Schistosoma mansoni granuloma in late evolutive phase, in a case of tumoral form in man


ABSTRACT

Introduction: Authors describe human schistosomal granuloma in late chronic phase, from the morphological and evolutionary viewpoints. Methods: The study was based on a histological analysis of two fragments obtained from a surgical biopsy of peritoneum and large intestine of a 42-year-old patient, with a pseudotumoral form mimicking a peritoneal carcinomatosis associated to the schistosomiasis hepatointestinal form. Results: Two hundred and three granulomas were identified in the pseudotumor and 27 in the intestinal biopsy, with similar morphological features, most in the late chronic phase, in fibrotic healing. A new structural classification was suggested for granulomas: zone 1 (internal), 2 (intermediate) and 3 (external). Conclusions: Regarding granuloma as a whole, we may conclude that fibrosis is likely to be controlled by different and independent mechanisms in the three zones of the granuloma. Lamellar fibrosis in zone 3 seems to be controlled by matrix mesenchymal cells (fibroblasts and myoepithelial cells) and by inflammatory exudate cells (lymphocytes, plasmocytes, neutrophils, eosinophils). Annular fibrosis in zone 2, comprising a dense fibrous connective tissue, with few cells in the advanced phase, would be controlled by epithelioid cells involving zone 1 in recent granulomas. In zone 1, replacing periovular necrosis, an initially loose and trabecular connective neoformation, housing stellate cells (fibroblasts and myoepithelial cells) and by inflammatory exudate cells (lymphocytes, plasmocytes, neutrophils, eosinophils). Accessory cells in zone 2 may be the main mediator for granulomatous response and for immune modulation. Fibroblasts and eosinophil granulocytes may, in certain cases, prevail over the other cells, so that granulomas arise almost exclusively eosinophilic as it is seen during the early acute phase of schistosomiasis toxemic form. In other cases, neutrophil granulocytes may prevail, developing an abscess (more common in paracoccidioidomycosis; rare in schistosomiasis); exceptionally mast cells and basophils may also contribute as accessory cells. Fibroblasts embedded are constantly found in an extracellular matrix composed of fbronectin, laminin, glycosaminoglycans and collagen.

INTRODUCTION

Granulomas are nodular formations in which exudate cells are displayed in particular arrangements to the inflammatory focus, forming structures with peculiar aspects and architecture which frequently allow to diagnose the disease even without finding its causal agent. Macrophages and their derivatives (remarkably epithelioid and giants cells) and lymphoid cells and their derivatives (particularly plasma cells) comprise the required cells. Accessory cells are composed of eosinophil granulocytes which may, in certain cases, prevail over the other cells, so that granulomas arise almost exclusively eosinophilic as it is seen during the early acute phase of schistosomiasis toxemic form. In other cases, neutrophil granulocytes may prevail, developing an abscess (more common in paracoccidioidomycosis; rare in schistosomiasis); exceptionally mast cells and basophils may also contribute as accessory cells. Fibroblasts embedded are constantly found in an extracellular matrix composed of fbronectin, laminin, glycosaminoglycans and collagen.

Granulomatous reaction in schistosomiasis is a late T-dependent hypersensitivity response to soluble egg antigens (SEA) released by a mature miracidium (this becomes mature in six to eight days after egg deposition), which induce a marked Th2 protective immune response, which guides the granulomatous reaction. Recent studies have indicated that IL-13 receptor α2 (IL-13 R α2) secreted by TCD4+ cells type 2 is the main mediator for granulomatous response and for hepatic fibrosis, which are two fundamental lesions in schistosomiasis.

METHODS

This study was conducted in two fragments obtained from a surgical biopsy of lesions from the large intestines and from the peritoneum (B-12.369/03), of a 42-year-old male patient (E.F.S), admitted at Hospital da Criança São José, Contagem/MG with a tumor form of schistosomiasis mansoni, with clinical signs and symptoms of intestinal obstruction, mimicking neoplasms. Fragments were whitish, irregular, with a firm consistency and measured 4.0 x 2.5 x 1.5cm (peritoneum) and 1.0 x 0.8 x 0.4cm (large intestines), respectively. After fixed in formaldehyde (10%), they were processed and included in paraffin blocks. Blocks were cut at 4 micrometers which were stained by hematoxylin and eosin (HE), Gomori trichrome and by picrosirius red, according to the technique recommended by Junqueira et al.11. A yellowish fragment of a hepatic biopsy, measuring 0.2 x 0.4cm, was examined.
RESULTS

At the late evolutive phase, a large number of granulomas has three distinct and morphologically different zones: I) zone 1, central; II) zone 2, intermediate; III) zone 3, peripheral, which modifies, due to the time of evolution.

