Fibro Adipose Vascular Anomaly: A Rare and Often Misdiagnosed Entity

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Introduction

Fibro adipose vascular anomaly (FAVA) is a type of vascular malformation which is classified under the International Society for the Study of Vascular Anomalies (ISSVA) in the provisionally unclassified vascular malformation group.¹,² FAVA tends to occur more in young females and lower extremities. Clinical, imaging findings, and histopathological correlations are needed for better understanding of the disease and to differentiate it from other vascular malformations. No single treatment is available for FAVA. A combination of sclerotherapy, intralesional steroids, cryotherapy, or ablation therapy can be tried. If there is restriction in movement surgical cut down/resection can be done.

Case Report

A 14-year-old female referred from the orthopaedics department for the evaluation of swelling over the posterior aspect of the left lower limb in suspicion of venous malformation (VM). The patient had swelling over the calf and dorsal aspect of the left foot for the past 10 years which was insidious in onset, gradually progressive, and associated with severe constant pain at the local site with restriction in movement. No history of trauma, fever, or similar family history was observed. There was past history of sclerotherapy and Achilles tendon release 2 to 3 years back, however, there was no much improvement. Routine blood investigations including coagulation profile, erythrocyte sedimentation rate, and C-reactive protein were within normal limits. On clinical examination, the left lower limb was thinner compared with the right (Fig. 1). There were multiple areas of soft swelling over the posterior aspect of the left leg and lateral aspect of the dorsum of the left foot. Palpation of the local site had tenderness and mild increase in temperature. There were restricted dorsiflexion movements and the tendency of toe walking. Motor power in the lower limb and the overlying skin was normal. The scar mark of the tendon release

Abstract

Fibro adipose vascular anomaly (FAVA) is a rare type of vascular malformation with distinct clinical features. The authors here discussed the clinical, imaging, differential diagnosis, histopathological features, and treatment options of FAVA along with an illustrative case. It is important to know about this uncommon entity as this can be misdiagnosed due to the overlapping clinical features with other common entities. It is a benign condition with no proven malignant potential. There are no guidelines regarding the best treatment option.

Keywords

Fibro adipose vascular anomaly vascular malformation venous malformation

Case Report

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procedure was also noted along the posterior part of the ankle.

The patient was extensively investigated with all imaging modalities during the course of her disease without a definitive diagnosis. Radiograph of left leg revealed few abnormal soft tissue radiopacity over the posterior aspect with dense amorphous calcification (►Fig. 2A, B). Mild thinning of the fibula is seen due to long-standing pressure remodeling from the swelling. Ultrasound of the local site showed heterogeneous predominantly echogenic well-circumscribed non-compressible soft tissue lesions in the intramuscular and subcutaneous plane in the posterior aspect of the left leg (►Fig. 2C, D). There were few echogenic foci noted giving posterior acoustic shadowing suggestive of calcifications (►Fig. 2E). Few anechoic serpiginous channels were also noted with minimal color filling and low-velocity venous type of waveform on spectral imaging (►Fig. 2F).

Non-contrast computed tomography (NCCT) revealed well-defined lobulated heterogenous hyperdense mass lesions in the intramuscular and subcutaneous plane. Areas of fat attenuation replacing the lateral and medial head of gastrocnemius, soleus, and plantaris muscles were seen. Multiple clustered areas of heterogenous calcification were seen within the hyperdense lesions (►Fig. 3A, B).

Magnetic resonance imaging (MRI) of the left lower limb showed well-defined lobulated predominantly T2W and short tau inversion recovery (STIR) hyperintense lesions in the subcutaneous and intramuscular plane. There was fatty replacement of the lateral and medial head of gastrocnemius, soleus, and plantaris muscles. Multiple hypointense areas were also noted within the lesion corresponding to calcifications. There are grouped muscle atrophy of the posterior compartment as compared with the right side. On post gadolinium, T1-weighted sequence dilated dysplastic venous channels were seen in the lesion. The lesions showed moderate enhancement in the late arterial phase which was persistent in late venous phase too (►Fig. 3C–F). Based on the clinical and imaging features, FAVA was suggested.

The ultrasound-guided biopsy was performed using an 18-gauge needle. Histopathological examination showed fibro-collagenous tissue cores with fibro adipose tissue and containing several large and few small-caliber vessels (►Fig. 4). Anastomosing retiform to ectatic vascular spaces

Fig. 1 Clinical image of patient anterior (A) and posterior view (B), showing relative atrophy of left leg (asterisk). The lesion is predominantly involving the posterior aspect of the leg, also extending to the dorsum of the foot (black arrow). Scar of previous tendo-achilles release is seen (white arrow).

