Collagenous Fibroma (Desmoplastic Fibroblastoma): Different Localization. A Case Report

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Collagenous fibroma is an extremely rare benign soft tissue tumor which arises in the subcutaneous tissue or muscle. We report a rare case of a collagenous fibroma in a 44-year-old woman who presented with a mass in the anterior chest wall. Microscopically, the tumor was hypocellular and consisted of diffuse bland spindle to stellate–shaped cells embedded in a collagenous stroma. Immunohistochemically, the tumor cells were diffusely and strongly positive for vimentin. Some cells contained alpha smooth muscle actin. They were uniformly negative for S-100 protein and CD34. After surgical excision, no recurrence or metastasis was observed during a 6 month follow-up period. A discussion of the histopathological features and a review of the literature is given.

Key Words: Collagenous fibroma, Desmoplastic fibroblastoma, Soft tissue tumors, Immunohistochemistry

A distinctive fibrous soft tissue tumor, originally named as desmoplastic fibroblastoma, is a benign soft tissue lesion with low cellularity, abundant collagen and large fibroblasts with reactive appearance. This tumor was first proposed by Evans in 1995 as desmoplastic fibroma and later renamed as collagenous fibroma by Nielsen et al. Collagenous fibroma (CF) typically arises in the subcutaneous tissue or muscle. Dermal involvement is rare. There is a predilection for adult males. This entity has been reported in various locations, including the upper extremities, posterior neck, upper back, lower extremities, abdominal wall and hip. However, few cases have been reported in the head and neck region, including three cases in the oral cavity, one case in thyroid, and one case in parotid gland. In addition, CF (desmoplastic fibroblastoma) of pectoralis minor muscle has also been reported.

We describe the clinico-pathological features of CF which arose within pectoralis major and illustrate the difficulties that may be encountered in differentiating it from other benign soft tissue tumors.

CASE REPORT

A 44-year-old woman presented with a slow-growing mass on the upper chest. She had first observed the mass 1 year previously, which gradually enlarged. Her past medical history was unremarkable. The patient underwent an excisional biopsy. The tumor was easily removed in the surgical excision. Macroscopically, the excised specimen
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contained well demarcated subcutaneous mass measuring 3x2.5x1cm. The cut surface showed a yellowish-white solid tumor without necrosis.

The excised specimen was fixed in 10% neutral formalin and embedded in paraffin. Sections were stained with hematoxylin–eosin. Histologically, the tumor was well-circumscribed, hypocellular and characterized by a proliferation of large, stellate and spindle cells set in a densely fibromyxoid stroma (Figure 1,2). Small areas of the tumor had keloid-like thick bands of collagen, dispersed between the neoplastic cells. Neither cellular atypia nor mitotic figures were observed. There was no tumor necrosis. Blood vessels were scarce and inconspicuous. Immunohistochemically, spindle-to stellate-shaped tumor cells showed diffuse and intense reactivity for vimentin (Figure 3). Tumor cells were focally positive for muscle-specific actin. Immunostaining for desmin and S-100 protein were negative.

**Figure 1.** Low-power photomicrograph of the tumor demonstrating a well-circumscribed fibrous tumor (H&E, original magnification x40)

**DISCUSSION:**

CF (desmoplastic fibroblastoma) is a rare benign soft tissue tumor morphologically characterized by a hypocellular proliferation of stellate and spindle-shaped fibroblasts separated by a collagenous or myxocollagenous matrix. The tumor usually occurs in the subcutaneous tissue but approximately 25% involve skeletal muscle. The lesion presents as a firm, well circumscribed, painless mass of long duration and is commonly located in the subcutaneous tissue or skeletal muscle in adults. The tumors range in size from 1 to 20 cm. The tumors are firm and tender. There is a predilection for adult males. We believe that the case described here presents similar characteristics to the one described by Evans. The tumor has a wide anatomical distribution including arms, shoulders, feet or ankles, legs and hands. Case reports in the literature indicate additional sites including thyroid and parotid glands, palate, skin, spinal epidural region and pectoralis minor (Table 1).