Central zone (zone 1)

In early phases, it is generally the longest and the less stained both by HE and by special staining such as Gomori trichrome and picrosirius. It frequently houses the embryonated egg or its shell, whether or not surrounded by an eosinophilic amorphous area of necrosis, and by a thin fibrillar substance, mixed with few starry cells resembling fibroblasts or by cells with fusiform nuclei and macrophages (Figures 1A and 1B). This amorphous eosinophilic material, also seen in the early evolutive phase of the acute phase of schistosomiasis (necrotic-exudative granulomas) around the egg, known as Hoeppli phenomenon, contains egg antigens, host immunoglobulins and fibronectin (non-collagen glycoprotein) with a high molecular weight produced by fibroblasts and by other mesenchymal cells. In this phase, zones 2 and 3 are already distinct, as it will be discussed later. In granulomas with no eggs, the necrosis zone, responsible for the Hoeppli phenomenon is not identified. The same occurs when they are centered by the empty shell or egg debris.

In older granulomas (posterior phase to that previously described), they enter the productive phase. In this phase, the necrosis zone tends to disappear and the egg or its shell is partially or completely involved by one or more Langhans giant cells or foreign body giant cells (Figures 1C and 1D). This aspect also became evident during the acute phase in granulomas with 70 to 90 days of evolution. When stained with picrosirius, thin conjunctive fibers, stained in green, start to appear intermingling fibroblasts and histiocytes, indicating a type-III collagen. In the third phase, zone 1, the central one, is replaced by a homogeneous little cell mass, stained in dark green by Gomori, semicompact with empty spaces or completely compact. Therefore, centrifugal fibrosis occurred from inside to outside. Few or no fibroblasts and histiocytes are seen replaced by a dense fibrous nodule (Figure 1E). These results seem to confirm findings described by Al Adnani of decreased fibronectin, due to decrease or absence of cells which are able to produce it (macrophages and fibroblasts).

Intermediate zone (zone 2)

Its thickness is variable and it is present and is part of acute-phase granulomas in the acute phase of schistosomiasis. In the same initial period, between 45 and 78 days after infection, it appears as a crown completely involving zone 1. In this phase it is mainly composed of epithelioid histiocytes. As time passes, cellularity is reduced and gives rise to an acidophilus zone to HE and greenish stain by Gomori. It tends to be a layer composed of collagen fibers, with a variable thickness, sometimes homogeneous or composed of round and parallel fibers type-3 prevailing over those of type-1 collagen (Figures 1A, 1B, 1F and 1G). Based on the early constitution of epithelioid histiocytes, the resulting collagen is thought to be produced mainly by these cells. These morphological data are in accordance with the findings of Linder et al., i.e., the absence of fibronectin from the reduction or the absence of cells (macrophages and fibroblasts) producing this glycoprotein. We believe this range, which is initially cellular and eventually practically fibrous and acellular, might play an important role as a barrier to the SEA diffusion through egg pores to limit the lesion.

External zone (zone 3)

According to our experience with acute toxemic form in humans, this zone is well delineated mainly in granulomas with over 120-150...
days of evolution (granulomas in fibrotic healing phase). It is mainly formed by type-I collagen fibers and a smaller amount of type-III collagen, well evidentiated by picrosirius and by Gomori trichrome stainings, forming thicker fibers, parallel and round with a few overlapped cells interwoven with twines, resembling druses in onion scales, externally involving zone 2 (Figures 1A, 1B, 1F and 1G). In older and smaller granulomas when zone 2 tends to vanish or disappear, it may directly involve zone 1 (Figures 1E and 1H). Sometimes, when there is a confluence of granulomas, it starts to form a single mantle involving or limiting two or more granulomatous formations. This external zone (zone 3) is thought to be produced by mesenchymal cells (fibroblasts, myofibroblasts, histiocytes, plasmocytes and eosinophils) present and abundant in recent granulomas (necrotic-exudative granulomas and exudative granulomas). Therefore, it has a centripetal formation (from outside to inside).

