Bazex Syndrome in Lung Squamous Cell Carcinoma: High Expression of Epidermal Growth Factor Receptor in Lesional Keratinocytes with Th2 Immune Shift

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Abstract
An 82-year-old Japanese man was referred for detailed examination of hyperkeratotic erythematous plaques on his palms and soles for 6 months. Two weeks before his first visit, he had undergone lung lobectomy for right lung squamous cell carcinoma (SCC). Laboratory findings showed elevations of eosinophil counts, serum IgE, thymus and activation-regulated chemokine, SCC antigen, and soluble interleukin-2 receptor levels. Histological results of a skin biopsy involving the left palm showed psoriasiform dermatitis. Before lung lobectomy, the hyperkeratotic erythematous plaques on the palms and soles and the erythemas on the trunk and extremities were difficult to treat with topical steroids. After lobectomy, the skin symptoms dramatically and rapidly subsided with topical steroids. Therefore, we diagnosed Bazex syndrome (BS), also known as acrokeratosis paraneoplastica, as a paraneoplastic cutaneous disease in lung SCC. The mild eosinophilia subsided and levels of SCC antigen, IgE,
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and soluble interleukin-2 receptor were reduced. BS is a paraneoplastic cutaneous disease characterized by acral psoriasiform lesions associated with an underlying neoplasm. In a previous report, a shift to the Th2 immune condition was found in patients with non-small cell lung cancer, as shown in our patient. Epidermal growth factor receptor (EGFR) is also known as tumor growth factor-α receptor; it is increased in psoriatic keratinocytes. In our case, EGFR expression increased in lesional keratinocytes 2 weeks after surgery and decreased 4 weeks after surgery. We speculate that a shift to Th2 immune reactions in lung SCC may be the pathogenesis of BS, whereby lesional keratinocytes highly express EGFR in parallel with disease activity.

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An 82-year-old Japanese man was referred for detailed examination of hyperkeratotic erythematous plaques on his hands (palms) and feet (soles) for 6 months. Two weeks before his first visit, he had undergone lung lobectomy for right lung squamous cell carcinoma (SCC). He had a history of tongue and gastric cancer, both of which were in remission. On admission, he had hyperkeratotic erythematous plaques on his palms and soles (Fig. 1). Multiple itchy, coalescing, scaly, papular erythemas, 20 mm in diameter, were on his trunk and extremities but did not cover more than 90% of his entire body surface area, which is the definition of erythroderma. Transverse abdominal folds were not spared, indicating that the “deck chair” sign characteristic of papuloerythroderma of Ofuji was negative. Laboratory findings were as follows: white blood cell count, 9.4 × 10³/μL (normal range, 3.6–9.3 × 10³/L); eosinophils, 9.4%, 8.3 × 10²/μL (normal range, 0.6–8.0%); hemoglobin, 12.6 g/dL (normal range, 13.8–16.9 g/dL); C-reactive protein, 0.11 mg/L (normal range, 0.0–0.2 mg/L); creatinine, 0.74 mg/dL (normal range, 0.7–1.1 mg/dL); alanine aminotransferase, 17 U/L (normal range, 8–48 U/L); IgE, 4,686 IU/mL (normal level, <295 IU/mL); thymus and activation-regulated chemokine (TARC), 3,984 pg/mL (normal level, 450 pg/mL); SCC antigen, 15.4 ng/mL (normal level, <1.5 ng/mL); and soluble interleukin-2 receptor, 1,050 U/mL (normal range, 145–519 U/mL). His Brinkman index score was 1,600 and he stopped smoking after the diagnosis of lung cancer. Histological results of a skin biopsy involving the left palm revealed parakeratosis, acanthosis, elongated rete ridges, and perivascular lymphocyte infiltration in the upper dermis. The granular layer remained and exocytosis of lymphocytes was found (Fig. 2a, b). Biopsy involving erythema of the left forearm revealed an eczematous reaction. Contrast-enhanced computed tomography did not detect additional internal malignancies. Before lung lobectomy, the hyperkeratotic erythematous plaques on the palms and soles and the erythemas on the trunk and extremities were difficult to treat with topical steroids. After lobectomy, the skin symptoms dramatically and rapidly subsided with topical steroids. There was no history of palmoplantar keratosis. Potassium hydroxide preparation and microscopic examination of the soles did not reveal any tinea infection. No drugs were suspected of causing adverse cutaneous reactions. Therefore, we diagnosed Bazex syndrome (BS), also known as acrokeratosis paraneoplastica, as a paraneoplastic cutaneous disease in lung SCC. Mild eosinophilia and elevated SCC antigen, IgE, and soluble interleukin-2 receptor levels improved to 8.9% (4.8 × 10²/μL), 13.2 ng/mL, 1,957 U/mL, and 486 U/mL, respectively; however, elevated serum TARC levels remained as high as 4,277 pg/mL 6 months after lung lobectomy.

