Nodular fasciitis (NF) is a benign reactive or inflammatory condition of mesenchymal fibroblasts. Rapid growth and histological features make nodular fasciitis a mimicker of malignancies such as sarcomas, however, nodular fasciitis is a benign lesion and does not metastasize. Although the etiology of nodular fasciitis is uncertain, histopathologically, it bears a close resemblance to organizing granulation tissue, and myofibroblastic proliferation may be initiated by a local injury or local inflammatory process, which supports a reactive proliferation theory triggered by trauma. Surgical excision is recommended for diagnosis and treatment to exclude malignancy. We are hereby reporting a case of nodular fasciitis on the front surface of the neck, 86 years old patient. Tumor grew rapidly and reached 10 cm., and which was clinically suspected as a malignancy located. An ultrasound examination was performed. Based on the clinical and ultrasound data, it was decided to perform an extended biopsy to establish the final diagnosis it. Tumor has been removed as a solid mass. The material was sent for histopathological examination. Based on the history, clinical picture, and histopathological examination, the lesion was diagnosed as nodular fasciitis. After surgery patient expressed satisfaction with the result of treatment. In cases of nodular fasciitis, proactive efforts are needed to perform histopathologic evaluation and treatment because nodular fasciitis tends to grow rapidly.
being the least common. In adults, the upper extremities are the most commonly affected site [12].

There is no standard treatment for nodular fasciitis, treatment options for nodular fasciitis depend in part on the size and location of the tumor. Treatment is usually surgical excision; however, large infiltrative lesions may be difficult to manage. Surgical excision is recommended for diagnosis and treatment to exclude malignancy. Laser treatment can be effective in decreasing the size of the lesions through tissue contraction. A carbon dioxide (CO2) laser used in a pinhole pattern is a treatment option for tumors on the face or another area [13]. For the nonsurgical treatment of nodular fasciitis on the face, triamcinolone intralesional injection (TA ILI) can be used on the basis of the fact that, nodular fasciitis is a benign reactive or inflammatory condition of mesenchymal fibroblasts [14]. If the nodular fasciitis is large or on the face, a corticosteroid injection at the site may help resolve the tumor. We describe a case of nodular fasciitis on the front surface neck that demonstrated partial spontaneous regression and treated by using surgical method.

Case Report

Patient, 86 years old, was taken to the Dept. of Oral and Maxillofacial Surgery of 1-st Yerevan State University Hospital Complex “Heratsi” by an ambulance team on 09.08.2018, with complaints of bleeding from the neoplasm located on the front surface of the neck. According to the patient, the tumor first noticed about 3–4 months ago. Tumor grew rapidly and reached 9–10 cm. History indicated the thyroid gland disease, about which the patient took a medication every day (presumably euthyrox). During external examination, a palpable dense consistency neoplasm was observed, covered with a dense hemorrhagic crust about 2–3 cm, blackish in color (Figure 1). There was a slight bleeding from the periphery, the blood was dark, stagnant. During palpation, the tumor was painless, partially moveable. The color of the overlying skin was not changed, with the underlying tissues not soldered. Submandibular lymph nodes were not palpated.

An ultrasound examination was performed, on the basis of which it was revealed that the thyroid gland is diffusely altered, there are nodes in the right lobe of the thyroid gland. In the area of the right lobe of the thyroid gland, a massive heterogeneous well vascularized neoplasm of about 10–13 cm was found with clear boundaries, located extracapsularly (Figures 2 & 3). Based on the above clinical and ultrasound data, it was decided to perform an extended biopsy to establish the final diagnosis. During the preoperative examinations of the associated diseases, a diffuse nodular goiter, chronic heart failure of the II-degree, chronic obstructive pulmonary disease, and asthma was detected. Only after the general somatic stabilization of the patient was an operation planned.

Figures 2 & 3: An ultrasound examination in the area of the right lobe of the thyroid gland, a massive heterogeneous well vascularized neoplasm of about 10-13 cm was found with clear boundaries, located extracapsularly.
and retreating from the palpable tumor boundaries by 2 cm. A skin flap was cut, including a hemorrhagic crust. In a blunt way, the tumor was mobilized and exfoliated from the surrounding tissues. All the vessels involved were ligated and hemostasis was achieved. Tumor has been removed as a solid mass. After antiseptic preparation, hemostasis was achieved and, the wound was sutured layer by layer (Figures 4-8). The material was sent for histopathological examination. Macroscopically: there was 4,0x4,0x2,5 cm dense consistency neoplasm, with smooth whitish edges. In the section there was a cavity which extended from the skin to the base of the neoplasm, filled with a homogeneous pink mass, with the presence of the surrounding yellow-whitish tissues.

