Interdigitating dendritic cell sarcoma of the liver and lung: a case report with morphological and immunohistochemical features of tumor

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ABSTRACT

Interdigitating dendritic cell sarcoma is extremely rare neoplasm that mainly occurs in the lymph nodes. Only 45 cases have been reported in the literature to date. We report a case of this sarcoma arising from the liver and lung, a previously unreported site for this neoplasm. An 13-year-old girl deteriorated rapidly after artificial abortion and died 7 weeks later. Autopsy showed markedly enlarged liver and lung with numerous nodules up to 0.5 centimeters in diameter. Microscopically, nodules were composed of large pleomorphic cells that were immunohistochemically positive for proteins S-100 and vimentin, some of them expressed positivity to fascin and CD68, with a rich small CD3 positive T lymphocytic infiltrate around them. Based on these findings, the present case was diagnosed as interdigitating dendritic cell sarcoma, a neoplasm that remains a diagnostic and clinical challenge, because it can mimic a wide variety of other malignant tumors and tumor-like lesions.

KEY WORDS: interdigital dendritic cells, sarcoma, extranodal location.

INTRODUCTION

Accessory immune system contains two categories of cells: antigen-presenting (dendritic cells) and antigen-processing cells (macrophages). Dendritic cells are heterogeneous group of cells to which we include: Langerhans cells (skin, cervix, vagina, stomach and esophagus), dermal dendrocytes, follicular dendritic cells (FDS) and interdigitating dendritic cells (IDS). IDS are primarily located in T cell zone of lymphoid tissue (lymph nodes, thymus and lien). According to the WHO classification of hematopoetic and lymphoid tumor tissues, tumors of dendritic cells are classified as “neoplasm of histiocytes and dendritic cells origin”, which has a distinct category of dendritic cells neoplasm, which also includes sarcoma IDS [1]. Sarcomas of dendritic cells are very rare. The most often are diagnosed FDS sarcomas in lymph nodes [2], while extranodal location is extremely rare and is found in 1/3 of published cases [3-10].

IDS sarcomas (SIDS) are even more rare neoplasm which same as FDS sarcoma appears mostly inside lymph nodes [10]. Up to date literature described so far 45 cases of SIDS [11]. Extranodal localization of neoplasm is recorded in duodenum, spleen, thyroid, breast, palpebra, tibia, salivary glands, testicle [11-19]. In this paper we described clinical and pathological characteristics of a case of extranodal localization of SIDS in liver and lungs, entity that clinically was not recognized and had fast and fatal outcome.

CASE REPORT

In 19-year old women, following abortion due to unplanned pregnancy four times curettage was performed due to metrorrhagia, which was believed to be caused by the resida of placental tissue in uterine cavity. Curetment was not submitted to a pathohistological analysis. Patient complained to slackness, fast lost of body weight and high body temperature which was followed in early stage of disease with jaundice development. Routine laboratory finding established anemia and slightly elevated sedimentation. Day before death and four weeks following abortion patient developed melena, which was diagnosed via gastroscopy as ulcer and treated conservatively. Couple of hours following gastroscopy patient became unconscious, developed coma and in the end died.

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Autopsy findings
Liver was extremely enlarged. Lower liver edge reached upper pelvic edge. In liver and in the lungs numerous grayish-white nodules diameter to 5 mm surrounded with hyperemic zone were found. Bilaterally in lungs in lower lobes a single area without air, dark red color, on cut surface triangle shape with the tip of the triangle directed toward hilus and base to the surface of lungs. Below pleura and also in lung parenchyma numerous areas of infiltration with blood. Renal capsule was easy to remove, the surface was smooth. Renal cortex was widened and unclear picture. The brain was extremely swollen. Dark red blood clots were found in uterine cavity. Prominent cyanosis was found in other tissues and organs.

Pathological findings
Microscopic examination of liver tissue revealed that tissue structure was impaired with infiltrate of atypical cells places among sinusoids (Figure 1A.), while in portal area and acini, tumor cells there are foci in form of larger and bigger aggregates (Figure 1B.). Inside acini tumor aggregates did not demonstrate zonal predilection. Atypical cells were of very polymorphic shape and size, abundant cytoplasm, round and oval nuclei, among them there were multinuclear giant cells and those with eccentric lobulated and deeply notched nuclei, with small invisible nucleoli (Figure 1A.). Numerous pathological mitoses are visible (do 10/10 HPF). Aggregates of neoplastic cells are surrounded with reactive cells, lymphocytes, mononuclear and multinuclear histiocytes, plasma cells, and scant polymorphonuclear leukocytes. Surrounding hepatocytes demonstrate reactive changes. Liver tissue was distorted in site of nodular aggregates. Hyperemia was observed in lungs with accentuated capillary thrombosis and focally hemorrhagic.
infiltration of parenchyma (Figure 2A.). In both lower lobes, macroscopically observed area of hemorrhagic infarction was confirmed microscopically. Below pleura and focally in interstitium visible nodular aggregates of atypical cells with identical morphology (Figure 2B.) same as those described in liver. Acute tubular necrosis was found in kidneys. In other tissues besides hyperemia areas of focal hemorrhagic infiltration are found. Lymph nodes are not affected the tumor process. The endometrium has only basal layer which is moderately infiltrated by lymphocytes, histiocytes, plasma cells and eosinophils. Endocervix was chronically inflamed.

*Immunohistochemical findings*

Immunohistochemically, tumor cells demonstrated strong, uniform expression of S-100 protein (Figure 3.) and vimentin (Figure 4.), focal positivity to fascin (Figure 5.) and CD 68 (Figure 6.), while they were negative to CD15, CD30, CD43RO, CD43, CD23, CD21, CK116, PLAP, ALK, HMB45, EMA, CD20, CD79a, Cyclin D1, lysosim, actin and desmin. Surrounding tumor cells there was variable number of small CD3 positive T lymphocites (Figure 7.).

**DISCUSSION**

Our case presents primary localization of SIDS in lungs and liver, form that so far has not been presented in literature. The disease developed in 19 female patient developed without any symptoms and they appeared for the first time during her last month of life and overlapping with period follow- ing abortion. SIDS is diagnosed post mortem, following autopsy, based on morphological and immunohistochemical finding of tumor tissue. Interdigitating dendritic cells sarcoma is a very rare neoplasm of antigen presenting cells. So far 78 cases of SIDS has been published worldwide. Mostly it was a nodal localization of the tumor in cervical lymph nodes with secondary spread to other tissues and organs (bone marrow, bones, lien, liver, lings, ovary and skin). The disease was rarely accompanied with appearance of systemic symptoms (loss of body weight, increased body temperature and/or anemia). Most often it develops asymptomatically. Most of the patients were adults (age range was 6-87), more often male gender (M: F=1,5:1), average survival period to 15 months following diagnosis [12]. The disease was of unknown etiology [10]. There were couple of cases of close sarcoma of follicular dendritic cells there was and infections with EBV or HHV-8 [10]. Extranodal localization of SIDS was described in 1/3 of cases [12]. The optimal therapy for this malignant disease (radiotherapy or standard chemotherapy) still has not been determined due to small number of diagnosed cases and on
the other hand to relatively fast lethal outcome which does not leave enough time for diagnosis and study of the disease.

CONCLUSION

Above described case is interesting for three reasons. First of all, SIDS is a very rare neoplasm. Second one, coincidence of abortion and primary disease. And in the end, according to our knowledge this is the first published case of primary localization of the SIDS in lungs and liver.

DECLARATION OF INTEREST

The authors declare no conflict of interest.

REFERENCES