Papular mucinosis: A report of two cases

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Abstract

Papular mucinosis (PM) (lichen myxoedematous) is a unique, chronic idiopathic disease characterized by lichenoid papules or nodules due to dermal mucin deposition and a variable degree of fibrosis. PM is a quite rare disease of unknown etiology, with fewer than one hundred and fifty cases reported. In this paper, we present two cases of PM with no associated monoclonal gammopathy in two male patients aged 75 and 38 years, for its rare occurrence in the literature.

Keywords: Cutaneous mucinosis, papular mucinosis, paraproteinemia, lichen myxoedematous

Introduction

Mucinoses are a heterogeneous group of disorders in which abnormal amount of mucin accumulates in the skin, either diffusely or locally. Cutaneous mucinoses may be listed as primary, in which mucin deposition is the major histologic property resulting in clinically characteristic lesions, and secondary, in which mucin represents an associated histologic finding. The classification of cutaneous mucinoses is complicated because the pathogenetic mechanism of mucin accumulation is not fully understood.

Case Reports

Case 1
The first case was a 75-year-old man. He presented with asymptomatic, flesh-colored papules mainly distributed over his left leg (Figure 1). These papular lesions have raised five months ago. He had type 2 diabetes mellitus and gonarthrosis in his medical history. Laboratory studies and thyroid function tests were normal. The patient had left knee operations eight times due to recurrent prosthetic infection. Histopathological examination revealed hyperkeratosis, atrophic changes, vascular proliferation and widespread mucin accumulation in the papillary dermis (Figure 2). The myxoid appearance in the dermis was found to be mucin accumulation with toluidine blue staining (Figure 3). The final diagnosis was papular mucinosis (PM).

The department of hematology was consulted for investigation of monoclonal gammopathy. The patient’s protein electrophoresis and bone marrow biopsy were normal. Monoclonal gammopathy was not detected. Systemic treatment, such as melphalan, thalidomide, interferon

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alpha, autologous stem-cell transplantation, cyclophosphamide, plasmapheresis or intravenous immunoglobulin (IVIG) was not planned by hematologists. The radiation oncology department was consulted for radiotherapy. Radiotherapy was not planned because it could increase leg edema and affect prosthetic infected tissue. Narrow band ultraviolet (UV) B therapy was recommended to the patient but he refused treatment because he could not move. 0.1% retinoic acid and 0.1% diflucortolone valerate 10 mg were given twice daily. The patient’s consent to publication was obtained.

**Case 2**

The second case was a 38-year-old man. He presented with itchy, flesh-colored papules mainly distributed over his right back. Dermatologic examination revealed papular and nodular lesions on the back, especially over the right scapular area (Figure 4). His medical and family history was unremarkable. Thyroid function tests were normal. Skin biopsy was performed with the preliminary diagnoses of PM, eruptive collagenoma and lipoid proteinosis on the back of the two lesions. Histopathological examination revealed that starting from just below the epidermis, the upper dermis was myxoid in character and this view was confined to the upper dermis. There were hyperkeratosis, atrophic changes, vascular proliferation and diffuse mucin accumulation in the papillary dermis (Figure 5). The myxoid appearance in the dermis was found to be mucin accumulation with alcian blue staining (Figure 6). The final diagnosis was PM.

Hematology consultation was requested for investigation of paraproteinemia. The patient’s protein electrophoresis was normal. Bone marrow biopsy was not needed for diagnosis. Our patient was clinically diagnosed with localized papular mucinous lesions with the presence of lesions, histopathological evidence, and without monoclonal gammopathy and thyroid disease. The patient received 0.1% betamethasone ointment treatment. Significant regression of lesions was observed after two months of follow-up. The patient’s consent to publication was obtained.
PM (lichen myxoedematous) is a unique, chronic, idiopathic disease characterized by lichenoid papules, nodules due to dermal mucin deposition and a variable degree of fibrosis. Montgomery and Underwood classified four kinds of PM in 1953: a generalized lichenoid eruption, later named scleromyxedema, a localized or generalized lichenoid plaque form, a discrete papular form and an urticarial plaque form. PM and lichen myxoedematoses have often been used as indiscriminately synonyms in the literature. PM is generally associated with monoclonal gammopathy.

PM is a quite rare disease of unknown etiology with fewer than one hundred and fifty cases reported in the literature. PM contains two clinicopathologic types: a diffused papular and sclerodermoid form (named scleromyxedema) and a localized papular form. The distinction between these two forms is important because treatment approaches differ. The most important point of differentiation is the extensiveness of skin involvement. Scleromyxedema differs from other skin mucinosis by four diagnostic findings: generalized papular and sclerodermoid eruption, dermal mucin accumulation with fibroblast reproduction and fibrosis, and monoclonal gammopathy without thyroid disease. In the classification by Rongioletti and Rebora, which has been widely accepted in recent years, cutaneous mucinosis forms have been described as scleromyxoedema (generalized papular), localized, and atypical forms. Systemic mucin deposition may occur with gastrointestinal, pulmonary, renal, cardiac, and neurological involvements. Monoclonal gammopathy and paraproteinemia are detected in the vast majority (83.2%) of scleromyxoedema cases. 10% of cases can progress to multiple myeloma.

Diagnostic criteria for localized papular form include papular or nodular eruption, mucin accumulation, fibroblast reproduction at different grades without monoclonal gammopathy and thyroid disease. Lesions show slow progression with no systemic involvement. However, spontaneous recovery is extremely rare. We also accepted our cases as localized papular form. The localized form has five subtypes: discrete PM (DPM), which can appear anywhere on the body; acral persistent PM, which only affects the extensor surfaces of the hands and wrists; self-cure adolescent and adult type PM; infantile PM and, nodular form. Our patients were considered to have the DPM subtype of PM. DPM may be associated with hepatitis C virus (HCV) or human immunodeficiency virus (HIV) infections. We did not detect these diseases in our patients. About 20 cases of DPM have been described that were not associated with HCV or HIV infection.

Scleroderma (systemic sclerosis), sclerema, eruptive papular xanthoma, lichen amyloidosis, lichen planus and lichenoid drug eruption should be considered in the differential diagnosis of scleromyxedema. In particular, the existence of papules in linear arrays is a useful practical sign. In addition, papules not present in scleroderma are distinctive for scleromyxedema.

The treatment of PM therapy is very challenging because the literature about the treatment is restricted to case reports and series. Melphalan, thalidomide, high-dose dexamethasone, methotrexate, cyclophosphamide, chloroquine, retinoids, chemotherapeutic agents, psoralen plus ultraviolet A (PUVA), interferon alpha, radiotherapy, plasmapheresis, IVIG, and autologous stem cell transplantation are possible treatment options. The side effects limit the use of these treatments. The first suggested treatment is melphalan therapy (an alkylating agent). Successful results with IVIG and thalidomide have been reported in the literature. Retinoids inhibit mucin secretion by inhibiting fibroblast proliferation. PUVA and electron beam irradiation outcomes should also be discussed. The contribution of long-term remission of autologous stem cell transplantation to these cases has also been controversial in recent years. Favorable results of steroid treatment have been reported. Steroid therapy is thought to target both the production of paraprotein through its immunosuppressive and anti-fibroblast effects. We did not consider systemic treatment in our cases with limited skin involvement. We observed regression of lesions with high-potent topical corticosteroids.

As a result, PM is rare; it must be considered in the differential diagnosis of patients with skin-colored papular lesions.
Ethics
Informed Consent: It was taken.
Peer-review: Externally peer-reviewed.

Authorship Contributions

Conflict of Interest: No conflict of interest was declared by the authors.

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