**Fibrotic granulomas**

Since the disease evolves to the chronic form, as a result of the modulation, granulomas tend to slowly reduce its size, until they become a small fibrotic nodule, consisting almost exclusively of collagen fibers, paucicellular or practically acellular, similar to a scar (Figure 1C). This aspect is seen in lesions with years of evolution. The transformation occurs by replacing zone 1 with an amorphous acidiphilic mass, compact and more stained, which lies in the center of the lesion (Figures 1E and 1G), followed by the disappearance of zone 2, previously well-delineated and by the withdrawal of zone 3. It eventually ends up as a lesion either regular or not, consisting exclusively of collagen fibers, predominantly of type 1, when fibronectin is no longer detected.

Granulomas of the large intestine have the same characteristics described in pseudotumoral form.

Histological analysis of hepatic biopsy has shown a granuloma in fibrotic healing phase, with the external zone delineated by parallel and circular connective fibers like druses in onion scales, as a result of modulation, and a discreet inflammatory infiltrate of mononuclear cells in portal tracts, regardless the granulomatous lesion. Portal fibrosis caused by *Schistosoma mansoni* eggs was also discreet. The Symmers-Bogliolo form of disease was discarded. Hepatocytes have shown a marked and diffuse non-alcoholic steatosis.

**DISCUSSION**

Examining a large number of granulomas in the peritoneum (around 203) and in the serosa and external muscular tissue of the large intestine (27), in the same individual, E.F.C., 42 years old, from Belo Horizonte (MG), with a tumor or pseudotumoral form of schistosomiasis mansoni, provided a favorable view of the morphology of granulomas in their several evolutive phases in humans.

Both in the peritoneum and in the large intestines (LI), granulomas were in several evolutive phases (recent, old and scarred), which indicates an active long-lasting disease, probably lasting for years and modulated, with egg deposition at different stages. Although worms or their remains were not identified in the sections examined, eggs are believed to have been deposited *in situ*, both in the peritoneum and in the LI. In this case, we would have to assume that single and mated females would have gone from their natural habitat to the peritoneum, through aberrant venous branches or connections between mesenteric and peritoneal branches. Massive and ongoing embolization of several eggs is unlikely.

Morphological studies conducted, following the evolution of hepatic granulomas since their formation until long-lasting chronic forms, in patients who developed the chronic benign form, asymptomatic or oligosymptomatic, authors recommended classifying them in four phases: 1) granulomas in the necrotic-exudative phase; 2) in the exudative phase; 3) in the productive phase; and 4) in the fibrotic healing phase. Raso et al., concluded that, in humans, as well as in mice, acute forms of granulomas are significantly larger than those chronic forms. According to Andrade & Warren, and corroborated by other authors, it is correct to say that a decreased volume of granulomas is seen during the chronic phase of schistosomiasis.

Volume reduction and changes in histological aspects of granulomas start to show up during the acute phase of the disease. Between 43 and 70–78 days after infection, granulomas were shown to be in the necrotic-exudative phase, being large, coetaneous, with a generally extensive necrosis zone around an egg, and more externally a cellular exudation zone composed of macrophages, fibroblasts and lymphocytes. Granulomas with no necrosis zone are termed exudative granulomas and sometimes they are termed eosinophilic granulomas when polymorphonuclear eosinophils prevail. In addition, according to these authors, in this phase, granulomas may be found with three distinct zones: 1) central with necrosis around the egg; 2) intermediate predominantly composed of epithelioid histiocytes forming a crown around the necrosis zone; and 3) a more external zone comprising a mixture of cells, with the predominance of eosinophilic granulocytes. The following phase, which starts around the 70th - 78th day after infection, is when the granuloma undergoes changes in its size and constitution, and assumes the type of granulomas from the productive or fibrotic phase. This aspect between the productive phase and the fibrotic healing phase is thought to correspond to an intermediate or transitional phase between the chronic and the acute toxemic forms. After 9 to 18 months practically all granulomas were in the fibrotic healing phase, some of them were represented only by a scar.

