Association between adiposity and systemic atherosclerosis: a protocol of a cross-sectional autopsy study

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INTRODUCTION

Ischaemic heart disease and stroke were the leading causes of death worldwide in 2015 and were estimated to be responsible for 14.8% and 11.7% of all deaths, respectively. In 2012, 14.1 million people died from these diseases.1 2 Moreover, ischaemic heart disease is the leading cause of DALYs (Disability-Adjusted Life Years) in the world (2342 per 100 000 cases), and the stroke is the third leading cause of DALYs (1998 per
100 000 cases). Therefore, the identification of risk factors is important in preventing atherosclerosis and decreasing the cardiovascular disease burden.

Measures of adiposity, such as anthropometric and thoracic and abdominal visceral fat measurements, have been associated with atherosclerosis. However, most of the previous studies have investigated the association between anthropometric and/or visceral fat measurements with systemic atherosclerosis using imaging methods such as ultrasound, CT, angiography and echocardiography. In a review study of Alexopoulos et al., all studies that investigated the association of visceral adipose tissue with atherosclerosis used imaging methods.

Several disadvantages of these methods are well known, for instance, angiography is an invasive method that exposes the individual to radiation; CT also exposes patients to ionising radiation and the exact anatomical borders of structures in non-contrasted exams are difficult to define using imaging methods, and paracardial fat is difficult to delimit by echocardiography. Although postmortem examination provides a more accurate measure of atherosclerosis and visceral fat due to the direct assessment of these structures, few autopsy studies compared atherosclerosis to anthropometric measurements and visceral fat. In addition, none of these studies evaluated atherosclerosis at sites other than coronary arteries. Most of these studies also had small sample sizes and were limited to middle-aged or young adults.

We aim to investigate the association between adiposity measured by the weight of the visceral fat and anthropometric measurements with morphometric measures of atherosclerosis in the aorta, coronary, carotid and cerebral arteries in a large sample from different age groups.

METHODS AND ANALYSIS

Study design
A cross-sectional observational study is designed.

Study setting
This study will be conducted at the Sao Paulo Autopsy Service (SPAS), Laboratory of Cardiovascular Pathology and the Pathophysiology in Aging Lab from the Department of Pathology, University of Sao Paulo, Sao Paulo, Brazil.

Since 1931, the SPAS has performed autopsies of individuals who died in the city of Sao Paulo and who have had an unknown natural cause of death. It performs more than 13,000 autopsies per year. In 2004, we started a successful initiative at the SPAS to investigate brain ageing and neurodegenerative diseases. The Brain Bank of the Brazilian Aging Brain Study Group (BB-BABSG) has collected more than 3,000 brains over the last decade to provide the materials and data for a number of publications in high-quality international journals. We will extend the work of the BB-BABSG to evaluate the atherosclerosis process.

Study participants
A convenience sample of 240 deceased individuals referred to the SPAS, and whose next of kin (NOK) agrees to participate will be included in this study.

Inclusion criterion:
A. Individuals aged 30 years or more.

Exclusion criteria (Figure 1):
A. The NOK of the deceased provides inconsistent data during the clinical interview.
B. The NOK of the deceased does not have at least weekly contact with the deceased.
C. There are visible evidence of factors modifying anthropometric measures, such as ascites, severe abdominal distension and pregnancy.
D. Loss of 10% or more of usual weight during the 6 months prior to death.
E. Arteries or visceral fat of the individuals are retained at the autopsy by the pathologist.
F. Postmortem interval (PMI) >24 hours.
G. Signs of body autolysis, according to the Crossley criteria.

Outcome measures
The primary outcome of interest will be atherosclerosis that is measured by the stenosis index in coronary, carotid and cerebral arteries and by the degree of atherosclerosis in the aorta. The secondary outcome will be the number of atherosclerotic plaques in cerebral and coronary arteries. All outcomes will be measured after autopsy.

