal, specialist in epidemiology, Municipal Health Director, Isle of Youth, Cuba.
Original Scientific Articles

Jorge Bacallao Gallestey, professor, Higher Institute of Medical Sciences of Havana, Atherosclerosis Research and Reference Center, Havana, Cuba.

Rosa M. Licourt Pérez, specialist in family medicine, Dr. Juan M. Páez Incháustegui Teaching Polyclinic, Isle of Youth, Cuba.

Pedro Mulet Pineda, specialist in family medicine, Dr. Juan M. Páez Incháustegui Teaching Polyclinic, Isle of Youth, Cuba.

Luisa Diéguez Pérez, specialist in family medicine, Leonilda Tamayo Matos Teaching Polyclinic, Isle of Youth, Cuba.

Israel Velásquez Batista, specialist in family medicine, Dr. Juan M. Páez Incháustegui Teaching Polyclinic, Isle of Youth, Cuba.

Maria C. Hernández Rivero, specialist in family medicine, Orestes Falls Oñate Teaching Polyclinic, Isle of Youth, Cuba.

Walfrido Caballero Odio, nurse, Orestes Falls Oñate Teaching Polyclinic, Isle of Youth, Cuba.

Pedro Urra González, Director, National Medical Sciences Information Center and founder and Director of INFOMED, Havana, Cuba.

Néstor Rodríguez Triana, Informatics Department, Héroes del Baire Hospital, Isle of Youth, Cuba.

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