**Figure 2.** Photomicrograph illustrates hypocellular lesion, with widely spaced bland cells embedded in a fibromyxoid stroma (H&E, original magnification x200)

**Figure 3.** The cells in the tumor showed strong expression of vimentin (Immunohistochemical stain, original magnification, x100)

The differential diagnosis includes a variety of benign and low-grade lesions, predominantly fibrous lesions. CF should be differentiated from fibromatosis, which has a high risk of local recurrence when simple local excision is performed. Fibromatosis is more cellular and shows short fascicular arrangement of tumor cells and shows greater infiltration at the periphery of the lesion than CF. By contrast, the nuclei in CFs exhibit no particular alignment or polarity and the tumor is hypocellular overall.
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Table 1: Clinical data on 8 patients with collagenous fibroma

<table>
<thead>
<tr>
<th>Source/case no</th>
<th>Year of publication</th>
<th>Age, y/Sex</th>
<th>Localization</th>
<th>Tumor Size</th>
<th>Follow up</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shimayama T et al. Case 1</td>
<td>2005</td>
<td>48/F</td>
<td>Palate</td>
<td>6 cm</td>
<td>No evidence of disease/4 year</td>
</tr>
<tr>
<td>Hao S et al. Case 2</td>
<td>2003</td>
<td>45/M</td>
<td>Pectoralis minor muscle</td>
<td>9.5 cm</td>
<td>NP</td>
</tr>
<tr>
<td>Mesquita et al Case 3</td>
<td>2001</td>
<td>35/F</td>
<td>Palate</td>
<td>5 cm</td>
<td>No evidence of disease/6-months</td>
</tr>
<tr>
<td>Wilson et al Case 4</td>
<td>2001</td>
<td>88/M</td>
<td>Thyroid gland</td>
<td>4 cm</td>
<td>No evidence of disease/a few months</td>
</tr>
<tr>
<td>Kamimura et al Case 5</td>
<td>2000</td>
<td>20/F</td>
<td>Thoracic spinal epidura</td>
<td>NP</td>
<td>No evidence of disease/6 years</td>
</tr>
<tr>
<td>Ide F et al Case 6</td>
<td>1999</td>
<td>50/F</td>
<td>Parotid gland</td>
<td>5 cm</td>
<td>No evidence of disease/2-months</td>
</tr>
<tr>
<td>Weisberg et al Case 7</td>
<td>1999</td>
<td>67/M</td>
<td>Forehead</td>
<td>2 cm</td>
<td>No evidence of disease/2-months</td>
</tr>
<tr>
<td>Present case Case 8</td>
<td>2006</td>
<td>44/F</td>
<td>Pectoralis major muscle</td>
<td>3 cm</td>
<td>No evidence of disease/6 months</td>
</tr>
</tbody>
</table>

NP indicates not provided

CFs may grossly mimic neurofibromas because of their well-circumscribed appearance, but they are generally negative for S100 protein immunohistochemically. Occasional tumors may show faint positivity, but always to a lesser degree than is seen in neurofibromas. In contrast to CF, neurofibroma is composed of cells with a wavy configuration which are often dispersed in a myxocollagenous stroma.2,5

Calcifying fibrous pseudotumor, which affects children and young adults, is distinguished from CF by psammomatous calcifications and by the presence of lymphoplasmacytic infiltrate.13 Calcifications and lymphoplasmacytic infiltration were not present in the current case.

Solitary fibrous tumor is a rare mesenchymal neoplasm, primarily described in visceral pleura. Histopathologically, solitary fibrous tumors are well-circumscribed or encapsulated and composed of bland spindle cells with some hypercellular areas and myxoid to collagenized background. Areas with a hemangiopericytomatous vascular pattern and focally increased cellularity are present in some cases. Solitary fibrous tumors are much more cellular than CFs and invariably express CD34.14

Elastofibroma almost always occurs in the lower subscapular area and is characterized by elastic fibers, some of which show branching. These fibers, that are absent in CF, are highlighted with an elastin stain.4

Low-grade fibromyxoid sarcoma is typically more cellular than CF. Its characteristic feature is the whorling growth pattern, often seen at the transition zone. In contrast to CF, its stroma may be hypocellular or hypercellular with areas showing myxoid change.15

Nodular fasciitis of long duration may resemble this lesion but usually is more cellular than CF. Compared to CF, nodular fasciitis usually demonstrates storiform areas, interconnecting bundles, myxoid areas or focal mucinous microcyst formation and numerous slit-like spaces. These spaces often contain erythrocytes, along with scattered lymphocytes and macrophages.3,6,12

Ultrastructural studies on CF have yielded fibroblastic or myofibroblastic lineage and they are diploid on flow cytometry.13 Seiot et al. have recently published two such tumors with aberrations of 11q12, similar to those described for fibroma of the tendon sheath.16 However, it is still unclear if this lesion is a reactive process or a true neoplasm.

In summary, awareness and recognition of CF as a distinct pathologic entity should be emphasized due to its tendency to mimic others. In the present case, the tumor had a benign histology and surgical excision was curative. Histopathologic examination and immunohistochemical analysis may help in the diagnosis of CF and the treatment of choice is en bloc excision. Prognosis is excellent and no recurrence is expected.

REFERENCES

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