Data collection
The NOK of the deceased will answer a semistructured clinical interview that will be administered by trained nurses concerning sociodemographics (age, gender, race, marital status, schooling and socioeconomic status (SES)), frequency of his/her previous contact with the deceased (weekly or daily) and cardiovascular risk factors (hypertension, diabetes mellitus, coronary artery disease (CAD), heart failure, dyslipidaemia, stroke, smoking, alcohol use and physical inactivity). Schooling will be quantified in years of formal education. SES will be evaluated using a Brazilian Scale that considers the quantity of household goods and the education level of the household owner, classifying in eight ordinal categories. These categories will be grouped to represent the upper, medium and lower social classes. Smoking and alcohol use will be classified in never, current or former. Physical inactivity will be defined as the individual who does not perform any physical activity. Frequency of physical activity per week will be also determined. The main cause of death and the PMI will be obtained from the autopsy report.

Anthropometric and visceral fat measurements
Anthropometric and visceral fat measurements will be obtained by trained persons following standardised procedures.
Weight will be measured in kilogram (kg) without clothes in the supine position using a calibrated electronic scale. Height will be measured in centimetre (cm) in the supine position using a stadiometer. After obtaining these measurements, the body mass index (BMI) will be calculated by dividing the weight (in kg) by the square of the height (in metres). Neck circumference (NC) will be measured in the region above the cricoid cartilage. The waist circumference (WC) measurement will be performed in the region of the umbilicus, and the hip circumference (HC) will be performed in the largest area on the great trochanters. The abdominal subcutaneous tissue thickness (ASTT) will be measured at the abdominal midline, 4 cm above the umbilicus. All measurements will be in cm, using an inelastic tape.

Individuals with a similar BMI may have different fat distributions. Thus, the waist-to-hip ratio (WHR) has been used as an additional measure of adiposity. The WHR and the waist-to-height ratio (WHtR) will be calculated by dividing the WC by the HC and by the height, respectively.

A body shape index (ABSI) will be calculated as WC/(BMI^{2/3}×height^{1/2}) using WC and height in metres, as suggested by Krakauer and Krakauer who determined that ABSI was a predictor of mortality. In addition, ABSI has been known to have a stronger association with total and cardiovascular mortality compared with other anthropometric measures in older adults.

The heart and pericardial fat (which consists of epicardial and paracardial fat) will be washed in running water to remove clots, fixed in 70% alcohol for 1–14 days and weighed. Epicardial fat is located at the outer wall of the myocardium, and paracardial fat is located in the mediastinum outside of the parietal pericardium. Omental, mesenteric, mesocolon and perirenal fat will be dissected and weighed (figure 2). All visceral fat will be weighed using a calibrated electronic scale, and the measurements will be expressed in grams. Samples of visceral fat with dimensions of 0.5×1.5 cm

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<td>Post mortem interval greater than or equal to 24 hours or cadaver in autolysis</td>
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![Figure 1](http://openheart.bmj.com/) Study outline.

will be frozen in liquid nitrogen and stored in the freezer at −80°C, and other samples with dimensions of 1.5×1.5 cm will be fixed in 10% formalin for future studies.

**Standardisation of pericardial fat fixation**

To evaluate the effect of different fixation times on pericardial fat weight, we collected pericardial fat from 20 individuals and we weighed the fat in natura and after fixation in 70% alcohol for 1, 3, 7 and 14 days. Bland-Altman plots were used to analyse the agreement between the weights of the fresh and fixed pericardial fat. We plotted the difference between the weights of the fresh and fixed pericardial fat against their means, and it was within the agreement limits (figure 3). We also calculated the correlation between the weights of the fresh and fixed pericardial fat, and they were high for all the different times (r>0.99).

**Evaluation of systemic atherosclerosis**

Carotid arteries will be dissected from the base of the aortic arch and stored in 70% alcohol for 24 hours. Gelatine will be injected inside the vessel lumen to prevent artery collapse, and the artery will be further fixed with 10% formalin. Subsequently, the common and internal carotid arteries will be cut transversely at 5 mm intervals, and five sections will be selected: (1) a section with the largest lumen obstruction of the common carotid artery, (2) the section of the common carotid artery located 1 cm below the carotid bifurcation, (3) the section located at the bifurcation, (4) the section of the internal carotid artery located 1 cm above the carotid bifurcation and (5) the section containing the largest lumen obstruction in the internal carotid artery. These cross-sections will be photographed with a stereomicroscope (Nikon SMZ 1000; Nikon Inst., New York, USA); the area will be delimited by the outer vessel wall.
and the lumen area will be measured using an image processor (Image J) (figure 4). For each segment, the stenosis index will be calculated by subtracting the lumen area from the outer area and dividing it by the outer area. The quotient will be then multiplied by 100.36