Fig. 2 Radiograph of left leg anteroposterior (A) and lateral (B) view showing lobulated soft tissue lesion (black arrow) in the calf region with dense amorphous calcification (asterisk). Thinning and scalloping of fibula without obvious bony destruction is seen. Ultrasound images (C–E) showing lobulated predominantly echogenic lesions with no compressible venous spaces. Presence of calcification (white arrow) with posterior acoustic shadowing seen within the lesion. Color Doppler (F) image showing minimal venous flow.
were lined by flattened endothelium. Aggregates of lymphocytes, perivascular mononuclear cell infiltrate, and hemosiderin pigment were noted. The final diagnosis of FAVA was confirmed.

The patient was explained about the disease and kept on regular follow-up. She was advised for symptomatic treatment in the form of local alcohol injection, cryotherapy, and surgical resection.

Discussion

Alomari et al in 2014, first time described a clinical condition named FAVA, with distinct clinical, radiological, and histopathologic features. FAVA consist of abnormal fibrofatty infiltration of muscles, venous abnormality in the form of phlebectasia, contracture of the affected extremity, and continuous pain. Histologically many reports mentioned similar pathological lesions termed as intramuscular hemorrhage causing contracture, leading to toe walking and equinus deformity.

International Society for the Study of Vascular Anomalies (ISSVA) classified vascular malformation into vascular tumors (neoplastic), vascular malformations (non-neoplastic), and unclassified anomalies. In the 2018 revision of ISSVA classification, FAVA was first time included in the provisionally unclassified vascular anomalies category. This category also includes other conditions like intramuscular hemangioma, angiokeratoma sinusoidal hemangioma, and acral arteriovenous tumor, etc.

Various published literature on FAVA with patient’s age, sex, site of involvement, symptoms, referring diagnosis, and treatment are compiled in Table 1.

FAVA is usually sporadic and most commonly caused by somatic mutation involving PIKC3A (phosphatidylinositol-4,5-bisphosphate 3-kinase) gene. The same gene is seen in most cases of isolated lymphatic and veno-lymphatic vascular...
Table 1 Demographic and clinical features of FAVA in published literature

<table>
<thead>
<tr>
<th>S. no</th>
<th>Authors</th>
<th>Year</th>
<th>Total number of patients (Lesions)</th>
<th>Male</th>
<th>Females</th>
<th>Age (Years)</th>
<th>Lower extremity/ Upper extremity/Trunk</th>
<th>Predominant site</th>
<th>Symptoms</th>
<th>Referring diagnosis</th>
<th>Management</th>
</tr>
</thead>
</table>
| 1.    | Alomari et al\(^2\)   | 2014 | 18                                | 5    | 13      | 0–28        | 15/3/0                                  | Calf: 12         | Pain: 15  
Limited dorsi flexion: 8  
Superficial phlebectasia: 2  
Disuse atrophy: 3  
Cutaneous lymphatic vesicles: 1 | VM-7  
Hemangioma-6  
AVM-5 | Surgery-8 |
| 2.    | Fernandez et al\(^4\) | 2014 | 1                                 | 1    | 0       | 10          | 1/0/0                                  | Calf             | Pain and Equinus deformity                                                | Sclero + Surgery    |                    |
| 3.    | Shaikh et al\(^8\)    | 2016 | 20 (26)                           | 6    | 14      | Mean age at first procedure 15.8 y, range 8–30 y | 24/1/1                                  | Calf: 8          | Pain: 26  
Functional restrictions: 21  
Skin hyperesthesia: 14 | Prior Surg + Cryo-7  
Prior Sclero + Cryo-9  
Prior Steroid injection + Cryo-6  
Additional sclero during cryoablation-2  
Surgery post-cryoablation-2 |                    |
| 4.    | Erickson et al\(^9\)  | 2017 | 2                                 | 2    | 0       | 7–8         | 1/1/0                                  | Foot: 1          | Pain: 2  
Deformity: 2  
Functional impairment: 1 | Sclero + Lumbar sympathectomy + Sirolimus-1  
Surg + Sirolimus-1 |                    |
Contracture: 4  
Swelling: 1 | VM-7  
VaM-6  
Hemangioma 3  
AVM-1  
Cav Lym-1  
FAVA-1 | Sclero + Cryo-1  
Cryo-4 |                    |
| 6.    | Cheung et al\(^11\)   | 2020 | 19 (28)                           | 4    | 15      | 0–14        | 0/28/0                                 | Forearm or wrist:12  
Hand or digit/thumb: 10  
Axilla, arm, or elbow: 6 | Pain: 15  
Contracture: 13  
Swelling: 9 | VM-7  
VaM-6  
Hemangioma 3  
AVM-1  
Cav Lym-1  
FAVA-1 | None-3  
Embo-10  
Surg-10  
Sclero + Surg-3  
Surg + Sclero-1  
Sclero + Cryo-1 |                    |
| 7.    | Amarneh and Shaikh\(^12\)| 2020| 38                                | 7    | 31      | 0–30 (mean 12) | 36/1/1                                  | Calf: 22         | Pain: 38  
Functional impairment: 38  
Swelling: 14  
Contracture: 8  
Paraesthesia: 10 | VM-10/37  
VaM-9/37  
Hemangioma-3/37  
AVM-3/37  
VL.M-2/37  
LM-1/37  
CVLM-1/37  
Soft tissue tumor-1/37  
muscle strain-1/37  
FAVA-6/37 |                    |

