Interdigitating Dendritic Cell Sarcoma

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Abstract
Interdigitating dendritic cell sarcoma (IDCS) is an extremely rare dendritic cell tumor with slightly more than 100 cases reported in the English literature. This report discusses a case of localized IDCS involving cervical lymph nodes and provides a literature review of clinicopathologic aspects and treatment outcomes. (J Natl Compr Canc Netw 2015;13:128–132)

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Learning Objectives
Upon completion of this activity, participants will be able to:
• Evaluate the patient demographics, clinicopathologic features, and therapeutic outcomes of documented IDCS cases
• Discuss challenges in the accurate pathologic diagnosis, staging, and treatment of patients with IDCS
**Case Report**

An 81-year-old Caucasian man presented with a 4-month history of a gradually enlarging and painless left neck mass. He had no systemic or constitutional symptoms. His medical history included poorly differentiated squamous cell carcinoma of the scalp, which was treated with wide local excision and skin graft 2 years prior to presentation, and high-grade papillary urothelial carcinoma of the bladder without muscle invasion 1 year prior, which was treated with transurethral ablation. He was otherwise in good health. Physical examination revealed a 3-cm, nontender, immobile mass in the left posterior triangle of the neck. No axillary, supraclavicular, or inguinal lymphadenopathy was present, and no organomegaly was identified. CBC count and results of renal and liver function tests were within normal limits. Results of the HIV antibody test were negative. An excisional biopsy of the neck mass revealed interdigitating dendritic cell sarcoma (IDCS) with lymph node involvement. A PET scan performed a month after the excision revealed persistent fluoro-deoxyglucose (FDG) avidity in an area of the left posterior triangle of the neck measuring 1.08 cm in maximum transverse diameter, with a maximum standardized uptake value of 3.14 (Figure 1). The presence of residual tumor was the likely cause of persistent FDG avidity at the surgical site; local radiation or reexcision was recommended. A left modified radical neck dissection was performed to exclude a metastasis from prior squamous cell carcinoma of the scalp; 53 lymph nodes were removed, 1 of which showed IDCS and the remainder were benign. The patient was offered local radiation postsurgery, but refused adjuvant therapy. The patient has been disease-free for more than 18 months.

**Pathology**

Histologic sections of the lymph node showed diffuse replacement of the normal architecture by a spindle cell neoplasm, with effacement of subcapsular sinuses and extension into perinodal fat (Figure 2A/B). Neoplastic cells were disposed in fascicles with a focal storiform pattern, had mostly large spindle-shaped vesicular nuclei, occasional prominent nucleoli, and abundant eosinophilic cytoplasm with well-defined cellular borders (Figure 2C). In other areas, tumor cells had epithelioid or histiocytoid cell morphology or had convolutions resembling Langerhans cells. Occasional bizarre, binucleated, or multinucleated (some Langhans-like) forms were present in addition to a few cells with hyperchromatic nuclei. Rare biobed nuclei resembling Reed-Sternberg cells were seen (Figure 2D). Mitotic activity was increased (29 mitoses per 50 high-power field) with many atypical mitoses and few areas of tumor necrosis.

![Figure 1](image1.png) PET scan revealing an FDG-avid residual lymph node in the left posterior triangle of the neck at the previous surgical site (arrow).

![Figure 2](image2.png) (A) Lymph node with paracortical and interfollicular involvement by interdigitating dendritic cell sarcoma, with effacement of normal architecture and extension into adjacent adipose tissue (hematoxylin-eosin, original magnification x20). Occasional residual disrupted cells are seen (arrows). (B) The disrupted follicles are highlighted by CD21, not expressed in the tumor cells (CD21, original magnification x20). (C) The neoplastic cells have variable size and predominantly spindle cell morphology, and show nuclear and cytoplasmic S100 labeling (inset: S100, original magnification x200). Note the intermixed small lymphocytes and plasma cells in the background. (D) Neoplastic cells are predominantly epithelioid; some histiocytoid or Langerhans cell-like and rare multinucleated forms are seen in a hyalinized background (hematoxylin-eosin, original magnification x600).
Immunohistochemical studies showed that neoplastic cells strongly and diffusely expressed vimentin, S100 (cytoplasmic and nuclear pattern; Figure 2C), and p53, and were negative for CD1a, CD3, CD4, CD15, CD20, CD21, CD23, CD30, CD34, CD35, CD43, CD45, CD68, CD163, lysozyme, pancytokeratin, CK 5/6, EMA, p63, HMB-45, Melan-A, D2-40, SMA, and desmin. The proliferation index determined with Ki-67 labeling averaged 20% (range, 10%–40%). Background lymphocytes were mostly of T-cell lineage, and residual B cells were sparse.

