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Citation
Bartenstein, Diana W., Taylor M. Coe, Samantha C. Gordon, Alison M. Friedmann, Maryanne M. Senna, Cassandra M. Kelleher, Cristina R. Antonescu, Rosalynn M. Nazarian, and Elena B. Hawryluk. 2017. “Lipofibromatosis-like neural tumor: Case report of a unique infantile presentation.” JAAD Case Reports 4 (2): 185-188. doi:10.1016/j.jdcr.2017.09.004. http://dx.doi.org/10.1016/j.jdcr.2017.09.004.

Published Version
doi:10.1016/j.jdcr.2017.09.004

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CASE REPORT

Lipofibromatosis-like neural tumor: Case report of a unique infantile presentation

Diana W. Bartenstein, BA,a,b Taylor M. Coe, MD,c Samantha C. Gordon, MD,d Alison M. Friedmann, MD,e Maryanne M. Senna, MD,b Cassanne M. Kelleher, MD,c Cristina R. Antonescu, MD,f Rosalynn M. Nazarian, MD,g and Elena B. Hawryluk, MD, PhDb
Boston, Massachusetts, and New York, New York

A 14-month-old boy presented with a slow-growing, asymptomatic back plaque, which was biopsied and found to have S100 positivity, sparse CD34 staining, and no significant mitotic activity, nuclear pleomorphism, or necrosis; genetic workup found LMNA-NTRK1 gene fusion, overall consistent with lipofibromatosis-like neural tumor (LPF-NT). LPF-NT is rare, with 14 cases previously reported, and our patient is the first report of this diagnosis in infancy. This case report and literature review includes comparison of similar diagnoses including lipofibromatosis, low-grade malignant peripheral nerve sheath tumor, infantile fibrosarcoma, and dermatofibrosarcoma protuberans and serves to aid detection of LPF-NT presenting in pediatric patients by highlighting similarities and differences that should prompt consideration. LPF-NT shows locally aggressive behavior only and should not be confused with conditions that have potential for distant spread. However, case reports of metastasizing LMNA-NTRK1 tumors draw into question whether growths with this gene fusion exist on a spectrum of disease severity. Our patient was treated with wide local excision and has developed no complications or evidence of recurrence with 6 months of follow-up time. (J Am Acad Dermatol 2018;4:185-8.)

Key words: infantile mesenchymal tumor; lipofibromatosis-like neural tumor; pediatric skin tumor.

INTRODUCTION

Infantile mesenchymal tumors can range from benign to malignant, and proper diagnosis is crucial for patient management and counseling. Clinical appearance is not sufficient for diagnosis, and histopathology must be performed to determine tumor type. Lipofibromatosis-like neural tumor (LPF-NT) is a recently defined entity that commonly shows infiltrative growth and spindle cells arranged in streaming fascicles, which is similar to lipofibromatosis, but the tumor is distinguished by S100 protein reactivity and NTRK1 gene rearrangements. Clinically, differential diagnoses other than lipofibromatosis include peripheral nerve sheath tumor, dermatofibrosarcoma protuberans, infantile fibrosarcoma, hamartoma, myofibroma, vascular plaque such as arteriovenous malformation, congenital nevus with proliferative nodules, and melanoma.

Abbreviations used:
LPF: lipofibromatosis
LPF-NT: lipofibromatosis-like neural tumor
FISH: fluorescence in situ hybridization
REPORT OF A CASE

An otherwise healthy 14-month-old boy presented for evaluation of an asymptomatic truncal ‘birthmark’ that slowly grew and changed color. He was born at 40.5 weeks via cesarean section after an uncomplicated pregnancy. Results of lower back ultrasound scan and radiography performed on the fifth day of life, because of to a small tuft of hair over the lower lumbar spine, were unremarkable. His parents reported the tumor to be present at birth, and his pediatrician documented a quarter-sized plaque at his 2-month visit. At age 9 months, the tumor measured 1.5 × 2 cm with central clearing. The plaque was not pruritic, painful, or friable.

With presentation to the dermatology department at age 14 months, physical examination found a 3 × 3.5-cm violaceous, hyperpigmented, atrophic plaque on his left lower back (Fig 1). It contained 2 prominent erythematous firm nodules, the larger nodule measuring 1.5 cm. Magnetic resonance imaging found a well-defined 3.8- × 3.4- × 0.6-cm discoid mass involving the skin and subcutaneous tissue with predominant T2 hyperintensity, intermediate T1 signal, and a small internal fat signal component.

Histology found a deep dermal and subcutaneous spindled-cell neoplasm with fascicular growth and infiltration into the adipose tissue but no significant mitotic activity, nuclear pleomorphism, necrosis, or hemangiopericytoma-like vascular proliferation (Fig 2). Tumor cells displayed focal S100 protein reactivity and very focal to weak CD34 staining but were negative for desmin, smooth muscle actin, epithelial membrane antigen, and anaplastic lymphoma kinase. Cytoplasmic NTRK1 immunohistochemistry showed diffuse positive staining (Fig 3), and fluorescence in situ hybridization (FISH) studies with custom Bacterial Artificial Chromosomes (BAC) probes found NTRK1 breakapart. Further fusion FISH assays showed LMINA-NTRK1 fusion, whereas testing for ETV6 or EWSR1 gene rearrangements was negative. Overall findings were most consistent with a diagnosis of LPF-NT.

