Blastic Plasmacytoid Dendritic Cell Neoplasm in a Young Female: Rare and Challenging

Abstract
Blastic plasmacytoid dendritic cell neoplasm (BPDCN) is a rare hematopoietic malignancy with an aggressive behavior, seen commonly in the elderly age group and usually involves the skin. Evaluation, diagnosis, and treatment pose unique challenges with poor outcomes. We report a case of an 18-year-old female who presented with lymphadenopathy and bicitopenia. The diagnosis of BPDCN was an initial challenge. She thereafter received intensive chemotherapy followed by an allogenic partially mismatched sibling donor hematopoietic cell transplantation, only to relapse in a few months. Beginning from the diagnosis, the management of this rare disorder and its relapse until her last followup provided several learning opportunities.

Keywords: Blastic plasmacytoid dendritic cell neoplasm, CD123, hematopoietic cell transplant

Introduction
Blastic plasmacytoid dendritic cell neoplasm (BPDCN), formerly known as primary cutaneous natural killer/T-cell lymphoma, has now been classified as a separate entity in the 2016 World Health Organization (WHO) myeloid neoplasms classification. [1,2] Blasmatoid dendritic cell has been described as the cell of origin.[1] BPDCN can be observed in all age groups, but it is more common in elders.[3,4] The male-to-female ratio is 2:2:1.[4] After the first case reported in 1994, around 200 cases have been described in the literature.[4] It constitutes 0.44% of all hematologic malignancies in a year.[4] Majority cases present with cutaneous manifestations such as plaque or nodules, red to purple in color.[1,5] Its presentation is often confused with other dermatologic disease manifestations. The disease also involves the bone marrow, spleen, and lymph nodes.[5] BPDCN may coexist with myelodysplastic syndrome, and 10%-20% cases may progress to acute myeloid leukemia.[5,6]

Case Report
An 18-year-old adolescent female presented with bilateral glandular neck swellings of 15 days. No systemic symptoms were observed. On examination, the cervical lymph nodes were firm to hard and nontender. There was a single cutaneous maculopapular lesion, 2.5 cm × 2 cm in size over left anterior aspect of the thigh, reddish-purple in color which resolved spontaneously. The liver and spleen were not palpable. Investigations showed a hemoglobin level of 7.8 g/dl, white blood cell count of 29,000/cu mm, and platelet count of 59,000/cu mm, with 68% atypical cells on peripheral blood smear [Figure 1]. Organ functions were unremarkable. Imaging identified retroperitoneal lymphadenopathy. Bone marrow aspirate and biopsy showed a hypercellular marrow with 87% blasts suggestive of acute leukemia [Figure 2]. Flow cytometric analysis revealed tumor cells to be positive for CD4, CD56, CD123, CD304 (BDCA-4), CD117, CD7, CD45, and human leukocyte antigen (HLA)-DR. The cells were negative for CD13, CD33, CD10, CD19, CD20, cMPO, TdT, CD41, and CD61 [Figure 3]. These findings were suggestive of BPDCN, although CD303 (BDCA-2) was not done due to unavailability. She received four cycles of hyper-CVAD chemotherapy regimen, and bone marrow was in morphological remission after cycle IB. She underwent allogenic hematopoietic cell transplantation (HCT) with an 8/10 HLA antigen partially mismatched (high-resolution typing) sibling donor.

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sister. Standard myeloablative conditioning with total body irradiation (TBI) and injectable busulfan was administered, and CD34 cell dose of $6.0 \times 10^6$ cells/kg (with a CD3 cell dose of $0.54 \times 10^8$ cells/kg) was infused. She received graft-versus-host disease (GVHD) prophylaxis with posttransplant cyclophosphamide on day +3 and day +4 along with tacrolimus and mycophenolate mofetil from day +5 onward. Neutrophil and platelet engraftment occurred on day +15 and day +16, respectively. During her peritransplant period, she developed acute gut GVHD on day +15 which responded to standard methylprednisolone therapy. While on methylprednisolone, she had a reactivation of cytomegalovirus for which she received ganciclovir for 2 weeks followed by valganciclovir. She was on regular follow-up without significant symptoms or morbidity but developed asymptomatic medullary relapse 19 months posttransplant. Intensive treatment and alternative “palliative” options were discussed. The patient and family chose to pursue decitabine monotherapy with intent for disease control only due to financial constraints. The patient developed febrile neutropenia after two cycles of decitabine and eventually died due to intracranial hemorrhage thereafter, 24.5 months from the initial diagnosis.

**Discussion**

Management of this young patient provided three learning opportunities for the treating team: arriving at a diagnosis in an unusual presentation of a very rare disorder, clinical management of aggressive disease in the background of scanty literature evidence, and treatment of relapse in such a disease setting. Our patient had peripheral blood and bone marrow involvement with cervical and retroperitoneal lymphadenopathy at presentation. There was a single cutaneous lesion which resolved spontaneously before treatment. With respect to diagnosis, the 2008 WHO Classification identifies BPDCN as a neoplasm which typically expresses CD4 and CD56 in addition to at least 1 of the pDC-associated antigens – CD123, TCL-1, CD304, or CD303/BDCA2 – in the absence of lineage-specific markers. A diagnostic criteria with 5 point scoring system based on flow cytometry has also been mentioned wherein a score of more than 2 favors the diagnosis of BPDCN. To date, there has been no consensus regarding the standard treatment of this disease. Although the disease usually shows early clinical response, relapse is very common. Various chemotherapy combinations such as acute myeloid leukemia (AML)-based induction, cyclophosphamide, doxorubicin, vincristine and prednisolone (CHOP), hyper-CVAD, and steroids have all been used. In a retrospective analysis of BPDCN patients who were treated with hyper-CVAD, CR rate was found to be 90% and OS ranged from 23 to 29 months, which was superior to a CHOP-like regimen. A recent review recommends AML like induction therapy followed by consolidation with allogenic HCT, owing to its myeloid lineage. HCT in first complete remission (CR1) has been suggested for a durable outcome. As per the European Society for Blood and Marrow Transplantation, there was an increase in overall survival and no relapse within 27 months in patients who received allogenic HCT in CR1. In a case series of HCT in ten patients (6 allogenic HCT and 4 autologous HCT) with a median age of 28.5 years and conditioning regimen of cyclophosphamide with TBI, overall survival was improved with the use of myeloablative conditioning followed by HCT (31.5 months) as compared to chemotherapy alone (13 months).
Management of relapsed disease is also fraught with worse outcomes using conventional chemotherapy as well as newer reported therapeutic options such as hypomethylating agents, bendamustine, pralatrexate, and venetoclax. It is worth noting that tegraxofusp-ezrs (ELZONRIS, a Monoclonal Ab, SL-401, directed against interleukin-3 receptor alpha chain [CD123] along with a truncated diphtheria toxin fusion protein), an immunotoxin genre of targeted immunotherapy has been recently approved by the USFDA (December 2018) with a reported CR rate of around 55% with a duration of response of around 5 months. Ongoing clinical trials evaluating novel therapeutic modalities strategies such as CAR-T cells (e.g., NCT03203369 and NCT03485547) will shed more light on the management of this aggressive malignancy.

BPDCN carries a poor prognosis with a median survival of around 12–16 months, and overall survival depends on the age and clinical characteristics of the patient. This case report exemplifies a rare disorder presenting at an unusual age with challenges of the diagnosis and management that are unique. It is important to initiate the treatment early and consolidate it with HCT for longer remission.

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Conflicts of interest
There are no conflicts of interest.

References