(Continued)
malformations. Other pathology caused by the same genetic mutation includes Klippel-Trenaunay syndrome, megalencephaly-capillary malformation-polymicrogyria (MCAP), and CLOVES (congenital lipomatous overgrowth, vascular malformations, epidermal nevi, and skeletal anomalies) syndrome. 12,16,17

FAVA usually occurs in young adults with age group between 1 and 30 years. It is more common in females with male to female ratio of 1:4. Lower extremities are more commonly involved than upper extremities. Calf muscles are most commonly involved followed by the thigh. 3,12

Since FAVA is a new clinical entity and has overlapping features with other conditions like VM, vascular malformation, intramuscular hemangioma, and soft tissue tumor, diagnosis is often delayed and missed. 12

FAVA is a distinct entity characterized by abnormal fibro-fatty masses and infiltration in intramuscular as well as subcutaneous plane. There is presence of abnormal venous channels in form of phlebectasia within masses. The fibrotic process leads to the contracture of involved muscles resulting in restriction of movement. Since calf and gastrocnemius are most commonly involved region the fibrotic process leads to equinus deformity and toe walking. 3,4

The lesion is also associated with continuous pain which is multifac torial; caused by muscular contracture, neurogenic infiltration, and thrombophlebitis of VM or phlebectasia. 3,4,12

FAVA can be divided into focal mass-like lesion, focal infiltrative, or diffuse infiltrative type. 12 The patient usually presents with long-standing soft non-compressible swelling with restricted movement due to contracture and constant pain. There can be associated skin changes in some cases like hypopigmentation.

Table 1 (Continued)

<table>
<thead>
<tr>
<th>S. no</th>
<th>Authors</th>
<th>Year</th>
<th>Total number of patients (lesions)</th>
<th>Male</th>
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<th>Referring diagnosis</th>
<th>Management</th>
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<tbody>
<tr>
<td>8.</td>
<td>López et al 13</td>
<td>2020</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>24</td>
<td>0/1/0</td>
<td>Forearm</td>
<td>Pain, Functional impairment</td>
<td>VM</td>
<td>Prior Sclero + Surg</td>
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<tr>
<td>10.</td>
<td>Ferreira et al 15</td>
<td>2020</td>
<td>1</td>
<td>1</td>
<td>9</td>
<td>9</td>
<td>1/0/0</td>
<td>Calf</td>
<td>Pain</td>
<td>FAVA</td>
<td>Surgery</td>
</tr>
<tr>
<td>11.</td>
<td>Our case</td>
<td>2020</td>
<td>1</td>
<td>1</td>
<td>8</td>
<td>8</td>
<td>1/0/0</td>
<td>Calf</td>
<td>Pain, contracture</td>
<td>VM</td>
<td>Sclero + Surg</td>
</tr>
</tbody>
</table>

Abbreviations: AVM, arteriovenous malformation; Cryo, cryotherapy; CVLM, capillary veno-lymphatic malformation; Embo, embolization; FAVA, fibroadipose vascular anomaly; Sclero, sclerotherapy; Surg, surgery; VaM, vascular malformation; VLVM, veno-lymphatic malformation; VM, venous malformation.
There is no single definitive treatment of FAVA. The image-guided treatment option is local sclerotherapy of venous components. Intralesional steroid and alcohol injection, local cryotherapy, and nerve blocks for pain relief. Surgical excision is an option with procedure-related morbidity as most of these lesions are intramuscular. Debunking of mass with contracture release to improve deformity, movement restriction, and pain relief in selective cases. Sirolimus is a drug also used in the treatment of FAVA but not approved in less than 18 years. To summarize, treatment is mostly symptomatic to address patient predominant clinical problems.8,9,14

Conclusion

FAVA is an uncommon but distinct clinical entity with typical presentation in young females and mostly involves lower limb. The classical presentation is long-standing swelling with constant pain, contracture, restricted movement, and deformity. However, because of the rarity of cases, recently described condition, overlapping features with other common entities, and lesser awareness, it is often misdiagnosed. The multimodality approach and patient education play a major role in management.

Declaration of Patient Consent

The authors certify that they have obtained all appropriate patient consent forms. In the form the patient(s) has/have given his/her/their consent for his/her/their images and other clinical information to be reported in the journal. The patients understand that their names and initials will not be published, and due efforts will be made to conceal their identity, but anonymity cannot be guaranteed.

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Date

Conflict of Interest

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References