The tumor was excised with 1-cm margins. The patient had a temporary vacuum-assisted closure to allow for confirmation of clear margins by formalin-fixed and paraffin-embedded pathology evaluation, and the defect was repaired with bilateral V-Y advancement flaps. The patient has no complications or evidence of clinical recurrence with 8 months of follow-up. Three months after excision, magnetic resonance imaging showed no definitive evidence of residual or recurrent tumor; this finding serves as a postoperative imaging baseline.

DISCUSSION

Lipofibromatosis (LPF) tumors, first described in 2000 by Fetsch and colleagues,1 are rare,
slow-growing soft tissue masses composed of adipocytic and fibroblastic elements. In reviewing 827 fibromatous tumors from more than 30 years of pathology data, these authors identified 45 cases that they proposed to be classified as LPF. LPF is distinguished from infantile and juvenile fibromatosis by its predominantly adipocytic composition and differentiated from fibrous hamartoma of infancy by its lack of immature mesenchymal tissue. Although LPF tumors are likely to present with ill-defined margins and infiltrative growth, they are distinguished from malignant lesions by their absent-to-rare mitotic figures and mild atypia.

In 2016, Agaram and colleagues identified LPF-NT as a related tumor distinguished by S100 protein reactivity, which is indicative of neural differentiation. FISH analysis found that 10 of these 14 initially reported LPF-NT cases contained NTRK1 gene rearrangements, including TPR-NTRK1, TPM3-NTRK1, and most commonly, LMNA-NTRK1. In contrast, 25 typical LPF tumors showed S100 protein reactivity or NTRK1 gene abnormality.

Our case represents the first infantile LPF-NT, as previous patients presented between the ages of 4 and 38 years. This patient had tumor admixture with adipose tissue, which is a key feature of LPF, but relative paucity of fat compared to previously described cases resulted in his tumor appearing fibrous clinically. Identification and report of further LPF-NT cases is required to determine whether this particular presentation is associated with his young age, as infants are known to demonstrate different fat distributions than older patients, as opposed to those characteristic of LPF-NT. Nevertheless, we recommend that LPF-NT be included in the differential diagnosis for patients presenting with masses that are clinically concerning for fibrous as well as fatty tumors. Table I provides an overview of key clinical and histopathologic features that differentiate LPF-NT from related tumors. LPF-NT is associated with locally aggressive behavior only and should not be confused with tumors that distantly metastasize. However, 2 sarcoma cases with the same genetic fusion raise concern for potential disease evolution over time. Agaram and colleagues describe a 37-year-old patient whose tumor was stable for more than 20 years before growing rapidly. Histopathologic and genetic evaluation found S100 positivity, focal CD34 positivity, and LMNA-NTRK1 gene fusion, features that are characteristic for LPF-NT but not generally found in sarcoma, but the tumor was distinguished as malignant by high mitotic activity and necrosis. Wong and colleagues report a 7-month-old patient with a “purplish lesion on his right buttock” found to have LMNA-NTRK1 gene fusion and CD34 positivity, without report of S100 protein reactivity. This tumor was diagnosed as infantile fibrosarcoma because of its rapid growth, friability, vimentin positivity, and mitotic figures.

### Table I. Key features of LPF-NT and related tumors

| Feature                          | Lipofibromatosis-like neural tumor | Lipofibromatosis | Low-grade malignant peripheral nerve sheath tumor | Infantile fibrosarcoma | Dermatofibrosarcoma protuberans |
|----------------------------------|-----------------------------------|-----------------|-----------------------------------------------|------------------------|--------------------------------|
| Atypia                           | Low                               | Low             | Nuclear atypia present                        | High                   | Variable                       |
| Mitotic rate                     | Low                               | Low             | Low                                           | High                   | Variable                       |
| Immunohistochemistry             | S100 Positive                      | S100 Negative   | Positive                                      | S100 Negative          | Negative                       |
| CD34                             | Focal to multifocal positivity    | Variable        | Positive                                      | Variable               | Strongly positive              |
| SOX10                            | Reported NTRK1 rearrangement       | Not reported    | Variable                                      | Not reported            | Negative                       |
| Reported genetic mutations       | Local recurrence with incomplete excision | Local recurrence with incomplete excision | Potential for distant metastasis; 50% occur in patients with neurofibromatosis type I | Potential for distant metastasis | Potential for distant metastasis |
| Natural history                  |                                   |                 |                                               |                         |                                |

*One report of balanced translocation (4;9;6).^1^ One report of LMNA-NTRK1 fusion.º

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although *LMNA-NTRK1* gene fusion and CD34 positivity are not typical for this diagnosis. Both sarcoma patients went on to have pulmonary metastases, and the younger patient also suffered progression to his S5 vertebral body and acetabulum. Although these tumors were histopathologically distinct from LPF-NT, overlapping genetic features raise concern for the possibility that over time, LPF-NT may accumulate genetic abnormalities and increasingly aggressive clinical characteristics.

This case is reported to increase awareness of LPF-NT, as identification of additional cases will improve understanding of this rare dermatologic condition. Surgical margin guidelines are not available for these tumors, and LPF and LPF-NT have been found to show local recurrence when incompletely excised,\(^1,^8\) which prompted our use of 1-cm clinical margins and confirmation of histologic clear margins by formalin-fixed tissue before repair. In the absence of long-term follow-up data, we suggest early diagnosis, complete excision, and close follow-up for those affected.

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