Case Report

Solitary nodule on the back: A diagnostic challenge on cytology

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ABSTRACT

Evaluation of a solitary lesion on the back can be a diagnostic challenge on cytology. However, based on the histological examination, differential diagnosis can be narrowed down, but still, sometimes, it becomes difficult to make a definitive diagnosis, even on biopsy findings. In such cases, the role of immunohistochemistry (IHC) becomes important. Herein, we present the case of a 49-year-old female who presented with a solitary nodule on the back, where fine-needle aspiration cytology followed by histology and IHC helped in making a definitive diagnosis of pseudolymphoma (lymphocytoma cutis), thereby differentiating it from lymphoma. This differentiation is crucial for early and accurate diagnosis, treatment, and prognosis of the patient.

Key words: Pseudolymphoma, Lymphoma, Cytology, Biopsy, Solitary nodule

Lymphocytoma cutis (LC), also termed as cutaneous lymphoid hyperplasia or Spiegler-Fendt sarcoid, results from various antigenic stimuli [1-4]. This entity is basically considered as a pseudolymphoma or collection of reactive polymorphic lymphoid cells where there is an accumulation of lymphocytes underneath the skin, thereby mimicking cutaneous lymphoma on histopathology [3-5]. Hence, it becomes important to establish the polymorphic nature of the lymphoid cells through immunohistochemistry (IHC) in these lesions.

Here, we describe the case of a 49-year-old-female patient who presented with a solitary nodular swelling at the back and was finally diagnosed with LC based on the histopathological and IHC evaluation. This diagnosis is of utmost importance keeping in view the management and prognosis of the patient.

CASE REPORT

A 49-year-old female presented to the hospital with a complaint of swelling on the left mid-back region. The swelling was mildly itchy, started 1 year back, and was slow growing. The patient did not have any history of any triggering agents such as an insect bite, drug intake, and vaccination.

On examination, a well-defined nodulocystic swelling was noted on the mid-back region toward the left side (Fig. 1a). The surface was smooth, however, the overlying skin showed mild discoloration. Clinically, the swelling was well-defined and considered benign in nature. General and systemic examination of the patient did not reveal any abnormality.

Complete blood count and other biochemical parameters were within normal limits. The patient was sent for fine-needle aspiration cytology (FNAC) with a provisional clinical diagnosis of epidermal inclusion cyst.

Fine-needle aspiration was done, and smears prepared were highly cellular, comprising many small to medium and round to oval-sized mononuclear cells with scant cytoplasm, round to oval nuclei, finely dispersed chromatin, and prominent nucleoli (Fig. 2a-b), also noted were elongated spindled and plasmacytoid looking cells with granular chromatin and a moderate amount of pale to dense cytoplasm seen singly scattered and few embedded in a pink matrix (Fig. 2c). The background showed a focal pink matrix, mixed inflammation, and occasional cystic macrophage admixed with blood. In view of cytological findings, a definitive diagnosis was difficult to be given, hence, the two possible differential diagnoses of adnexal neoplasm and the lymphoproliferative lesion were given. Subsequently, the excision of the lesion and its histopathological correlation was advised.

Complete excision of the lesion was done and sent for the histopathological examination. Fig. 1b shows the post-surgical picture of the lesion, with a scar mark. On gross examination, a single skin covered tissue measuring $2.8 \times 1.5 \times 1 \, \text{cm}$ was received (Fig. 1c). The external surface was partly skin covered and the cut surface showed gray-white areas along with focal yellow fatty areas. The specimen was cut into two bits and both the bits were sent for processing.

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Histopathological examination stained sections showed epidermis with its appendages and connective tissue stroma. The upper dermis was highly cellular with a clear zone of connective tissue separating the cells in the dermis from the epidermis (Fig. 3a). The cells were predominantly lymphocytes with few plasma cells, eosinophils, and histiocytes (Fig. 3b). These lymphoid cells were predominantly arranged in a dense sheet-like pattern; however, a focal follicular pattern was also noted (Fig. 3b). Although the upper dermis was majorly involved by these lymphoid cells, infiltration into the deeper dermis and subcutaneous tissue was also seen. Based on the clinical findings and histology, a final diagnosis of cutaneous lymphoid hyperplasia (LC) was made.

However, the important point was to differentiate this entity from cutaneous lymphoma, and hence, establishing the polyclonality of lymphocytes was required. IHC with LCA, CD3, and CD20 was done (Fig. 4a-d). The results showed positivity with all these antibodies, ruling out lymphoma and helped in achieving a final diagnosis of LC. The patient was given topical steroids and was advised to be on regular follow-up.

DISCUSSION

LC is also known as cutaneous pseudolymphoma, cutaneous lymphoid hyperplasia, or Spiegler-Fendt sarcoid [2-4]. It is a reactive process where there is a polyclonal T- or B-cell proliferation. It is a rare lesion presenting either as localized solitary lesion or in a disseminated form as multiple shiny plaques. This is usually noted in young adults with female preponderance. The disseminated form is also termed as military LC and is usually seen in elderly individuals [3-6]. This entity has a benign course and few other terminologies which have been used to describe it are sarcoma cutis, lymphadenosis benigna cutis, and actinic reticuloid.

Considering the etiological factors, most cases of LC are idiopathic [4-7]. Second, reactive responses may be responsible for its causation such as skin disease (allergic reactions and dermatitis); some viral infections (herpes simplex and molluscum contagiosum); tattoo dyes, trauma, vaccination, drug reaction (antibiotics, anticonvulsant, anti-TNFα drugs, etc.), or even with Borrelia burgdorferi infection [5-8]. Although the common site involved is skin; LC may be noted in the eyes, tongue, parotid gland, GIT, lung, kidney, and breast [6-9].

In our case, it was noted in a middle-aged female patient who presented with a discrete, slightly pigmented swelling with itching in the left mid-back region. There was no definite etiological factor associated with its causation in our patient.
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Once a diagnosis of LC is made, management involves the complete surgical excision of the lesion along with the application of steroids post-surgery; other methods include cryosurgery, superficial radiotherapy, psoralen and ultraviolet A (PUVA), hydroxychloroquine, and photodynamic therapy [7-12]. If a definite cause like Borrelia infection is noted, it should be treated specifically for it. In our patient, complete surgical excision was done, and the patient was given topical steroids for local application.

Establishing a definitive diagnosis of LC is very important not only in terms of management purpose but also in long-term follow-up of the patient. It is important to note that, even it is a reactive process, follow-up of the patients is very crucial as there have been very rare cases of LC evolving into cutaneous lymphoma.

CONCLUSION

The present case highlights the importance of a very simple investigative process, that is, FNAC in directing a path for a provisional diagnosis of a pseudolymphomatous condition. This initial diagnosis must be backed up by the histopathological correlation along with the establishment of polyclonality in these reactive lymphoid cells. The differentiation of LC from true lymphoma is further important in planning the management and follow-up of the patient.

REFERENCES


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