In this work, recent granulomas (probably 30 to 40 days after egg deposition in the tissue) in the necrotic-exudative phase, similar to what was seen on the 78th day of evolution of the acute toxemic form, were in the necrosis zone (zone 1), sometimes vast, involving the embryonated egg, characterized by an amorphous eosinophilic material, known as Hoeppli phenomenon, interspersed by thin fibers and a few fibroblast-like cells. Necrosis zone was generally less extensive than that commonly seen in the acute toxemic phase up to the 78th day after infection. Some interpret it as the result of an Ag-Ab reaction, as it depends on egg specific antigens (soluble egg antigens) which are highly immunogenic due to the presence of carbohydrate epitopes, LDN, LDN-F and LDN-DF.

In time, zone 1, until then represented by necrosis, is gradually replaced by a fibrosis, originally loose and then dense, carried out by cells which are morphologically similar to fibroblasts. Therefore, they would account for centrifugal fibrosis, inside out, from the central zone which was previously represented by necrosis. Thin type I1 conjunctive fibers, stained in green by picrosirius, interspersing fibroblasts and histiocytes, at a later phase it is replaced by a homogeneous mass, little cellular, with few fibroblasts and histiocytes and finally by a compact
dense fibrous nodule. These results seem to confirm what Al Adnani\textsuperscript{13} reported about the decrease in fibronectin due to the reduction or absence of cells able to produce it (macrophages and fibroblasts)\textsuperscript{15}.

Around zone I of necrosis, a round wall is formed eminently cellular at first, composed mainly of epithelioid macrophages (zone II) which acts as a barrier trying to prevent the expansion of antigens secreted by the egg. These macrophages, at least in the beginning of the granuloma evolution (by around 70-78 days of acute form of infection), are epithelioid and tend to lie in palisade arrangement around zone I, forming a round mantle, which acts as a wall, preventing the egg antigen diffusion. About the 78th day of infection (in the acute phase), as it also occurs in the chronic phase, some fuse to form giant Langhans or foreign body cells formation which try to involve and/or comprise the egg, sometimes reduced to a wizened shell, with no miracidium and no power of secretion.

In time, the wall, which was prominently cellular, is replaced by a practically acellular compact formation, comprising predominantly type I collagen fibers, with a variable thickness, composed of round and parallel fibers. This collagen is thought to be produced in this intermediate zone 2, by epithelioid macrophages. These morphological data explain findings of Linder \textit{et al}.,\textsuperscript{15} i.e., the absence of fibronectin from the decrease or absence of cells (macrophages and fibroblasts) producing this protein glycoprotein. We may consider that this range, initially cellular and then practically fibrous and acellular, continues to act as a barrier to the diffusion of SEA antigens, with the purpose of limiting the lesion, mainly in granulomas with more time of evolution.

According to our experience, the external zone (zone 3) is well delineated after 120-150 days of infection\textsuperscript{4}. It is mainly formed by type I collagen fibers and a smaller amount of collagen type 3, forming thicker fibers, which are parallel and round, with a few interposed and interwoven cells (connected by links similar to those of a chain) or resembling druses in onion scales, externally involving zone 2. In smaller and older granulomas when zone 2 tends to disappear, it may directly involve zone 1. According to our findings, zone 3 is produced by fibroblasts, with a possible participation of other cells characteristic from the conjunctive tissue or which migrate to the focus and which are present in recent granulomas.

In this case, the pseudotumoral form is associated with the hepatointestinal form of schistosomiasis. Furthermore, our results clearly demonstrate the modulation of immunopathological response and that the signs and symptoms of the pseudotumoral form of the disease disappeared with the chemotherapy (praziquantel).

Modulation of granulomatous response was extensively investigated in mice. In 1964, Andrade & Warren\textsuperscript{4} demonstrated a decrease in the size of granulomas formed around \textit{Schistosoma mansoni} eggs that occurs at the chronic phase of disease, as well as alterations in the cellular and non-cellular constituents which take part in this inflammatory process. This phenomenon in the decreased granuloma size observed in late infection has been called \textit{modulation or immunological modulation}. The phenomenon seems to be dependent of cell-mediated immunity and has been corroborated by several other authors, as it can be seen in Boros' review\textsuperscript{37}.

