Classical Hodgkin Lymphoma Presenting as Cutaneous and Soft Tissue Mass Lesion: An Enigmatic Presentation Posing a Diagnostic Challenge—A Case Report and Review of Literature

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Abstract
Hodgkin lymphoma (HL) is primarily a nodal disease. Cutaneous involvement of HL as a presenting feature is extremely rare. Skin involvement is usually seen as a metachronous involvement in the course of disease and is associated with poor prognosis. Primary skin and soft tissue involvement can be construed as nonhematological, inflammatory, or infective etiology. We report a 14-year-old girl with fever, weight loss, lymphadenopathy, and multiple papular lesions over the right chest wall as initial manifestation of HL, posing a diagnostic challenge. In view of stage IVBE, patient was managed with intensive chemotherapy regimen and is currently free of disease at 6 months of follow-up.

Keywords ► cutaneous  ► extranodal  ► Hodgkin lymphoma

Introduction
Hodgkin lymphoma (HL), previously known as Hodgkin disease, is a monoclonal lymphoid neoplasm, originating most often from B lymphocytes. It is characterized by the orderly spread of disease from one lymph node group to another and usually begins in lymph nodes.\(^1\) Extrannodal invasion of adjacent tissue is observed in up to 15% of cases.\(^2\) Cutaneous involvement of HL as a presenting feature is extremely rare. Skin involvement is usually seen as a metachronous involvement in the course of disease and is associated with poor prognosis. We report here an interesting case of HL in a 14-year-old girl, with fever and lymphadenopathy associated with skin and soft tissue involvement as initial manifestation of HL, which posed a diagnostic challenge. Informed consent was obtained from the parents.

Case Report
A 14-year-old female presented with history of intermittent fever, weight loss, and progressively increasing bilateral neck masses over a period of 6 months. She also reported skin lesions over the right chest wall for 3 months and cough associated with breathlessness for about a month. There were no bone pains or symptoms suggestive of aerodigestive tract involvement.

Examination findings revealed the patient to be thin built, anxious, with pallor and mild respiratory distress, bilateral firm, nontender cervical (level 2–4), supraclavicular and axillary lymphadenopathy. The large conglomerate of fused right cervical nodes measured 7x4 cm. There were multiple papules and nodules present over entire right half of chest wall involving breast, infra-axillary, and scapular regions and...
extending to right shoulder and neck, few lesions were ulcerative with purulent discharge (►Fig. 1). Systemic examination revealed reduced air entry over right chest.

She was initially investigated elsewhere and in view of generalized lymphadenopathy and right-sided pleural effusion, she was clinicoradiologically diagnosed as pulmonary tuberculosis and had received 5 months of antitubercular therapy. Her skin lesions were suspected as herpes zoster infection and treated with acyclovir. She was unresponsive to both of the therapies, instead the lymph node mass and skin lesions progressed, for which patient was referred to our institute for further management.

At our institute, patient was investigated and initial complete blood counts, erythrocyte sedimentation rate (24 mm), liver and renal function tests were normal. Biopsy from the cutaneous lesion revealed unremarkable epidermis with underlying dermis showing infiltrate of polymorphous population of lymphocytes, histiocytes, and eosinophils. Numerous interspersed large neoplastic cells were seen monolobated to polylobated nuclei having vesicular chromatin and prominent nucleoli (►Figs. 2A and B). On subsequent immunohistochemical staining, the large scattered neoplastic cells were positive for CD30, PAX-5 (heterogenous; ►Figs. 2C and D) while negative for CD20, LCA, CD3, CD15, BCL-6, and EBER-ISH. Overall histopathological features were conclusive of classical HL, nodular sclerosis. The concurrent biopsy from lymph node also revealed HL involvement. Staging investigations revealed the bone marrow (aspirate and biopsy) to be uninvolved by HL, while positron emission tomography computed tomography (PET-CT) scan (►Fig. 3) showed metabolically active multiple bilateral cervical, supraclavicular, subpectoral, axillary, internal mammary, paratracheal, retrocaval, aortocaval, and para-aortic adenopathy, with extranodal disease involving bone (osteodestructive lesion involving head and shaft of right humerus), soft tissue deposits (right chest wall and right upper arm), right pleural deposits, and right pleural effusion with collapse, hence conforming to stage IVBE. The cross-sectional images reveal that the involvement was contiguous.
Subsequently, the patient was started on Adriamycin, bleomycin, vinblastine, dacarbazine chemotherapy protocol. However, disease progressed and after one cycle she was put on escalated bleomycin, etoposide, Adriamycin, cyclophosphamide, vincristine, procarbazine and prednisolone chemotherapy. Subsequent evaluation with PET-CT showed partial response (50–60%) after two cycles and complete response (including skin, except for residual pleural effusion) after four cycles (Fig. 3C). Hence, further consolidation with high-dose chemotherapy followed by autologous peripheral blood stem cell rescue was undertaken. After the completion of treatment, the patient is free of disease at 6 months of follow-up.

### Discussion

HL is primarily a nodal disease and pattern of spread is usually contiguous, spreading from one LN (lymph nodal) region to the next along the lymphatic system. Extranodal invasion of adjacent tissue is observed in up to 15% of cases, while hematogenous spread is seen in 5 to 10% of cases. We have described an interesting case of HL with multiple extranodal sites of disease with significant involvement of skin and soft tissue.

