Primary systemic amyloidosis, acquired cutis laxa and cutaneous mucinosis in a patient with multiple myeloma

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Abstract: A 57-year-old woman presented with periorbital ecchymoses, laxity in skin folds, polyneuropathy and bilateral carpal tunnel syndrome. A skin biopsy of the axillary lesion demonstrated fragmentation of elastic fibers, but with a negative von Kossa stain, consistent with cutis laxa. The diagnosis of primary systemic amyloidosis was made by the presence of amyloid material in the eyelid using histopathological techniques, besides this, the patient was also diagnosed with purpura, polyneuropathy, bilateral carpal tunnel syndrome and monoclonal gammopathy. She was diagnosed as suffering from multiple myeloma based on the finding of 40% plasma cells in the bone marrow, component M in the urine and anemia. The patient developed blisters with a clear content, confirmed as mucinosis by the histopathological exam. The final diagnoses were: primary systemic amyloidosis, acquired cutis laxa and mucinosis, all related to multiple myeloma.

Keywords: Amyloidosis; Cutis laxa; Mucinoses; Multiple myeloma

INTRODUCTION

Systemic amyloidosis may occur associated to clonal lymphoid diseases in which immunoglobulins are produced, including non-Hodgkin’s lymphoma, Waldenström’s macroglobulinemia and multiple myeloma (MM).

Acquired cutis laxa (CLA) is rare and may appear at any age. There are reports of its association with plasma cell dyscrasias, including MM.1 The link between CLA and MM remains uncharted, however, it is believed to be immune mediated.2 There is no synchronism between the courses of MM and CLA.

Mucinosis may also occur in cases of paraproteinemia, such as MM.

The case reported here features primary systemic amyloidosis, acquired cutis laxa and cutaneous mucinosis in a patient with MM.
CASE REPORT

A fifty-seven year old, female informed the presence of bilateral eyelid hyperchromia for the last fifteen years, followed by an increase in palpebral volume that started three years after the first symptoms. Six years ago, she observed cutaneous laxity in skinfold areas and bilateral palpebral ptosis. In the last four years, the patient reports pain and paresthesia in a stocking-like pattern affecting the lower limbs. The patient denied any familial history of cutaneous lesions.

She had a previous history of bilateral carpal tunnel syndrome and blepharoplasties performed in two different occasions, 2008 and 2009.

Dermatologic examination of eyelids showed hyperchromia, increase of local volume, cutaneous laxity and bilateral ptosis with periorbital ecchymoses (Figure 1). We observed cutaneous laxity, hyperchromic papules and comedones over the skinfolds, such as the armpits, cervical and inframammary regions (Figure 2). Biopsies of both axillary and palpebral lesions were performed.

Histopathological examination of the axillary lesion did not show any significant alteration with hematoxiline-eosine stain, detecting only one epidermic cyst, corresponding to the observed papule. Orcein staining showed the fragmentation of elastic fibers and von Kossa staining was negative, i.e., without evidence of calcium, thus establishing the diagnosis of acquired cutis laxa (Figure 3).

Histological exam of the palpebral material showed an amorphous eosinophilic substance, located on the superior and vascular dermis, positive for red Congo staining and with green birefringence to polarized light, which confirmed the diagnosis of amyloidosis (Figure 4).

Laboratory exams demonstrated normocytic and normochromic anemia, an inversion of the albumin-globulin ratio, IgG-Kappa monoclonal gammopathy, Bence-Jones proteinuria and 40% of plasma cells in the bone marrow specimen, prompting a diagnosis of multiple myeloma.

The patient evolved with citrine-content blisters on the inframammary region. A plaque, with severe edema, located on the left arm was biopsied, showing dermic edema with moderate fibroblast proliferation, some of which were stellate (Figures 5A and 5B). Colloidal iron staining determined the presence of mucin, and the diagnosis of papular mucinosis, which was not in agreement with the clinical presentation, thus making it difficult to classify the disease in this case (Figures 5C and 5D). A negative result for red Congo stain discarded bullous amyloidosis.

The final diagnoses were, accordingly, primary systemic amyloidosis, acquired cutis laxa and cutaneous mucinosis, all dermatologic manifestations associated to MM.

DISCUSSION

The diagnosis of primary systemic amyloidosis was made based on the following findings: presence of amyloid material on the perivascular dermis; purpura – including in the periorbital area, considered a marker for this disease; polyneuropathy; bilateral carpal tunnel syndrome; inversion of the albumin-
globulin ratio and IgG-Kappa monoclonal gammopathy. The patient was diagnosed with MM by presenting 40% of plasma cells in the bone marrow exam, urinary M component and anemia.