Figure 7: Clinical appearance after removal of nodular fasciitis.

Figures 8: Nodular lesion after being surgically removed: there was 4,0x4,0x2,5 cm dense consistency.

Figure 9: Accumulation of hemosiderin.

Microscopically: peripheral hypercellular zones were observed on sections, with S and C shape solid areas of rounded and elongated moderately pleomorphic cells located in different directions, with hyperchromic and weakly cytoplasmatic nucleus, having vesicular chromatin, pronounced nucleoli. There were also hypocellularity areas with the presence of keloid type collagen, extensive areas of hyalinosis, sometimes in the middle there were cavities filled with blood, diapedesing hemorrhage, accumulation of hemosiderin and hematomidin, mixed inflammatory cellular infiltration, including giant cells, focal infiltration of lymphocytes, hyperemic vessels and a lot of vascular fissures. In one of the sections, an opening of the cystic cavity was observed, with blood content, skin manifestation, tissue detritus and leukocytes. In other sections, epidermal hyper keratinization, acanthosis, dermal fibrosis, and ganglion-type cell accumulation were found. There was also a lymphoplasmacytic focal infiltration of muscle tissue. This histological picture is most comparable to the diagnosis of nodular fasciitis (Figures 9-11). Based on the history, clinical picture, and histopathological examination, the lesion was diagnosed as nodular fasciitis.

Figure 10: Foci of fibrosis and hyalinosis.

Figure 11: Hyper cellular areas.

Discussion

Nodular fascitis is a rare benign soft tissue tumor, most commonly afflicting the soft tissues of upper extremity followed by trunk, head, and neck [15]. Oro-facial lesions involve the skin of the face, parotid gland, buccal mucosa, labial mucosa, and tongue[16]. Lesions are small, solitary, and are commonly located on extremities occasionally on the trunk and infrequently on the head and neck. Nodular fascitis is also known as pseudo sarcomatous fasciitis, pseudosarcomatous fibromatosis and infiltrative fasciitis. The etiology is still unknown. It is considered to occur due to unusual proliferation of myofibroblasts triggered by local injury or inflammatory process [16].

Previously nodular fascitis was thought to be reactive in nature. Infection and hormonal influences have also been suggested as initiating factors [2, 17, 18]. However, genetic research proves that nodular fascitis undergoes a clonal expansion process [8]. It is commonly seen in the 40-50-year age group with almost equal distribution amongst both genders. It almost always poses a considerable diagnostic challenge. However, fuzzy cytoplasmic borders, multiple fragile cytoplasmic processes, and prominent nucleoli favor a diagnosis of nodular fascitis [19, 20]. Nodular fascitis can be differentiated from other facial mesenchymal tumors, including sarcomatoid carcinoma, fibrosarcoma, leiomyosarcoma, and neurofibroma, based on histology, molecular biology, and immunohistochemistry [21].
In cases of nodular fasciitis, proactive efforts are needed to perform histopathologic evaluation and treatment. The histological appearance of nodular fasciitis can be described as deceptive, due to its non-specific and, even though well demarcated, infiltrative nature. The lesion also has a rich vascularity and is mitotically active. This rapid growth and histological features make nodular fasciitis a mimic of malignancies such as sarcomas [22, 23]. A nodular fasciitis should be surgically excised to obtain complete cure as the chances of local recurrence are almost nil (<1%) [24].

We report a rare case of nodular fasciitis of neck in an 83-year-old lady. Clinical, laboratory, radiological methods were used in the examination of patients. Aspiration of the swelling did not show any contents. To establish a final diagnosis, we conducted a histological examination. Tumor has been removed as a solid mass. The material was sent for histopathological examination and the lesion was diagnosed as nodular fasciitis. After surgery patient expressed satisfaction with the treatment and with good long-term results. In cases of nodular fasciitis, proactive efforts are needed to perform histopathologic evaluation and treatment because nodular fasciitis tends to grow rapidly.

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Conflicts of Interest

None.

Protection of Human and Animal Subjects

The authors declare that the procedures followed were in accordance with the regulations of the responsible Clinical Research Ethics Committee Yerevan State Medical University after M. Heratsi (protocol N23.11.2018) and in accordance with those of the World Medical Association and the Helsinki Declaration.

Consent Statement

Written informed consent was obtained from the patient for publication of this case report and accompanying images.

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