Cutaneous involvement of HL is extremely rare occurring in 0.5 to 3.4% of cases and is more frequently reported in non-HL. Although first reported in 1904 by Grosz et al., less than a hundred cases have been reported in literature, and this is the youngest case reported to the best of the author's knowledge (Table 1). Various types of skin lesions have been reported; these may be nonspecific findings due to paraneoplastic syndrome or vasculitis, like pruritis, urticaria, hyperpigmentation, or ichthyosis, or due to infiltration of skin by disease. The latter may present as nodules, papules, plaques, ulcers, and/or erythroderma, most of which we observed in the present case with biopsy proof of tumor infiltration.

Three different pathologic mechanisms have been described in the spread of cutaneous and extranodal HL, of which the most frequent is retrograde lymphatic spread followed by local tumor extension from an underlying nodal or extranodal site. The third mechanism includes hematogenous spread that if extensive has been seen to be associated with poor outcome. Our patient had multiple supradiaphragmatic nodal disease and extranodal disease involving upper end of right humerus, the skin and soft tissue of the right chest wall (up to the midline), right upper arm, and right pleura.

Skin involvement could also occur from direct extension of same side internal mammary or intercostals lymph nodal disease, which is often present along with chest wall involvement. Involvement of skin by HL is usually associated with more advanced disease as was observed in the present case; patient had stage IV disease that was refractory to first-line chemotherapy.

Interestingly, unilateral lung disease has been seen to occur from involved same-sided hilar adenopathy rather than from hematogenous spread. Pleural effusions occur in ~13% of cases and are usually negative for malignant cells.

Single osseous involvement is most likely the result of local spread rather than hematogenous process and is usually a late manifestation. It is believed that local extension from adjacent lymph nodes does not alter staging, although extranodal disease resulting from local infiltration (lung, bone, etc.) is regarded as stage IV as in the present case despite absence of widespread dissemination.

We believe that delay in diagnosis occurred in the present case due to two confounding clinical features that were

### Table 1

<table>
<thead>
<tr>
<th>Author (y)</th>
<th>No</th>
<th>Age (y)</th>
<th>Gender</th>
<th>Symptoms (No)</th>
<th>Duration (mo)</th>
<th>Stage</th>
<th>Treatment</th>
<th>Response (No)</th>
</tr>
</thead>
<tbody>
<tr>
<td>White and Patterson (1985)</td>
<td>16</td>
<td>16–63</td>
<td>Male (9) Female (7)</td>
<td>Lesion on Chest (11) Neck (3) Scalp (2)</td>
<td>32.1 (mean)</td>
<td>IV (7) III (4) II (5)</td>
<td>Multiple regimens</td>
<td>Death due to disease (11) Death due to unrelated cause (1) Lost to follow-up (4)</td>
</tr>
<tr>
<td>Jurisić et al (2005)</td>
<td>1</td>
<td>77</td>
<td>Female</td>
<td>Nodule arms and abdomen</td>
<td>6</td>
<td>II B</td>
<td>CVPP</td>
<td>No complete resolution</td>
</tr>
<tr>
<td>Rubenstein and Duvic (2006)</td>
<td>3</td>
<td>Male (1) Female (2)</td>
<td>Nodule over chest (n=2) Rash on trunk (n=1)</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td>Isao et al (2007)</td>
<td>1</td>
<td>44</td>
<td>Male</td>
<td>Nodule on back</td>
<td>32</td>
<td>IV B</td>
<td>Multiple regimens</td>
<td>Died</td>
</tr>
<tr>
<td>Dhull et al (2012)</td>
<td>1</td>
<td>22</td>
<td>Female</td>
<td>Fungating mass over mid-chest</td>
<td>24</td>
<td>IV B</td>
<td>ABVD</td>
<td>Disease free</td>
</tr>
<tr>
<td>Khawandanah et al (2014)</td>
<td>1</td>
<td>46</td>
<td>Male</td>
<td>Rash on chest and neck</td>
<td>3</td>
<td>IV E</td>
<td>AVD/Bend/GCD</td>
<td>Disease free 8 mo</td>
</tr>
<tr>
<td>Goyal et al (2014)</td>
<td>4</td>
<td>50 21 25 28</td>
<td>Female Male Male Male</td>
<td>Nodules over neck Ulcer over sternum Nodule on breast Rash arms and legs</td>
<td>1 6 12 4</td>
<td>IV B IV B IV B IV B</td>
<td>ABVD-RT ABVD-RT ABVD ABVD</td>
<td>Disease free 3 y Recurrence in 6 mo Disease free Disease free 2.5 y</td>
</tr>
</tbody>
</table>

Abbreviations: ABVD, Adriamycin, bleomycin, vinblastine, and dacarbazine; Bend, bendamustine; CVPP, cyclophosphamide, vincristine, procarbazine, prednisone; GCD, gemcitabine, carboplatin, dexamethasone; No, number of participants; RT, radiotherapy.
pleural effusion and skin lesions, both of which are uncommon primary clinical presentations of HL. With modern therapy, advanced HL has 5-year failure-free survival rate of ~80%.10 Hence, it is important for the oncologist and pathologist to suspect and diagnose skin lesions due to HL, especially in a pediatric population.

Conclusion
To conclude, cutaneous involvement, although rare, is a well-recognized manifestation of extranodal HL and can be seen on primary presentation that may mimic nonhematological soft tissue neoplasm. Hence, a diligent histomorphological examination along with intensified chemotherapy regimen is of paramount importance due to distinct therapeutic and prognostic implications.

Declaration of Patient Consent
The authors certify that they have obtained all appropriate patient consent forms. In the form, the patient has given her consent for her images and other clinical information to be reported in the journal. The patient understands that name and initials will not be published and due efforts will be made to conceal identity, but anonymity cannot be guaranteed.

Financial Support and Sponsorship
Nil.

Conflicts of Interest
There are no conflicts of interest.

References