Granular Cell Tumor of the Scrotum in a Black African: Case Report of a Very Rare Condition

Edoise M. Isiwele1*, Theophilus Ugbem2, Terence A. Azeke1

1Department of Urology, University of Calabar Teaching Hospital, Nigeria
2Department of Histopathology, University of Calabar Teaching Hospital, Nigeria
3Department of Anatomic Pathology, Irrua Specialist Teaching Hospital, Nigeria

*Corresponding author: Isiwele EM, Department of Urology, University of Calabar Teaching Hospital, Nigeria. Tel: +234-8039677046; Email: eddieisiwele@yahoo.com


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Abstract

Granular Cell Tumor (GCT) of the scrotum is very rare. Generally, granular cell tumors are more common in Blacks. However, this may not be the case with scrotal granular cell tumors. We report the first case seen in a native Black African patient to our knowledge. The 31-year old man presented with a nodular scrotal mass found on histology to be GCT. On immunohistochemistry it was found to have strong reactivity to S-100 and Vimentin. A strong index of suspicion is needed in making a clinical diagnosis, as the lesion can easily be missed.

Keywords: Black African; Granular Cell Tumor (GCT); Immunohistochemistry; Scrotum; Vimentin

Case Description

A 31 year old male Nigerian presented to the Urology Clinic of University of Calabar Teaching Hospital, Nigeria with history of a small and firm growth on the underside of his scrotum of 5 months’ duration. He had not noticed any appreciable change in its size since it was first noticed. It was not associated with pain or pruritus and there were no similar lesions elsewhere. There was no history of preceding trauma or infection at the region affected. On physical examination, there was a dark colored, non-tender, nodular mass, 0.6 cm in diameter, on the lower aspect of the left hemi-scrotum with well-defined edges. Other aspects of scrotal examination were essentially normal. Results of routine investigations were essentially normal with scrotal ultrasound revealing a rounded homogenous solid mass 0.54 × 0.44 cm on the left side of the median raphe with a suspicion of neuroma. The mass was subsequently excised with a rim of about 1 cm of normal tissue. Histological analysis showed irregular groups of polyhedral cells with indistinct borders and abundant eosinophilic granules in the cytoplasm and relatively small hyperchromatic nuclei that are centrally placed (Figure 1). Immunohistochemistry revealed the tumor to be strongly positive both for S-100 and Vimentin (Figure 2). Post-operative condition was satisfactory and follow up 12 months’ post-excision has shown no recurrence.

Introduction

Granular Cell Tumor (GCT) is a tumor commonly affecting the head and neck but which can affect any part of the body. They are soft tissue tumors thought to be of neural origin [1,2]. Since 1926 when Abrikossoff first described the condition, some few hundred cases have been reported in literature [3,4]. Most cases are benign with less than 1-2% of cases being reported to be malignant. While benign cases are usually small sized and slow growing, the malignant varieties rapidly grow and assume large proportions and also rapidly metastasize [5-7]. Granular cell tumor of the scrotum is very rare with only 11 cases reported so far in literature. In Nigeria, nine cases of granular cell tumor of the breast seen over a 10-year period were reported by Imam, et al. [8] in their study in Kano while a case of granular cell tumor of the mons pubis was reported by Aniebue and Olusina [9] in Enugu in 2009. No case of scrotal GCT has been reported in literature in the country, to our knowledge. This therefore represents the first case from Nigeria and indeed Africa being reported in literature.
Discussion

Granular Cell Tumor (GCT) has variously been referred to as Abrikosoff’s tumor, granular cell myoblastoma, granular cell neuroma, granular cell neurofibroma or granular cell schwannoma. It is a soft tissue neoplasm of neural origin that is distinct from neurofibromas, schwannomas or muscle tumors [10,11]. Scrotal GCT was first reported in literature by Ha and associates [12] in 1994 and since then this is the twelfth case being reported and the first in Africa (Table 1). Previous reports indicate all patients to have presented with painless nodules [11-13], with one case reported to have been ulcerated at presentation [14]. Nodule was noticed to have been present in various cases for periods ranging from 3 weeks to 35 years with no malignant case recorded so far.
### Table 1: Summary of scrotal granular cell tumors reported so far in literature. (Adapted from the work by Chen and colleagues [10])