Results of in situ hybridization for Epstein-Barr virus–encoded small RNA were negative, and B- and T-cell polymerase chain reaction clonality studies were polyclonal. The previous squamous cell carcinoma of the scalp was histologically different from the nodal tumor at review.

Discussion

Dendritic cells (DCs) are a heterogenous group of nonlymphoid, nonphagocytic immune accessory cells present in lymphoid and nonlymphoid organs. They are key antigen-presenting cells and initiators of the immune response. Four types of DCs exist in lymph nodes: follicular, interdigitating, Langerhans, and fibroblastic cells.1 DC neoplasms are rare and represent less than 1% of all lymph node tumors.2 These tumors were previously classified as lymphomas, sarcomas, or histiocytic neoplasms on histologic grounds alone, but recent diagnostic advancements have refined their classification. The WHO currently classifies these neoplasms based on their putative histogenesis and phenotype into myeloid-derived histiocytic neoplasms (histiocytic sarcoma), myeloid-derived dendritic neoplasms (Langerhans cell histiocytosis and sarcoma, IDCS, indeterminate DC tumor, and probably plasmacytoid DC tumor), and stromal-derived dendritic neoplasms (follicular DC sarcoma and fibroblastic reticular cell tumor).3

Interdigitating DCs (IDC) present antigens to T cells and are normally seen in T-cell zones of lymphoid organs, such as the paracortex of lymph nodes and the periarteriolar lymphoid sheaths in the spleen.4 IDCs are believed to originate from differentiation of marrow precursor cells; Langerhans cells can mature into IDC upon migration to lymph nodes, thus enhancing their antigen-presenting function.5–7

IDCS is a rare neoplasm. A recent comprehensive review of dendritic sarcomas by Saygin et al8 presented the demographics, clinicopathologic features, and therapeutic outcomes of 100 cases of IDCS. Patients are generally adult men with a median age at diagnosis of 56.5 years.8 Saygin et al8 reported that, of 39 patients, 16 (41%) were Asian, 19 (49%) were Caucasian, 2 (5%) were Hispanic, and 2 (5%) were African American. The most common presentation was painless localized lymphadenopathy, occurring in 47% of cases (sites in order of frequency: cervical, axillary, abdominal, inguinal, or mediastinal regions), and 25% of patients presented with isolated extranodal disease (the most common sites being liver, lung, spleen, bone marrow, and gastrointestinal tract); the remaining patients presented with both nodal and extranodal disease.8 Systemic symptoms (fever, night sweats, weight loss, and fatigue) were uncommon and mostly occurred in patients with both nodal and extranodal disease.8 The incidence of distant metastasis was 39%, with the most common sites of metastasis being lymph nodes (29%), lung (11%), liver (11%), and bone marrow (8%).8

The etiopathogenesis of IDCS is unknown. Epstein-Barr virus has been suggested as a causative factor in the pathogenesis of follicular DC sarcomas,9 but not in IDCS. In addition, human herpesvirus-8 genome has not been identified in IDCS.10 However, the association of IDCS with preceding or concomitant malignant neoplasms has been suggested in several case reports. Saygin et al4 reported occurrence rates of 12% and 9% in hematopoietic (eg, T-cell lymphoblastic lymphoma, low-grade follicular lymphoma, chronic lymphocytic leukemia/small lymphocytic lymphoid tumor [CLL/SLL], and large B-cell lymphoma) and solid organ malignancies (eg, liver, stomach, colon, brain, breast, and skin neoplasms), respectively, in patients with IDCS during their lifetime. This high frequency of concomitant malignancies may be coincidental and related to aging, or may support the hypothesis that DCs introduce tumor antigens to immune cells and that their dysfunction could result in a diminished response to the neoplasm. In the present case, the patient had a history of 2 solid organ malignancies (bladder and skin).

Previous case reports and small series have demonstrated a clonal relationship between low-grade B-cell lymphomas (follicular lymphoma and CLL/SLL) and histiocytic/DC tumors through identical
molecular alterations and monoclonal immunoglobulin heavy chain gene rearrangements. This finding of shared underlying molecular alterations in low-grade B-cell lymphomas and IDCS led to the concept of transdifferentiation of B-cell neoplastic cells into mature neoplastic DC, of which IDCS is common.

