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LETTERS TO THE EDITOR

Loss of histidine decarboxylase as a marker of malignant transformation and dedifferentiation of B-cells infiltrating the skin. A case report of a therapy-resistant multiple myeloma complicated by skin infiltration

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To the Editor

Multiple myeloma (MM) is a B-cell lymphoma of matured plasmacytic origin. There is a well described spectrum of cutaneous diseases in MM including cell infiltration and amyloid deposition [1]. Little is known about the basic processes involved that allow malignant plasma cells or lymphomas to grow outside the bone marrow environment. Several experimental models have been used to study this issue [2]. Hedvat et al. found altered gene expression profile pattern in plasma cells growing in extramedullary sites mostly involved in angiogenesis and adhesion [3]. Identification of other tumour-specific alterations required for extra-medullary growth would confer better understanding of tumour metastasis. In an earlier study authors found that histidine decarboxylase (HDC)- that is the only enzyme capable of histamine synthesis- is absent from B-lymphocytes infiltrating the skin in a B-cell chronic lymphocytic leukemia patient who developed cutaneous infiltrations in the course of the disease. In contrast to this, those cells that remained in the bone marrow had their HDC activity been preserved [4]. There is evidence that besides histamine being a well known mediator of allergic reactions, may also be involved in certain types of cell proliferations like wound healing, embryonic development and tumor growth [5,6]. There are data confirming that a functioning HDC gene is important in maintaining immune homeostasis [7]. The relation of histamine metabolism and metastasising human plasma cell malignancies has not been examined so far. The case reported here served an appropriate occasion for that.

Case report

A 70 year-old female multiple myeloma patient type IgG lambda, stage IA, developed purple coloured papular skin infiltrates involving both legs after 2 months ineffective thalidomide and 5 months melphalan plus prednisolone therapy (Figure 1).

The skin infiltrating cells were shown to be plasma cells (Figure 2). There was a significant difference in HDC content comparing the skin infiltrating plasma cells to bone marrow plasma cells: plasma cells infiltrating the skin became HDC negative during tissue invasion (Figures 3, 4). A very strong HDC positivity was found among bone marrow plasma cells identified by CD38 and CD138 in contrast to other mononuclear cells (Figures 5, 6).

Skin infiltrates came along with other signs of progression, like extension of bony lesions, fibronodular
lung manifestations and the development of heart and renal amyloidosis causing intractable oedema leading eventually to the patient’s death.

The disease was refractory to various chemotherapeutic protocols including continuous low dose Melphalan, cyclophosphamide, CVP, M2, VAD, high dose dexamethasone and local bone irradiation. The administration of the cytokine interferon alpha (IFN-α) had to be stopped because of intolerance. Twenty two months from diagnosis the patient died of cardiac and renal insufficiency.

**Laboratory results**

Initially WBC: 4.3 × 10⁹/l, PCV: 39%, Hgb 133 mg/dl, Platelet count 236 × 10⁹/l, bone marrow infiltration rate: 30%, IgG: 1925, IgA: 15, IgM: 28 mg/dl. Preterminally the degree of proteinuria was over 1.0 g/24 h, pancytopenia developed and the IgG concentration had also risen up to 3090 mg/dl.

(Between the two end points the M component was around 11 mg/dl for 8 months.) ECG showed low voltage. There were focal sparkling, pericardial fluid, dilated atria and concentric left ventricular hypertrophy seen on echocardiography as evidences of amyloidosis.

**Discussion**

The case presented here raises three questions: 1) the pathologic mechanism of epidermoinvasivity; 2) the role of histamine metabolism in the malignant process and 3) the therapy for secondary B-cell cutaneous lesions.

Secondary cutaneous infiltration developing in the course of a patient with B-cell neoplasm is a relatively rare event. In contrast to primary cutaneous infiltrations the secondary form in a
lymphoma patient usually means transformation of the disease, heralding a poor prognosis [4,8]. In all those cases had already been published, the absence of HDC in the cutaneous lesions was confirmed in contrast to lymphoma cells residing in the bone marrow, similarly to the case presented here. This phenomenon may be one of the alterations that allow the cells to become epidermotropic.

The presence of HDC in both benign and malignant cell proliferations has been well documented. Intracellular histamine may bind to binding sites other than histamine receptors. Ligands for the intracellular histamine receptor/binding sites appear to represent a new class of tumor promoting agents or perhaps might act as inhibitors [9]. The possible role of antihistamines and HDC inhibitors as antiproliferative agents is controversial and still under evaluation.Histidinol, an intracellular histamine receptor HIC antagonist, has an antiproliferative effect itself and also enhances cytotoxicity of anti-neoplastic drugs in combination. This effect is not uniform; it depends on cell type and dose [10]. Histamine potentiates alpha IFN-mediated antitumor effects, which may be due to histamine enhancement of NK and T-cell cytotoxicity [11]. This effect had been exploited in a trial where histamine was given to NHL patients previously treated with chemotherapy, to maintain remission [12]. IFN-α does not eliminate malignant cells but inhibits their overgrowth in culture [13]. This could be one possible explanation for the clinical finding in the B-CLL case with cutaneous infiltrations responding to INF-α as it has been reported earlier [4]. In that case the skin infiltrations disappeared within a month without healing of the leukemia process itself.

Conclusion

What we could learn from the case presented here is that epidermoinvasive plasma cells lose their HDC content. This is an acquired characteristic similar to the upregulation of adhesion molecules like CD44 which renders them able to disseminate and capable of skin invasion [14]. The acquired loss of HDC expression in the infiltrative cells points out that changes may occur not only on cell surface level-involving adhesion and angiogenesis-but even in the metabolism of such conservative molecules as HDC. Thus, extramedullary myeloma can be biologically altered from myeloma remaining in situ suggesting that the therapeutic approach should also be different. Hopefully further studies on histamine metabolism might lead to the development of more targeted therapy to lymphoma patients including those with cutaneous infiltrations.

Abbreviations

EBV positivity in primary cutaneous large B-cell lymphoma with immunophenotypic features of leg type: An isolated incidence or something more significant?

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To the Editor,
The WHO-EORTC classification divides cutaneous B-cell lymphomas into four categories of which primary cutaneous diffuse large B-cell lymphoma, leg type (PCLBCL, leg type) characteristically involves lower legs of elderly women and shows a predominance of confluent sheets of medium sized to large cells with round nuclei and prominent nucleoli resembling centroblasts and/ or immunoblasts. The neoplastic B-cells of PCLBCL, leg type have characteristic immunophenotype which readily differentiates them from other subtypes of primary cutaneous diffuse large B-cell lymphomas. The overall prognosis in this group is poor with a tendency to extracutaneous dissemination. An association of PCLBCL, leg type with Epstein Barr virus (EBV) has not been previously described. We report an unusual case of PCLBCL with immunophenotypic features of leg type with EBV positivity in the neoplastic B-cells by in situ hybridization. A fifty-five year old Caucasian male presented with a subcutaneous nodule on his right thigh which was comple-