Pediatric Pharyngeal IgD-positive Monoclonal Plasmacytoid and Plasma Cell Neoplasm

Shanxiang Zhang, MD, PhD* and Catherine Long, MD†

SUMMARY

Pediatric neoplasm with monoclonal proliferation of lymphoplasmycladoid lymphocytes and plasma cells is exceedingly rare and has essentially never been reported in immunocompetent children. Here, we report a previously healthy 13-year-old girl with a pharyngeal mass and enlarged cervical lymph nodes. The pharyngeal mass was composed of CD138+, CD79a+, MUM-1+, IgD+, CD20+, PAX-5+, CD43+, λ-restricted monoclonal plasmacytoid, and plasma cells. Scattered CD20+, PAX-5+ B cells were present in the background. The patient was treated as localized non-Hodgkin lymphoma (stage II) with cyclophosphamide, doxorubicin, vincristine, and prednisone and is in complete remission at 17 months from the last chemotherapy.

Key Words: immunocompetent children, neoplasm, lymphoplasmycladoid lymphocytes, plasma cells, immunoglobulin D positive

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CASE REPORT

A 13-year-old girl with a history of tonsillectomy at age 3 noticed an abnormal mass in her throat with occasional pain. Physical examination revealed a submucosal linear mass measuring 4 × 1 cm, located in right posterior oropharynx near the old tonsil fossa. An ENT physician at an outside institution performed an excisional biopsy of the mass. Upon pathology report from that institution she was referred to our tertiary care center [Indiana University Health (IUH)] for further evaluation. She had no fever, night sweats, or weight loss. Her past medical history was otherwise unremarkable. Her family medical history included cervical cancer in her maternal grandmother, thyroid cancers in the sister and niece of her maternal grandmother, and lymphoma (type unknown) in her paternal grandfather. There was no family history of immunodeficiency. A whole body computed tomography and positron-emission tomography scan revealed enlarged, bilateral, hypermetabolic, level II cervical lymph nodes, abnormal fullness, and intense metabolic activity in the lymphoid tissue in the tongue base, oropharynx, and nasopharynx. There was no evidence of metastatic disease outside of the neck. Her blood analysis showed normal complete blood count (white blood cells: 7.0 k/mm³, reference: 4.5 to 11.5 k/mm³; hemoglobin: 13.4 g/dL, reference: 12 to 15 g/dL; platelet: 195 k/mm³, reference: 150 to 450 k/mm³) and differential (5% neutrophils, 33% lymphocytes, and 9% monocytes), normal liver/renal functions, normal lactate dehydrogenase (162 U/L, reference: 100 to 242 U/L), and normal uric acid (5.4 mg/dL, reference: 2 to 7 mg/dL).

Review of the excisional biopsy at IUH revealed histologically unremarkable squamous mucosa overlaying marked proliferation of predominantly plasmacytoid lymphocytes and plasma cells. There were few scattered and rare aggregates of small lymphocytes (Fig. 1A). Few polyclonal B cells and immunophenotypically unremarkable T cells, normal liver/renal functions, normal lactate dehydrogenase (162 U/L, reference: 100 to 242 U/L), and normal uric acid (5.4 mg/dL, reference: 2 to 7 mg/dL).

Molecular study with multiplex PCR performed at IUH showed clonal Ig heavy chain γ rearrangement (Fig. 2). No cytogenetic karyotyping study was performed at either outside or our own institute. A descriptive diagnosis of monoclonal plasmacytoid and PCN, most compatible with extranodal MZL of mucosa-associated lymphoid tissue (MALT lymphoma) with plasmacytic differentiation was rendered. Subsequent bilateral bone marrow examinations performed at IUH revealed few scattered λ-restricted monoclonal plasmacytoid/plasma cells (supplemental Fig. 1). Supplemental Digital Content 1, http://links.lww.com/JPHO/A91, http://links.lww.com/JPHO/A92, http://links.lww.com/JPHO/A93, http://links.lww.com/JPHO/A94.

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IUH revealed no clonal Ig heavy chain \( g \) rearrangement (data not shown).

As pediatric MALT lymphoma has been reported in immunodeficient patients and gastric MALT lymphoma is associated with \textit{Helicobacter pylori} \((H. pylori)\) infection, the patient was evaluated for T-cell and B-cell subsets, Ig levels including IgG with subclasses IgA, IgM, and IgE, HIV, hepatitis panel, and \textit{H. pylori} infection. There was a borderline increase in her serum IgG antibody (1.03, reference range: <0.75) against \textit{H. pylori}. The patient had normal serum levels of IgA (112 mg/dL, reference: 47 to 317 mg/dL), IgM (72 mg/dL, reference: 56 to 242 mg/dL), IgE (14 kU/L, reference: 2 to 114 kU/L), and IgG (838 mg/dL, reference: 680 to 1531 mg/dL) including its subclasses. B-cell (355/mm\(^3\), reference 200 to 1259/mm\(^3\)) and T-cell (1524/mm\(^3\), reference: 1072 to 3890/mm\(^3\)) counts and their subsets were within normal reference range. Serum studies for hepatitis A, B, C, and HIV were all negative. There was no monoclonal protein by serum protein electrophoresis test.