Carpal tunnel syndrome appears in 25% of cases of primary systemic amyloidosis, and it is often bilateral. The peripheral nervous system may also be affected. Both conditions were detected in our patient.

Mucocutaneous alterations are present in 20 to 40% of the cases, frequently as the first signs of amyloidosis. The following may also occur: purpura (spontaneous or secondary to minimal trauma), waxy papules, plaques or nodules (these are the most distinctive lesions) on skinfolds and affecting the central area of the face, and macroglossia. Purpuric lesions are the most frequent ones, and may affect all the body, however the characteristic lesion is the periorbital one (“raccoon eyes”). Those are believed to be secondary to amyloid deposits on the vascular walls, creating capillary frailty.

Multiple myeloma rarely affects the skin. There are, however, several diseases that have cutaneous manifestations and that may be associated to multiple myeloma (Chart 1).

Mucinosis was yet another condition found in the context of paraproteinemia. This case was considered as a form of the illness that could not be included in any of the defined forms of classification of mucinoses, because, histopathologically the patient had an intense deposit of mucin, with moderate pro-
**CHART 1:** Cutaneous diseases associated to multiple myeloma

| Specific cutaneous manifestations: |  
|---------------------------------|---|
| • Extramedullary cutaneous plasmacytoma |  
| Almost always associated: |  
| • Scleromyxedema |  
| • POEMS syndrome |  
| • Schnitzler’s syndrome |  
| • Necrobiotic xanthogranuloma |  
| Frequently associated: |  
| • Normolipemic plane xanthoma |  
| • Buschke’s Scleroderma |  
| • Acquired deficiency of C1 esterase inhibitor angioedema |  
| Significant association (≥15% of cases): |  
| • Primary systemic amyloidosis |  
| • Erythema elevatum diutinum |  
| • Subcorneal pustular dermatosis |  
| • IgA Pemphigus |  
| • Pyoderma gangrenosum |  
| Occasionally associated: |  
| • Acquired cutis laxa |  
| • Sweet Syndrome |  
| • Hypergamma globulinemic purpura of Waldenström |  
| • Disseminated xanthoma |  
| • Small vessel cutaneous vasculitis |  
| • Acquired Bullous Epidermolysis |  
| • Paraneoplastic pemphigus |  

lleration of fibroblasts and without an increase in collagen, which would lead us to the diagnosis of papular mucinosis. However, we observed, clinically, a plaque with severe edema on the arm and the presence of inframammary blisters, which were discordant with the histopathological classification. Therefore, we might classify this case as atypical cutaneous mucinosis. There are no cases reported on the literature of cutaneous mucinosis manifesting clinically as blisters. Until the present, there is only one published report of papular mucinosis associated to primary systemic amyloidosis in a patient with paraproteinemia.5

Acquired cutis laxa has been described in patients with MM. In several patients with CLA, the involvement of the face and neck occurs first, with a cephalocaudal progression, such as in the case reported here. Despite the probability of visceral involvement in CLA, causing pulmonary emphysema, gastric fibromas and tracheobronchial malacia, no other organ was affected in our patient.

A possible differential diagnosis, in this case, is pseudoxanthoma elasticum (PXE), one of our first hypotheses. It was rejected by the absence of calcium on von Kossa staining. Other findings were also contrary to this diagnosis, such as the palpebral involvement (there are no reports on the literature of PXE affecting the eyelids); and the absence of signs that other organs were affected, such as, retinal angioid streaks.

There are no treatments to prevent the progression of cutis laxa, although the normal cicatrization process of these patients allows for surgical corrections.4 Due to the progressive nature of this disease, many interventions are required over time. Two blepharoplasties were performed in our patient, with subsequent relapse of the palpebral ptosis.

First-line treatment for MM was started with Bortezomib and Dexamethasone, followed by autologous bone marrow transplantation, with clinically important dermatological improvement.

This case presents a plethora of cutaneous findings (primary systemic amyloidosis, acquired cutis laxa and mucinosis), all linked to MM. There is not, at present, another description of all these findings on the same patient.

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How to cite this article: Lavorato FG, Alves MFGS, Maceira JMP, Unterstell N, Serpa LA, Azulay-Abulafia L. Primary systemic amyloidosis, acquired cutis laxa and cutaneous mucinosis in a patient with multiple myeloma. An Bras Dermatol. 2013;88(6 Suppl 1):S32-5.