PRIMARY LOCALIZED RETROPERITONEAL AMYLOID TUMOR WITH OSSEOUS METAPLASIA: CASE REPORT AND REVIEW OF THE LITERATURE

T. GHIASI,* M.D., P. RAOFIAN,**M.D., AND S. NORUZPOUR,*** M.D.

From the Department of Pathology, Ghaem Hospital, Mashhad University of Medical Sciences, Mashhad, and the **Sina Hospital, Reza Avenue, Mashhad, Islamic Republic of Iran.

ABSTRACT

Amyloidoma or amyloid tumor is a tumor-like localized deposit of amyloid encountered occasionally in association with multiple myeloma, various chronic inflammatory diseases and primary amyloidosis.

Amyloid tumors unassociated with plasmacytoma or other disease are extremely rare in soft tissues and few cases in various situations have been reported.

Histologic examination discloses amorphous eosinophilic material that stains positive with congo red preparation, and shows apple-green birefringence under polarized light. The deposits are surrounded by more or less prominent chronic inflammatory cells, especially plasma cells and giant cells. Cartilage formation and ossification, however, occur in some amyloid tumors.

Keywords: Tumor, amyloid, retroperitoneal; metaplasia, osseous.


INTRODUCTION

Amyloidoma or amyloid tumor refers to localized tumor-like deposits of amyloid which are associated with multiple myeloma, various chronic inflammatory diseases and primary amyloidosis.

Amyloid tumors are extremely rare in soft tissues without concomitant plasmacytoma or other diseases as only a few cases have been reported in the breast, eye, thigh, mediastinum, central nervous system, urethra, bladder, lung, thyroid, rectum, larynx, trachea, stomach and palate.

Case report

The patient is a 44 year old woman who noticed a slightly painful rounded mass in her abdomen 7 years ago which had begun growing since last year. All blood examinations, serum protein electrophoresis and immunoelectrophoresis and bone radiographs were normal. Bence Jones protein was negative in urine. We did not find any other abnormal findings in her physical examination. The patient underwent laparotomy, and a retroperitoneal and intrapelvic mass was discovered which had adhesion to the iliac veins.

Microscopically, deposits of amorphous eosinophilic material are seen which stain positive with congo red and have green birefringence under polarized light. Chronic inflammatory cells, especially plasma cells, surround the deposits and rarely cartilage or osseous metaplasia are seen.

Correspondence: T. Ghiasi, M.D., Department of Pathology, Ghaem Hospital, Mashhad, IRAN, 91735 Fax: (051) 809612 Tel: (051) 800001-9

* Assistant Professor of Pathology
** General Surgeon
*** Resident of Pathology

369
Pathologic findings

Macroscopically the mass was 14×10×8 cm in diameter, of hard consistency in most areas and had a brown to white color in cut surfaces (Figs. 1, 2). Microscopic examination showed lymphocyte and plasmocyte infiltration with fibrosis and deposits of eosinophilic material (Figs. 3, 4) which were positive with congo red staining and had green birefringence under polarized light (Fig. 5) and showed areas of osseous metaphasia (Fig. 6).

DISCUSSION

Amyloid tumor is referred to as tumor-like localized deposits of amyloid material which is associated with multiple myeloma and other various chronic inflammatory diseases. Amyloid tumor is also associated with primary amyloidosis without any other disease which is rare and usually seen as solitary or multiple nodules in genitourinary, gastrointestinal and respiratory tracts. These nodules are usually large, and present with signs and symptoms of pressure which compress adjacent organs. Typical examples of amyloid tumor are seen in the palate, stomach, ureter, larynx, trachea and especially in the lungs.

Amyloid tumor without concomitant plasmacytoma or other disease is extremely rare in soft tissues and only a few cases have been reported. One case of this tumor was in the soft tissue of the neck of a 46 year old man which had presented as a large and lobulated mass deep to the sternomcleidomastoid and internal jugular vein for a period of 6 years. The other case was in the femoral triangle of an 84 year old woman, 6.5 cm in diameter and grey to light grey in cut surfaces. Serum proteins were within normal limits in both of these cases. Other cases have been reported in the literature.

Lipper and Kahn have reported 4 cases, one in bone and the others in the right breast of a 57 year old woman, the abdominal wall of a 47 year old man and the left inguinal region in a 40 year old man.

Follow up of two of the three cases showed nothing. In other reports, other organs, for example the breast, eye, thigh, mediastinum, brain, urethra (17 cases), bladder, lung, thyroid and the rectum have been involved.

Amyloid tumor has also been reported in the eyelid and skin as small nodules.

Brown Stein and Helwig in 1970 reported 39 cases of primary localized amyloid tumor in the skin and since then there have been several reports of these tumors in the skin. Some authors think that gonococcal urethritis is a predisposing factor for amyloid tumor of the urethra. One case of localized amyloid tumor in the thyroid has been reported which was associated with Hashimoto's thyroiditis.

Macroscopically the tumor consists of a mass or lobulated
metachromatic with crystal violet staining. They are positive with congo red staining and have apple-green birefringence under polarized light. These deposits are surrounded by multinucleated giant cells and more or less chronic inflammatory infiltrates rich in lymphocytes and especially plasma cells. Differentiation between the infiltration observed in amyloid tumor, which is often associated with enormous plasma cells, from the tumoral infiltration present in plasmacytoma is important. The following findings disclose the former as being a reactive phenomenon:

1) Small and enormous blood vessels which are covered by monomorphous but large endothelial cells.
2) Mature plasma cells
3) The presence of other inflammatory cells
4) Russell bodies
5) To prove the polyclonality by immunohistochemistry.

Deposits of this material are fine and fibrillar and have different amino acids in each case.

Some believe that the precursor protein in localized cases is made by plasma cells which are adjacent or in the deposits even if they are few in number. On the other hand, in systemic cases the precursor proteins exist in the circulation and their site of production may be far from the place in which they are deposited.

Immunohistochemical evaluation has been done in only a few cases, but in almost all of them this protein was a light chain monoclonal immunoglobulin.

Kappa chain was present in two patients with localized skin and larynx amyloidosis, and lambda chain was reported in two patients with localized skin and lung amyloidosis.

The kind of precursor protein in localized amyloid tumors of the skin, cervical vertebrae, wrist and other joints is beta 2 microglobulin, although a small area may be alpha, positive.

Vessel walls are often diffusely thickened by deposition of amyloid. Microscopically, there is no difference between the amyloid tumor present in primary or secondary amyloidosis.

Elastin stain helps to differentiate this tumor from elastofibroma and absence of calcium separates this tumor from tumoral calcinosis.

Cartilage and bone formation sometimes occurs in amyloid tumor.

REFERENCES

Retroperitoneal Amyloid Tumor with Osseous Metaplasia