Sudden cardiac death in a woman with clinical evidence of coronary vasospastic disease and myocardial stunning. A necropsy insight into the etiology

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Abstract
This case shed some light on the etiology of severe coronary vasospastic disease. A 54 years old lady with metabolic syndrome had a terminal ventricular fibrillation after an episode of acute ischemia caused by a coronary spasm. At the autopsy study an inflammatory perivascular and mainly perineural infiltration of the autonomic fibres of the cardiac nerves, confirmed at immunohistochemistry, could suggest a potential susceptibility of the nervous fibres to evoke an intense coronary vasospastic response as a consequence of non-specific events.

Background
The etiology of vasospatic angina is not well known [1]. Coronary endodothelium and autonomic nerves dysfunction have been indicated as a possible mechanism of coronary vasospasm but these studies have had little clinical impact as no causes of this dysfunction if any, has been demonstrated [2-7].

Objective
We presented a case of clinical vasospatic angina associated with myocardial stunning, unfortunately ended with the sudden cardiac death of the patient whose subsequent necropsy examination revealed inflammatory infiltrate of the autonomic cardiac nerve, thus identifying a probable etiologic factor of the coronary vasospastic disease.

Methods and findings
A 52 year old woman was admitted to the emergency department 3 hours after the onset of chest pain. The patient was a mild obese black woman with a history of dyslipidemia, hypertension, type 2 diabetes mellitus, hiatal hernia and recurrent chest pain. A coronary angiography performed 2 years earlier had demonstrated the absence of significant coronary stenosis. The physical was unremarkable. The first electrocardiogram showed left bundle branch block, already present at a previous evaluation, but with deep negative T waves (>5 mm) discordant to QRS associated with mild ST elevation (3 mm) in leads V2-V3. Troponine I peak was at 0.43 ng/ml. The echocardiography demonstrated a left ventricle (LV) with a mid-septal akinesia and apical dyskinesia and with an ejection fraction of 50%. Such LV diskinesia completely reverted to normal 1 week after the recovery. The angiography confirmed the 2 years earlier angiographic findings. During the recovery the patient complained recurrent chest pain and such episodes were scarcely ameliorated by calcium channel blockers and nitrates. About 16 hours after the discharge, early in the morning at the awakening, as reported by relatives, the patient developed intense chest pain terminating after 20 minutes with a rapid resolution followed by syncope due to irreversible cardiac arrest. Autopsy was performed to clarify the cause of death.

Holter monitoring during the lethal syncope. In the first 16 hours the recording was unremarkable. At 4:08 a.m. the ST segment rose to a peak of 9 mm above the isoelectric line within 12 minutes and remained so elevated for a further 13 minutes, then dropped to 2 mm above the isoelectric line within 2 minutes. Repetitive ventricular ectopic beats started, leading immediately to ventricular fibrillation.

Autopsy findings. At autopsy, the heart (330 g of weight) showed a mild left ventricular hypertrophy and a marked but non occlusive coronarosclerosis. Histology showed myocardial signs of a recent acute ischemia and ventricular fibrillation and at the level epicardial layer of the heart a mild infiltration composed of mononuclear cells with a perivascular and perineural pattern of distribution. (Figure 1) These cells resulted positive to the immunostain for T lymphocyte marker CD 3. No expression for B lymphocyte and macrophage antigens (CD 20 and CD 68 respectively) were observed.

Discussion
The clinical history clearly depicts a malignant coronary vasospastic disease and the post mortem histology identifies an inflammatory perivascular and mainly perineural infiltration of the autonomic fibres of the cardiac nerves confirmed at immunohistochemistry that could suggest a potential susceptibility of the nervous fibres to evoke an intense coronary vasospastic response as a consequence of non-specific events. Several observations suggests that the autonomic nerves dysfunction is important in the development of coronary vasospasm [2-7] but they do not give any idea on why such an autonomic imbalance, if any, does occur.

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Our study demonstrated for the first time a concrete histopathologic alteration (perineural inflammation of the cardiac nerves) that may represent the pathologic substrate subtended to an autonomic imbalance potentially inducing a coronary vasospasm. This finding concurs and corroborate in terms of histology other studies of the literature that document low grade chronic inflammation by means of humoral markers (P-Selection adhesion molecules, elevated HESR) in patients with coronary spasm. As far as we know, all the histopathological studies reported in the literature regarding patients with coronary vasospastic disease, were performed on endomyocardial biopsies and documented a scant interstitial lympho-histiocite infiltration, mild fibrosis and occasional contraction band. Endothelial dysfunction associated or not to a frank obstructive plaque (in our case there was a subaclusive plaque in the left anterior descending coronary artery) has a common occurrence and does not cause clinically significant spasm, thus other factors are concomitantly needed to cause coronary spasm.

Why inflammation of the autonomic nerve can occur is unknown. However our patient was mild obese and diabetic. This metabolic condition favors silent inflammation in the tissue thanks mainly to hyper-insulinemia [8,9] triggered to insulin resistance. Thus it can be hypothesized that a control of both obesity and hyperinsulinemia through life style changes [10] may have a curative impact in the long term on the vasospastic disease by reducing silent chronic inflammation. Further studies are needed to test this hypothesis in patients with coronary vasospasm.

References