BS is a paraneoplastic cutaneous disease characterized by acral psoriasiform lesions associated with an underlying neoplasm. The skin of the palms and soles initially thickens when the neoplasm is not active. Subsequently, the trunk and extremities become involved, as shown in our patient [1, 2]. The pathogenesis of BS is unknown, but some immunological
mechanisms are hypothesized and serum levels of SCC antigen can parallel the severity of cutaneous lesions [3]; however, they can also parallel the severity of the primary malignancy as a tumor marker. In a previous report, a shift to the Th2 immune condition was found in patients with non-small cell lung cancer. Radical surgery treatment may improve the Th1/Th2 shift in patients [4], as shown in our patient, who had an eczematous reaction on the extremities and trunk with the increased serum IgE and TARC and increased eosinophil count. Serum IgE level and eosinophil count were decreased after surgery. Cigarette smoking increases some inflammation markers, including TARC [5]. Epidermal growth factor receptor (EGFR) is also known as tumor growth factor-α receptor; it is increased in psoriatic keratinocytes [6] and decreased in atopic dermatitis (AD) keratinocytes [7]. We compared the expression of EGFR in lesional keratinocytes between 2 and 4 weeks after lobectomy. Surprisingly, its expression was increased in lesional keratinocytes of the left palm 2 weeks after surgery and decreased in those of the left sole 4 weeks after surgery (Fig. 2c, d). From these results, we speculate that a shift to Th2 immune reactions in lung SCC, which is similar to that in AD patients, may be the pathogenesis of acral psoriasiform lesions in BS, whereby lesional keratinocytes highly express EGFR in parallel with disease activity; however, it is usually suppressed in AD patients.

Statement of Ethics

The authors have no ethical conflicts to disclose.

Disclosure Statement

The authors have no conflicts of interest to disclose.

References

Fig. 1. Clinical appearance during the first visit. Hyperkeratotic erythematous plaque on the left sole. Hyperkeratotic erythematous plaques developed on the bilateral palms and soles. Nails were thickened and fissures were found. Examination using potassium hydroxide on the scales was negative for a fungal infection.
Fig. 2. Histological and immunohistochemical findings during the first visit. a, b Skin biopsy of hyperkeratotic erythema on the left palm. Hematoxylin and eosin staining revealed parakeratosis, acanthosis, elongated rete ridges, and perivascular lymphocyte infiltration of the upper dermis. The granular layer did not disappear and exocytosis of lymphocytes was found. Original magnifications: ×40 (a) and ×100 (b). c, d Immunohistochemical staining with EGFR of hyperkeratotic erythema on the left palm 2 weeks (c) and on the left sole 4 weeks (d) after lobectomy for lung cancer. Lesional keratinocytes highly expressed EGFR 2 weeks after surgery, and its expression was decreased 4 weeks after surgery. Original magnifications: ×100 (c) and ×100 (d).