The patient was treated as stage II non-Hodgkin lymphoma \((\text{NHL})\) per POG9219 protocol (https://members.childrensoncologygroup.org/prot/ProtoString.jsp?ProtocolNum = 9219&Disease = NHL). A 6-week induction phase included vincristine 1.5 mg/m\(^2\) weekly for 6 doses, doxorubicin 40 mg/m\(^2\) and cyclophosphamide 750 mg/m\(^2\) on days 1 and 22, respectively, and prednisone 40 mg/m\(^2\)/d for 28 days. This was followed by a shorter consolidation phase consisting of a single dose each of vincristine 1.5 mg/m\(^2\), doxorubicin 40 mg/m\(^2\), and cyclophosphamide 750 mg/m\(^2\) along with 5 days of prednisone 40 mg/m\(^2\)/d. She was also treated with lansoprazole, amoxicillin, and clarithromycin for \textit{H. pylori} due to positive \textit{H. pylori} IgG antibody.

Her postchemotherapy computed tomography and positron-emission tomography scan performed 1 week after completing chemotherapy revealed resolution of previous disease. Since that time she has been followed clinically and is currently in complete remission at 17 months from her last chemotherapy.

**DISCUSSION**

Small B-cell lymphoma with plasmacytic differentiation is used to describe a neoplasm of small B cells, plasmacytoid lymphocytes, and plasma cells. The major differential diagnosis includes MZL with plasmacytic differentiation and LPL.\(^1\,5\) In the pediatric population, MZL has been rarely reported including MZL with plasmacytic differentiation. However, essentially all the reported MZL cases contained aggregates and sheets of CD20\(^+\) B cells.\(^3\,5\) Pediatric MZL tends to be IgD-negative and presents with localized disease (stage I).\(^5\) The lesion in our case involved the patient’s oropharynx and cervical lymph nodes (stage II). Histologically, the lesion was composed of sheets of IgD-positive, \(\lambda\)-restricted monoclonal plasmacytoid lymphocytes and/or plasma cells. There were rare follicles as indicated by CD21 and CD23 stains. CD20 and PAX-5 stains revealed few B cells. The B cells were shown to be polyclonal by FCA, although the viability of the sample was very low (approximately 25%). The few B cells in our case were CD43\(^+\), although the B cells in the reported pediatric MZL tend to be CD43\(^-\). There were no background progressive transformed germinal centers as...
reported in pediatric nodal MZL; or monocytoid B cells and lymphoepithelial lesions as commonly seen in extranodal MZL. The molecular test (PCR) demonstrated clonal Ig heavy chain γ gene rearrangements, which supported a clonal neoplastic process. Overall we favor this lesion to be a B-cell lymphoma with extreme plasmacytic differentiation, although the presence of few CD43+ B cells added the difficulty of recognizing this case as a B-cell neoplasm.

LPL is essentially an adult disease. To our knowledge there were only 2 reported pediatric cases of LPL/ LPL-like lesions which were both associated with Wiskott-Aldrich syndrome and were self-limited. Our patient did not have any history of recurrent infections and workup for possible immunodeficiency was negative. Similarly, PCN is exceedingly rare in children and is by current World Health Organization definition a neoplasm secreting heavy chain class-switched Ig. Morphologically, however, our case was most compatible with LPL or PCN. As pediatric LPL and PCN are either extremely rare or have not been widely accepted in literature, and LPL in adults is typically IgD-negative, we were hesitant to render a diagnosis of LPL or PCN for this lesion.

Plasmablastic lymphoma (PBL) is a diffuse proliferation of large neoplastic cells which morphologically resemble B immunoblasts, but with the immunophenotype of plasma cells. The tumor cells are positive for CD138, CD38, VS38c, MUM-1, CD79a, and are negative or only weakly positive for CD45, CD20, and PAX-5. The Epstein-Barr virus study (EBER) is commonly positive. PBL is an aggressive lymphoma typically with a high proliferation fraction. Patients with PBL usually have advanced disease (stage III or IV) and die within the first year after diagnosis. Although it is most commonly seen in HIV-positive individuals, PBL has been reported in immunocompetent individuals. However, almost all PBL reported in immunocompetent individuals is seen in adults, whereas pediatric PBL is reported only in HIV-positive children.

