Hepatocyte metaplasia in experimental chagasic pancreatitis: preliminary report

Metaplasia hepatocítica em pancreatite chagásica experimental: nota prévia

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Abstract: Beginning the study of chronic pathologic changes in pancreas of hamsters experimentally infected with Trypanosoma cruzi Vic strain, hepatocyte metaplasia was observed in one animal from infected group. This is the first report of oncocyes in Chagas’ disease, which could be due to aberrant regenerative response to pancreas inflammatory process.


Resumo: Iniciando estudo de alterações patológicas crônicas no pâncreas de hamsters experimentalmente infectados com a cepa Vic de Trypanosoma cruzi, metaplasia oncocítica foi observada em um dos animais infectados. Este é o primeiro relato de oncocitos na doença de Chagas, que poderiam decorrer de resposta regenerativa aberrante ao processo inflamatório pancreático.


Hamsters infected with Trypanosoma cruzi may exhibit histopathological changes similar to human Chagas’ disease, including pancreatitis. Following injury and regeneration, pancreas ductular cells may give rise to metaplastic hepatocyte-like cells. Oxyphilic, oncocytic, ductular, hepatocytic or hepatocyte-like metaplasia has been described in older people, in human chronic pancreatitis and diabetes mellitus, and in spontaneous or experimentally induced tumors.

The complete study was planned in order to include the acute (first 60 days after the initial infection) and the chronic phase of experimental Chagas’ disease up to 375 days. Just at the beginning of chronic pathologic and immunohistochemical evaluation in pancreas of hamsters experimentally infected with T. cruzi, oxyphilic metaplasia was observed in one of the infected animals. This is the first report of pancreas hepatocyte metaplasia in Chagas’ disease.

Experimental procedures were in accordance with the Principles of Laboratory Animal Care from the National Society of Medical Search, and the Guide for the Care and use of Laboratory Animals from the National Institute of Health. The animals were housed ten per rigid plastic cage, with hardwood chip bedding and a 12-hr light/dark cycle. A standard industrial pelleted diet (54% carbohydrate, 23% protein and 4% fat) supplemented with minerals, vitamins and amino acids, and tap water given ad libitum.

Ten young male non-isogenic Syrian golden hamsters, which had been infected, and reinfeeted 75 days after, by intraperitoneal route, with 2,000 trypomastigote forms of the T. cruzi Vic strain were sacrificed on the 105th day after initial infection. Ten weight and age-matched non-infected male hamsters constituted the normal control group studied.

The sacrifice and careful necropsy schedule included the 15th, 30th, 45th, 60th, 105th, 180th, 255th, 330th and 375th day, and the reinfections were scheduled for the 75th, 150th, 225th, and 300th days of experiment. Fragments of pancreas were collected, and buffered formalin-fixed, paraffin-embedded, 5 m sections were stained with hematoxylin-eosin and also submitted to immunohistochemical procedures. For immunohistochemical studies, the avidin-
biotin-peroxidase method, antibodies anti-\textit{T. cruzi} (1:3,000) and the islet cell specific antibodies: insulin (1:500), anti-glucagon (1:600), anti-somatostatin (1: 3,500), and anti-pancreatic polypeptide (1: 15,000) were employed. Statistical analysis was planned to include parametric and nonparametric methods.

At the 105th day of experiment, major evidence of parenchyma hypotrophy, fibrosis and inflammatory infiltrate, characterizing pancreatitis, were invariably found in the infected group, while only minor changes were observed among the animals from control group.

Hepatocyte metaplasia was seen in a peri-insular zone of an infected animal (Figure 1), while the phenomenon was not observed in normal control hamsters. In both groups, normal islet-cells were positive to insulin, glucagon, somatostatin and pancreatic polypeptide antibodies, while oncocytes were all negative (Figure 2). The PAP-anti-\textit{Trypanosoma cruzi} test was also positive in the pancreas sample where oncocytes were observed (Figure 3).

Hamster chronic pancreatitis, due to chagasic infection, was characterized by reduction in number and

Figure 1 - Parenchyma hypotrophy and lymphocytic infiltrate diffusely distributed, including the islet. Peri-insular changes characterizing hepatocyte metaplasia (arrows) are observed (HE, X100).

Figure 2 - On immunohistochemistry, normal islet-cells are positive to insulin (A), glucagon (B), somatostatin (C) and pancreatic polypeptide (D) antibodies, while oncocytes are negative (X400).
size of acini and chronic inflammatory infiltrate around pancreas lobules and ducts, in addition to PAP anti-T. cruzi positive test in the same pancreas section.

The absence of oncocytes in animals of the non-infected group could suggest that the pancreas hepatocyte metaplasia is related with an aberrant response to the inflammatory process in Chagas' disease. Nevertheless, at the 105th day of experiment, the number of animals already studied (n=60) was not large enough to enable such a conclusion. Moreover, since oncocyctic proliferation may become a more common event in association with animal aging11 and/or with chagasic reinfection, the remainder (n=155) normal control as well as infected hamsters will be appropriately evaluated after a longer period - up to 375th day post-inoculation - as previously scheduled.

Only for the purpose of this preliminary report, the oncycotic nature of pancreas changes was based on the presence of large mitochondria, in addition to abundant, almost homogeneous, and finely granular eosinophilic cytoplasm observed under light microscopy8 12. However, for more convincing evidence of the phenomenon, further evaluation of the oncocytes will include liver-specific markers and ultrastructural details.

The occurrence of pancreas hepatocyte metaplasia in Chagas' disease was not previously described. Although the authors consider it inadequate to extrapolate this very preliminary finding to human cases, the concern is an eventual relationship between Chagas' disease and pancreas oncocytes.

REFERENCES