Appearance and size of granulomas are very clearly described during transition from acute to chronic phases of human schistosomiasis in two papers: in the first, Raso & Neves\textsuperscript{4} reported a histopathological study on 36 hepatic biopsy specimens from patients with acute toxemic form of schistosomiasis mansoni, describing in detail the evolution of the schistosomal granuloma from 21 to 250 days. In the second paper, Raso et al.\textsuperscript{5} described and measured 286 granulomas from patients with acute schistosomiasis and 165 granulomas from patients in chronic phase. In their papers Raso et al.\textsuperscript{5} clearly described the phenomenon of granuloma immunomodulation in humans infected with \textit{Schistosoma mansoni}. Authors were able to define, in humans, the time of appearance, development and transformation from one to another of the four basic morphological types of granuloma (granuloma in necrotic-exudative phase, in the exudative phase, in the productive phase and in the cure by fibrosis). In 1976 Raso et al.\textsuperscript{5} were able to conclude that: a) granulomas show diverse development phases due to structural changes that occur during the evolution of the disease; b) granuloma size varies depending on the tissue or the organ where it is formed. It is larger in the liver, for example, than in the pancreas, intestines or testicles, etc.; c) there is a clear variation in size according to the stage of disease. They are much larger in the acute phase than in the chronic phase. As the disease progresses, there is an increasing tendency towards reduced volume; e) granuloma in humans, is much more developed than in experimental animals; f) granuloma falls under the influence of immunosuppressants.

There is, without doubt, replacement by connective tissue as time passes, until the transformation of granulomatous lesion to a dense, compact nodule in the liver and in other organs is complete. This replacement may go on for months or years.

Modulation of granulomatous response starts around 70-78 days after infection, and concludes, in some cases, within 150-day infection, when it is indistinguishable from other chronic forms\textsuperscript{5,5}.

An extremely important observation was provided by anatomo-pathological study of 12 cases of acute toxemic schistosomiasis, necropsied at the Department of Pathology of Faculty of Medicine, \textit{Universidade Federal de Minas Gerais}, in the 1950s-1970s by Bogliolo (4 cases) and Raso (8 cases). Granulomas of the liver, intestines, lungs, lymph nodes, etc. were maintained for a long time in the initial phase of development, in the necrotic-exudative phase. They were large granulomas with an extensive necrosis area around of the eggs, indicating an absence or a retardation of immunomodulation. This led one of us (Raso) to conclude that the cases with no modulation of granulomas evolved toward a fatal outcome. The same seems to have happened in the case recently described by Ferreira et al., 2012 of an acute form (Katayama syndrome) associated with Symmers form\textsuperscript{37}. Rather, the majority of patients who had serial biopsies immunomodulation of granulomas evolved as wells to healing, regardless of treatment administered.

In the acute form, an immunological imbalance is proved by changes in the cytokine profile\textsuperscript{19}.

On the other hand, a serial study on liver puncture-biopsies led us to conclude that in cases of modulation of granulomatous response, until the phase of cure by fibrosis, developed well: for one of the chronic forms or for a possible spontaneous cure, as it was observed in three cases, 13 years after the manifestations of acute toxemic form\textsuperscript{20}.

In the present case, we may state that there is no significant in the morphology of the granuloma of the peritoneum and of the wall of the colon. Modulation of the immune response occurred in both as well as it occurred in the liver.
As time goes time, both the human and in the experimental schistosomiasis have a decreased in the size of the granulomas and its replacement by fibrosis. In humans, fibrosis is predominantly consisted of type 1 collagen fibers; in mice, type 3 collagen fibers predominate. A total of 1,170 granulomas were measured on the 60th day after infection and were shown to have a larger size than those measured on the 90th day. Modulation of the immunological response was significantly more efficient for LE strain-infected mice than for those infected with SJ strain. The granulomas related to SJ were significantly larger (on the 60th and 90th days after infection)22.

Finally, in humans, unlike the claim Silva et al. in mice11, the immunomodulation occurs also outside of the liver (Figure 2).

FIGURE 2 - Granuloma in fibrotic healing phase, with external zone (Z3) delineated by parallel and circular connective fibers, stained by HE in liver (A), and stained by trichromic in large intestine (B) and from peritoneum (C) as a result of modulation, in the same individual, with pseudotumoral form of schistosomiasis mansoni.