<table>
<thead>
<tr>
<th>Case</th>
<th>Age (years)</th>
<th>Number of Tumors</th>
<th>Duration before presentation</th>
<th>Size (cm)</th>
<th>Presentation</th>
<th>Treatment</th>
<th>Recurrence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ha, et al. [12]</td>
<td>40</td>
<td>1</td>
<td>N/A</td>
<td>5</td>
<td>Nodule</td>
<td>Excision</td>
<td>N/A</td>
</tr>
<tr>
<td>Bryant [15]</td>
<td>36</td>
<td>1</td>
<td>1 year</td>
<td>0.4</td>
<td>Nodule</td>
<td>Excision</td>
<td>Nil at 6 months</td>
</tr>
<tr>
<td>Altman, et al. [16]</td>
<td>55</td>
<td>2</td>
<td>N/A</td>
<td>0.7, 1.1</td>
<td>Nodule</td>
<td>Excision</td>
<td>Nil at 22 months</td>
</tr>
<tr>
<td>Medina Perez (1999) [14]</td>
<td>19</td>
<td>1</td>
<td>Months</td>
<td>2</td>
<td>Skin ulcer</td>
<td>Excision</td>
<td>Nil at 24 months</td>
</tr>
<tr>
<td>Mendelez Lopez, et al. [17]</td>
<td>38</td>
<td>1</td>
<td>2 years</td>
<td>1</td>
<td>Nodule</td>
<td>Excision</td>
<td>N/A</td>
</tr>
<tr>
<td>Craig, et al. [4]</td>
<td>67</td>
<td>1</td>
<td>2 years</td>
<td>1.6</td>
<td>Nodule</td>
<td>Biopsy/ Excision</td>
<td>N/A</td>
</tr>
<tr>
<td>Godoy, et al. [1]</td>
<td>57</td>
<td>1</td>
<td>35 years</td>
<td>1.5</td>
<td>Nodule</td>
<td>Biopsy/ Excision</td>
<td>N/A</td>
</tr>
<tr>
<td>Sidwell, et al. [2]</td>
<td>6</td>
<td>1</td>
<td>Few months</td>
<td>0.5</td>
<td>Nodule</td>
<td>Excision</td>
<td>N/A</td>
</tr>
<tr>
<td>Chen, et al. [10]</td>
<td>89</td>
<td>1</td>
<td>3 weeks</td>
<td>1.5</td>
<td>Nodule</td>
<td>Excision</td>
<td>N/A</td>
</tr>
<tr>
<td>Richmond, et al. [13]</td>
<td>15</td>
<td>1</td>
<td>3-4 years</td>
<td>1</td>
<td>Nodule</td>
<td>Excision</td>
<td>Nil at 1 month</td>
</tr>
<tr>
<td>Djenic [11]</td>
<td>17</td>
<td>1</td>
<td>Over 10 years</td>
<td>2</td>
<td>Nodule</td>
<td>Excision</td>
<td>N/A</td>
</tr>
<tr>
<td>Current case</td>
<td>31</td>
<td>1</td>
<td>5 months</td>
<td>0.6</td>
<td>Nodule</td>
<td>Excision</td>
<td>Nil at 12 months</td>
</tr>
</tbody>
</table>

N/A – Information not available

It is essential that GCTs be confirmed with S-100 staining which is noted to be positive in all cases [17]. Positive staining with antiserum to S - 100 indicates that granular cell myoblastomas are of neural origin arising from Schwann cells [18]. Vimentin (fibroblast intermediate filament) which is the major intermediate filament found in non-muscle cells like fibroblasts and Schwann cells is also known to be expressed by GCTs [19,20]. Immunohistochemistry confirmed both in our current case. GCTs have been noted to be more common in blacks generally [10,21] with two-thirds of cases being reported in blacks [22]. However, aside the explicit mention of the patient with scrotal GCT being black in the reports by Craig [4] and Altman [16], this is the third mention of scrotal GCT in a black man and the first mention in a native African from Nigeria, which has the largest population of indigenous black men worldwide. Unanswered questions therefore include; do scrotal GCTs differ from GCTs found in other regions of the body, being less common in blacks or is it a case of the tumors being missed? Benign scrotal GCTs are noted to be small sized, very slow growing tumors which are frequently solitary and hardly presenting with other symptoms. There is thus a high likelihood that patients do not consider them a problem worth reporting. There therefore has to be a high index of suspicion for these tumors to be diagnosed, and especially so because of the small percentage of GCTs that are malignant. However, till date, no malignant case of scrotal GCT has been recorded. Generally, between 1-2% of cases of GCTs are reported to be malignant [23]. Recurrence rates of benign GCTs range between 2 and 8 %, even when the resection margins show no evidence of tumor; and rise to 20% and above when the surgical margins are positive for tumor [24]. Reported cases of scrotal GCTs have been followed up for variable periods ranging from 1-24 months with no evidence of recurrence. Our own patient has not shown any evidence of recurrence 12 months post operatively.

### Conclusion

Scrotal granular cell tumors are very rare. Benign disease is rarely associated with symptoms making it possible for the disease to be missed easily. Diagnosis therefore requires a high index of suspicion. The potential for it to be malignant makes it imperative for there to be histologic diagnosis. Complete excision with tumor free margins is the standard of treatment to reduce chances of recurrence.

### References