The circle of Willis (CW) will be removed from the base of the brain and processed similarly to the carotid arteries. Subsequently, the following arteries of the CW will be cut into 3 mm thick cross-sections: the anterior cerebral arteries, the anterior communicating artery, the middle cerebral arteries, the internal carotid arteries (segments close to the CW), the posterior communicating arteries, the posterior cerebral arteries and the basilar artery. The segment with the largest lumen obstruction from each artery will be photographed using the stereomicroscope. The stenosis index will be calculated similarly to the carotid artery.

The left, circumflex, left anterior descending and right coronary arteries will be dissected and processed similarly to the carotid and CW arteries. They will be cut into 5 mm thick sections. The stenosis index of the largest plaque will be also calculated as previously described. The presence of myocardial infarction will be carefully evaluated by a certified pathologist. Lesions will be classified in acute or chronic myocardial infarcts according to the criteria of Kumar et al.37 Left ventricular wall thickness will be measured 1 cm below the mitral valve in mm.

The number of atherosclerotic plaques in cerebral and coronary arteries will be counted to describe the extent of atherosclerotic disease.

We plan to evaluate atheroma plaque characteristics from the coronary, carotid and cerebral arteries. These samples will be submitted to the process of dehydration, diaphanisation and paraffin embedding. Then, the paraffin block will be cut into 4 µm thickness sections using a microtome (Leica RM 2145; Leica Microsystems Brazil). The slides will be stained using H&E and Verhoeff and photographed using a stereomicroscope. The evaluation of the plaque composition will be performed according to the American Heart Association criteria and classified in types I–VIII.38

The aorta (ascending, thoracic and abdominal) will be opened longitudinally. The degree of atherosclerosis and the presence of confluent lesions (those that extend around the circumference of the aorta) will be investigated. Plaques will be classified as grade 1 if the plaques are not confluent and there is no ulcerations and protrusions, grade 2 if there are confluent areas or/and an area of ulceration with minimal protrusion and grade 3 if there are confluent plaques, multifocal ulcerations or protrusions39 (figure 5).

Sample size calculation

The sample size was calculated on the basis of previous studies using the lowest correlation coefficient found in the literature from the study of Kortelainen and Särkioja16 (r=0.109 for the correlation of WC and perirenal fat). Assuming a power of 90% and α=0.05, in a two-tailed test, we will need a sample of 205 individuals. To better study different age groups, we will include at least 240 individuals in our sample.

DATA ANALYSIS

To describe the sample, we will use mean and SD or median and IQR for continuous variables and absolute
and relative frequencies for categorical variables. If the distribution is Gaussian, an unpaired t-test will be used to compare the anthropometric measures (BMI, NC, WC, HC, WHR, WHtR, ABSI and ASTT) and the weight of visceral fat (thoracic, abdominal and total) between genders; otherwise we will use the Mann–Whitney test. We will use one-way ANOVA to compare these measurements among age groups (30–49; 50–69; 70–89 and 90 years or more) and types of myocardial infarction (control, acute, and chronic) if the distribution is Gaussian, otherwise we will use the Kruskal-Wallis test. We will use Tukey’s HSD (honest significant difference) test in post hoc analyses to determine which mean value is significantly different among age groups and types of myocardial infarction.

We will use Pearson’s correlation test (for continuous variables with Gaussian distribution) or Spearman’s rank correlation coefficient (for continuous distribution with non-Gaussian distributed or ordinal variables) to compute the correlation of:

A. Weight of visceral fat with stenosis index in coronary, cerebral and carotid arteries.
B. Weight of visceral fat with degree of atherosclerosis in aorta.
C. Weight of visceral fat with number of plaques in coronary and cerebral arteries.
D. Anthropometric measurements with stenosis index in coronary, cerebral and carotid arteries.
E. Anthropometric measurements with degree of atherosclerosis in aorta.
F. Anthropometric measurements with number of plaques in coronary and cerebral arteries.
G. Weight of the heart and left ventricular wall thickness with the stenosis index in coronary arteries.
H. Weight of the heart and left ventricular wall thickness with number of plaques in coronary arteries.