Our}

### TABLE 1. Comparison of Marginal Zone Lymphoma, Lymphoplasmacytic Lymphoma, and Plasmablastic Lymphoma in Pediatric Population

<table>
<thead>
<tr>
<th></th>
<th>Nodal Zone Lymphoma</th>
<th>Extranodal Lymphoma</th>
<th>Lymphoplasmacytic Lymphoma</th>
<th>Plasmablastic Lymphoma</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Common affected sites</strong></td>
<td>Head and neck LN, lymphoid tissue</td>
<td>Ocular adnexa, salivary glands, skin</td>
<td>Generalized lymphadenopathy</td>
<td>Head and neck, skin</td>
</tr>
<tr>
<td><strong>Pathologic description</strong></td>
<td>Small to intermediate lymphocytes with scattered plasma cells, marginal zone expansion, PTGC features</td>
<td>Monocytoid cells, plasma cells, lymphoepithelial lesions, architectural distortion</td>
<td>Numerous plasma cells with scattered small lymphocytes</td>
<td>Large plasmablastic cells with conspicuous nuclei and basophilic cytoplasm, sheet-like or nest-like growth pattern</td>
</tr>
<tr>
<td><strong>Immunophenotype</strong></td>
<td>CD20+, CD43+ (~70%), CD5-, CD10-, BCL-6-</td>
<td>CD138+ and IgG+ plasma cells and CD20+, CD5−, CD10−, CD43− B cells</td>
<td>CD45+, CD5−, CD20−, VS38c+, MUM-1+, CD79a−/CD79a+, EMA−/EMA+, EBER−/CD56−, Ki-67 high (&gt;75%)</td>
<td>More males; predominantly HIV +; stage III and IV, rarely II</td>
</tr>
<tr>
<td><strong>Associated features</strong></td>
<td>Predominantly male; stage I, rarely II, III</td>
<td>Few with autoimmune disease; <em>Heliocobacter pylori</em> in gastric MZL; stage I, rarely II, III</td>
<td>Wiskott-Aldrich syndrome and 1 case also with Von Recklinghausen neurofibromatosis</td>
<td>Unknown</td>
</tr>
<tr>
<td><strong>Molecular markers</strong></td>
<td>Few trisomies 18, 3</td>
<td>Rare trisomy 3, <em>IGH-MALT1</em>, tetraploidy</td>
<td>Unknown</td>
<td>Possible t(8;14), <em>IGH/MYC</em></td>
</tr>
<tr>
<td><strong>Treatment</strong></td>
<td>Excision, CT, RT</td>
<td>RT, CT, excision</td>
<td>None</td>
<td>HAART, HARRT + CT, with or without RT</td>
</tr>
<tr>
<td><strong>Outcome/survival</strong></td>
<td>Excellent</td>
<td>Self-limited</td>
<td>Dismal, most died within 1.5 y</td>
<td>Dismal, most died within 1.5 y</td>
</tr>
</tbody>
</table>
patient was a previously healthy teenager who presented with a localized disease (stage II). The neoplasm she had showed very low proliferation index (<10%) as indicated by Ki-67 stain. The patient has been disease free for 17 months after receiving treatment for low-grade B-cell lymphoma. The initial clinicopathologic presentation as well as the disease response to the treatment makes the diagnosis of PBL highly unlikely. The major clinicopathologic features including treatment and prognosis for pediatric MZL, LPL, and PBL were summarized in Table 1. Other differential diagnosis includes atypical marginal zone hyperplasia with light chain restriction. These cases reported by Attygalle and colleagues all showed CD20+ B cells in the expanded marginal zone and follicular hyperplasia. These B cells were IgM, IgD-positive, and showed high proliferation index by Ki-67 stain. There was no significant plasmacytoid differentiation in the reported 6 cases. No evidence of clonality at the genetic level was demonstrated by PCR analysis.20 Our case instead showed predominantly IgD-positive, IgM-negative plasmacytoid, and plasma cells with both λ chain restriction by immunohistochemical stains and clonality by PCR analysis.

Here, we reported a highly unusual case of clinically stage II neoplasm composed of predominantly IgD-positive, λ-restricted monoclonal plasmacytoid, and plasma cells, morphologically most compatible with LPL or PCN, in a 13-year-old girl with no significant past medical history. The patient was treated per POG9219 protocol for localized NHL and remains in complete remission 17 months after her last chemotherapy. In patients with localized (stage I or II) NHL treated with POG9219, the 5-year event-free survival is 83.7%, with an overall survival of 96%. The standard follow-up includes complete blood count along with clinical history and physical examination. No imaging is necessary unless a relapse is suspected if the patient was in full remission at the end of therapy (https://members.childrensoncologygroup.org/prot/ProtInfo.asp?ProtocolNum = 9219&Disease = NHL).

Could this case represent an authentic pediatric LPL or PCN, or just a MZL with extreme plasmacytoid differentiation? In adults, the distinction between B-cell lymphoma with plasmacytic differentiation and PCN is critical as they require different treatment. In the pediatric population, MZL usually requires only local treatment with long-term follow-up.5 For our patient, long-term clinical follow-up may be helpful in the differential diagnosis. Report of other similar cases may also help to answer this question.

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REFERENCES

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