A diagnosis of IDCS can be challenging to a general pathologist and even a hematopathologist because of its rarity and morphologic similarity to a variety of primary and metastatic spindle cell neoplasms involving lymph nodes. Because of the morphologic overlap of histiocytic, Langerhans, follicular, reticular, and interdigitating tumors, ancillary studies are needed for a definitive diagnosis. Other tumors that require exclusion are spindle cell carcinoma, melanoma, and a variety of mesenchymal neoplasms. Approximately 11% of IDCS cases are misdiagnosed as lymphoma, melanoma, peripheral nerve sheath tumor, or malignant fibrous histiocytoma. Histologically, IDCS shows a diffuse proliferation of round-ovoid cells to spindle cells, with a variety of growth patterns, including sinusoidal, nesting, fascicular, and storiform, such as in the present case, and the proliferation can partially or completely replace the affected tissue. A collagenous or hyalinized background, with increased reticulin fibers surrounding individual cells, may be present. Microscopically, individual cells have a slender, spindled to plump (histiocytoid) appearance with ill-defined cell borders, abundant eosinophilic cytoplasm, and enlarged indented nuclei. The cells resemble Langerhans cells in microscopic appearance, but lack both Birbeck granules ultrastructurally and expression of CD1a and langerin. Occasional epithelioid cells, bizarre atypia, or Reed-Sternberg-like cells, as seen in the present case, may superficially resemble classical Hodgkin lymphoma. Necrosis is unusual and the mitotic rate is variable. Tumor cells are positive for S-100, vimentin, HLA-DR, and CD68, with CD68 weakly and variably expressed. In the present case, all histiocytic markers were negative. Variable staining has been reported for CD1a, CD1c, CD45RB, CD45RO, CD4, and CD14. IDCS is negative for CD21, CD35, CD3, and CD20. The Ki-67 labeling index usually ranges between 10% to 20%. IDCS may be difficult to distinguish from metastatic melanoma, which can have a similar immunohistochemical profile (S-100+, CD68+/−). The lack of other melanocytic markers establishes the diagnosis. The significant immunophenotype variability of DCs may reflect the multiple pathways postulated in normal DC development.

After establishing the pathologic diagnosis, staging should be completed, which includes imaging with CT scans and a bone marrow biopsy to evaluate for systemic disease. Primary bone marrow involvement seems to be less common in adults (11% in Saygin et al). Whether bone marrow biopsy is required in patients with localized disease who lack cytopenias and/or systemic symptoms is unclear. The role of PET scans in these rare tumors is also unclear. IDCS has been shown to be FDG-avid in some cases, and PET can be used to monitor treatment response. In the present case, PET scan detected residual disease and prompted surgical reintervention.

Currently, no standard of care exists for the treatment of IDCS. Localized disease is managed with surgery or radiation. Saygin et al reported no significant difference in survival among patients with localized IDCS who underwent surgery compared with those treated with nonsurgical modalities. Therefore, radiotherapy can be a reasonable alternative to surgery in select cases. However, the role of adjuvant radiotherapy after surgery is unclear. Chemotherapy is the preferred treatment for patients with metastatic IDCS. Combination chemotherapy regimens administered for aggressive lymphomas can be considered, such as CHOP (cyclophosphamide, doxorubicin, vincristine, prednisone), ABVD (doxorubicin, bleomycin, vinblastine, dacarbazine), and ICE (ifosfamide, carboplatin, etoposide). Patients with localized disease had significantly superior overall survival compared with those with metastatic disease (2-year survival rates, 68.1% vs 15.8%, respectively), compared with follicular dendritic cell sarcoma as stage did not significantly alter overall survival. The patient in the present case presented with localized IDCS that was treated with surgery alone and has been disease-free for more than 18 months. Young age (≤40 years), presence of intra-abdominal involvement, and combined nodal and extranodal involvement were associated with adverse outcomes (local recurrence, distant metastasis, or death), whereas gender, tumor size (≥6 cm), and histologic features such as presence of necrosis, high mitotic rate, and lymphoplasmacytic infiltration did not seem to predict recurrence or survival.
Conclusions

IDCS is an extremely rare tumor, with slightly more than 100 cases reported to date. The disease can be associated with other various malignancies (including hematolymphoid and solid tumors) occurring either synchronously or metachronously, which may create challenges for disease management. The histomorphology and immunoprofile help differentiate IDCS from other spindle cell neoplasms and metastatic melanomas. Patients with localized IDCS are treated similarly to patients with soft tissue sarcoma with primary surgical resection, and have better outcomes compared with those with metastatic disease. Young age, intra-abdominal involvement, and advanced stage are poor predictors of outcome. The role of adjuvant therapy remains unclear. Chemotherapy options for metastatic disease are controversial.

References


Instructions for Completion

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Posttest Questions

1. True or False: IDCS and follicular dendritic cell sarcoma have distinct histomorphologic features but similar biology and clinical course.
2. Patients with IDCS can also have preceding or concomitant hematopoietic and solid organ malignancies. The reported occurrence rate of these malignancies during the lifetime of IDCS patients is approximately:
   a. 10%
   b. 40%
   c. 70%
   d. 90%
3. Which of the following neoplasms should be considered in the differential diagnosis of IDCS?
   a. Spindle cell carcinoma
   b. Follicular dendritic cell sarcoma
   c. Melanoma
   d. Non-Hodgkin lymphoma
   e. Langerhans cell histiocytosis
   f. All of the above