Linear regression analysis will be used to assess the association of visceral fat, anthropometric measurements, sociodemographic data and cardiovascular risk factors with atherosclerosis in coronary, carotid and cerebral arteries and will use ordered logistic regression analysis to assess the association of visceral fat, anthropometric measurements, sociodemographic data and cardiovascular risk factors with the degree of atherosclerosis in aorta. Results will be adjusted for age, gender, schooling, smoking and physical inactivity; and the weight of visceral fat will be adjusted for height. The significance level will be set at 0.05 in two-tailed tests. All analyses will be performed using the Stata/MP V.14 (StataCorp LP, College Station, Texas, USA).

STRENGTHS AND LIMITATIONS OF THE STUDY
The main limitation of this study is the absence of clinical follow-up while the participant was alive. However, previous studies performed by our group demonstrated good reliability of our interview compared with a gold-standard outpatient evaluation. In addition, the main variables in this study (e.g., atherosclerosis, visceral fat and anthropometric measurements) will be directly measured and will not be affected by a lack of follow-up during the life of the participant. Another limitation is the convenience sample, since SPAS performs only autopsies on individuals who died of unknown natural causes of death; therefore, the sample will not be representative of all deaths in the city of Sao Paulo.

On the other hand, the structure provided by SPAS will allow us to get a large sample due to acceptance of NOK regarding tissue donation and participation in the study, good conditions of the deceased’s tissue and reliable information provided by the NOK. Furthermore, this study has several strengths. The most important one will be the large number of participants. In comparison, most previous autopsy studies evaluated a lower number of cases, ranging from 30 to 50 individuals, which is probably due to the decreasing number of autopsies performed worldwide. Another strength is the evaluation of older individuals while previous studies included a larger percentage of younger patients (Kortelainen and Särkioja: mean age 41 years (range: 19–63); Kortelainen and Särkioja: 35 years (range: 19–49) and Kortelainen and Särkioja: 17 years (range: 13–19)). The short PMI mean of 10.4 hours (4–20 hours) of the material collected by the BB-BABSG team allows for high-quality material for future genetic and proteomic studies. Moreover, our sample will be ethnically diverse, unlike previous autopsy studies for which the samples were mostly Caucasian. The present study also will perform morphometric measures of atherosclerotic plaques and objective measurements of visceral fat that are the gold standard compared with other clinical and imaging techniques. In fact, Kortelainen and Särkioja reported that the autopsy studies have the advantage of being able to measure the whole deposit of visceral fat and allows for complete assessment of the coronary tree. Additionally, our study will evaluate atherosclerosis not only in coronary arteries but also in the aorta, cerebral and carotid arteries, which is a unique aspect compared with other studies. Furthermore, most of the previous autopsy studies evaluated only abdominal visceral fat, while we also will evaluate pericardial fat.

Finally, we will use a unique autopsy service that already supported a variety of pathological projects to gather a high-quality and diverse tissue collection of arteries and visceral fat. This service also provides a comprehensive postmortem clinical evaluation. A large number of samples can be collected due to the great number of autopsies performed in our service. The stored fat and arterial samples will be an outstanding opportunity for future collaborative studies that will advance the understanding of the relationship between atherosclerosis pathophysiology and adiposity.

DISSEMINATION
After being informed about this study, the NOK of the deceased will sign a written informed consent. Results
Coronary artery disease

will be submitted for publication in a peer-reviewed journal and also presented at national and international conferences.

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Contributors The International Committee of Medical Journal Editors (ICMJE) criteria for authorship have been met. AN, CKS and CAP designed the study; AN, DSF, FMC, KCSS and AC were responsible for collecting and processing data. CKS, REPL, RELF, LTG, JMF, WJF and CAP were responsible for general supervision of the research group and support in acquisition and analysis of data. AN drafted the manuscript. AN and CKS performed the statistical analysis. All authors read, revised the manuscript critically, making significant improvements, and approved the final version to be published.

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Competing interests None declared.

Patient consent Obtained.

Ethics approval The study protocol was approved by the Ethics Committee of University of Sao Paulo Medical School (412/11), Sao Paulo, Brazil.

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REFERENCES


