Complete Remission of Recalcitrant Viral Warts under Oral Isotretinoin in a Patient with Low-grade B-cell Lymphoma

Alexandra Monastiri1, Panayiota Matsouka2, Efi Pasmatzi1, Maria Melachrinou3, Sophia Georgiou1, Elena Solomou2, Nicholas Zoumbos2 and Dionysios Tsambaos1

Departments of 1Dermatology, 2Hematology and 3Pathology, School of Medicine, University of Patras, PO Box 1413, Rio-Patras 265 04, Greece. E-mail: almonast@med.upatras.gr
Accepted November 4, 2004.

Sir,

Viral warts are the most frequent clinical manifestations of the ubiquitous human papillomaviruses (HPV) infection. Particularly in immunocompromised patients they may become extensive in both size and area of involvement, representing a frustrating problem, and can progress to squamous cell carcinomas (1, 2). We report herein the complete remission of recalcitrant and extensive warts in a patient with low-grade B-cell small lymphocytic lymphoma under treatment with oral isotretinoin (13-cis retinoic acid).

CASE REPORT

A 31-year-old HIV-negative Caucasian man with a 4-year history of both-sided submaxillary and cervical lymph node enlargement and an 8-year history of warty skin lesions presented to the outpatient Hematology Department at our University Medical Center. Physical examination revealed submaxillary, cervical/superclavicular lymph nodes in blocks of size 2 × 2 cm and 3 × 4 cm, respectively, whereas enlarged lymph nodes were palpable in the axillary and inguinal region, as well. Additionally, multiple, large and confluent hyperkeratotic viral warts were found on the dorsal aspect of both hands (Fig. 1A) and on the neck spreading over the beard area. Repeated earlier treatments with various modalities had resulted in a temporary remission of these skin lesions.

At admission, the counts of peripheral blood were: Ht 47, leukocytes 28000/dl; differential count: polymorphs 26%, lymphocytes 70%, monocytes 2%, eosinophils 6% and platelets 169,000/dl. Flow cytometry of peripheral blood revealed a B-clonal lymphocytosis (CD3+ 24%, CD20+ 58%, CD19+ 64%, CD20+/CD5+ 58%). Blood chemistry and serological tests were within normal limits. Whole body computed tomography confirmed the enlargement of peripheral lymph nodes. Histological examination of a skin biopsy revealed the typical features of common warts, i.e. hyperkeratosis and orthokeratosis with parakeratotic columns over the tops of the papillae, acanthosis, papillomatosis, keratohyalin condensation and perinuclear vacuolization, whereas in situ hybridization revealed the occurrence of HPV-2 DNA. The biopsy of an enlarged cervical lymph node revealed the morphological and immunohistochemical features of B-cell small lymphocytic lymphoma/chronic lymphocytic leukaemia (B-SLL/CLL) according to REAL and WHO classification with a CD20+, Bcl-2+, CD23+ and CD5+ phenotype. Bone marrow biopsy showed a nodular involvement from small B-cells with an identical phenotype and a low proliferation index (<5%).
Cytogenetic investigations revealed a normal karyotype (46,XY).

Our therapeutic decision was ‘to watch and wait’. However, during the subsequent 17-month follow-up without any treatment a gradual further enlargement of the affected lymph nodes and the occurrence of splenomegaly were observed. Thus, we decided to start treatment with rituximab (anti-CD-20 humanized monoclonal antibody) at a dose of 375 mg/m² by intravenous infusion weekly for 4 consecutive weeks, then monthly (six courses) and finally once every 3 months for six further courses as maintenance therapy. Therapy was well tolerated and side effects were limited to infusion-related symptoms such as fever, chills and flushing.

In contrast to the gradual regression of the lymph nodes of the patient seen under treatment with rituximab, his warty skin lesions clearly showed a marked progression; 8.5 months after the onset of rituximab monotherapy, the deterioration of his dermatological status prompted us to initiate oral administration of isotretinoin (1 mg/kg/day). Six weeks after the onset of isotretinoin treatment, an impressive therapeutic response of the patient to this retinoid was seen, characterized by a marked reduction in the number and size of the extensive viral warts; the latter completely disappeared after 10 weeks of oral treatment with this retinoid (Fig. 1B). The low CD4/CD8 ratio (0.7) observed in the peripheral blood of the patient before the onset of isotretinoin administration was found to remain unchanged upon completion of retinoid therapy. Apart from a marked cheilitis, the patient developed no significant clinical or laboratory retinoid-related side-effects. He has completed a 24-month follow-up without treatment with oral isotretinoin and shows no evidence of relapse of his skin lesions. Additionally, 10 months after completion of his maintenance rituximab therapy, he remains free of any recurrence of his lymphoma. The counts in his peripheral blood are normalized and the bone marrow infiltration by CD20+ B-lymphocytes has regressed to 10%. Flow cytometry of bone marrow lymphocytic populations revealed only 4% anti-CD20+ B-lymphocytes, whereas no CD20+ B-lymphocytes could be detected in the peripheral blood.

DISCUSSION

Rituximab is a chimeric human-murine IgG1 monoclonal antibody binding specifically to the B-cell surface antigen CD20 that is expressed in 95% of B-cell lymphomas and causes the eradication of the targeted CD20-positive lymphoma cells through the induction of complement-mediated and CD20-dependent cytotoxic effects and apoptosis (3). Recent clinical trials have demonstrated the efficacy of rituximab, administered as a first-line single agent, in the treatment of low-grade B-cell lymphoma with objective response ranging between 60% and 75% (4,5). Considering the promotion of B-cell maturation caused by isotretinoin in patients with common variable immunodeficiency (6) and the synergistic action of retinoid and rituximab on apoptosis induction of malignant B cells (7), it seems reasonable to suggest that isotretinoin may have enhanced the therapeutic effects of the latter on the lymphoma of our patient.

In view of the reported encouraging therapeutic results of oral retinoids in patients with extensive viral warts (8,9), we tried this at a dose of 1 mg/kg/day. The response of his skin lesions was impressive. To the best of our knowledge, it is the first time that oral retinoid therapy has led to the complete remission of extensive viral warts. Under oral administration of either etretinate or isotretinoin, a rapid favourable therapeutic response of the warts was observed but no complete remission was achieved, probably due to early dose reduction (8,10) or discontinuation of therapy because of the development of liposarcoma (9). In contrast to the relapse of viral warts observed in patients with epidermodysplasia verruciformis subsequent to discontinuation of etretinate treatment (11), our patient has completed a 24-month follow-up without any evidence of recurrence of his cutaneous lesions.

Isotretinoin is capable of dramatically affecting epithelial cell differentiation and proliferation (12). As HPV replication is related to the state of keratinocyte differentiation, it is possible that this retinoid may have inhibited the replication and assembly of HPV (13).
Although a spontaneous regression of the warts cannot be ruled out (14), it seems very unlikely in our patient in view of the extremely long duration of the recalcitrant skin lesions. The question as to whether rituximab (15) may have enhanced the therapeutic action of isotretinoin on the extensive viral warts remains to be elucidated.

REFERENCES