Rare skin cancers in general practice

Case study
Mr LA has long been troubled with actinic damage to his skin, especially his face. He has had many squamous cell carcinomas (SCCs) removed and many solar keratoses managed. On this occasion Mr LA had two actinic lesions on his left cheek that failed to respond to cryotherapy (Figure 1). A biopsy of each site produced a surprise. Histology of the superior lesion revealed sebaceous carcinoma (Figure 2). This is an uncommon yet aggressive cutaneous malignancy derived from sebaceous glands. The 5 year survival rate is 60–70%. The tumour was widely excised with a minimum 10 mm margin. A multidisciplinary approach resulted in a decision not to proceed to adjunctive radiotherapy. The wound was well healed by 8 weeks (Figure 3). Four years on there is no sign of local or regional recurrence (Figure 4).

Many sebaceous carcinomas occur on the eyelids where the outcome is often poor; and some patients are prone to multiple other cutaneous SCCs.

There is also a rare syndrome called Muir-torre of visceral neoplasms associated with sebaceous carcinoma on the skin. As this is an autosomal dominant condition, family history and counselling is an essential part of management (enquire about family history of internal malignancies). A family member’s diagnosis can be important for other family members and offers screening for internal and cutaneous malignancies.

Mr LA’s tumour reminds us that among the basal cell carcinomas (BCCs), SCCs and melanomas removed in large numbers in Australia every day, there are unusual malignancies that we may come across from time-to-time. While we have large studies comparing management options in the more common cutaneous malignancies, it is rarely possible to have large management trials of tumours that none of us see frequently. Treatment is less clear and needs several good minds working together.

The relative severity of some of the rare tumours is tiered and summarised in Table 1. Sarcomas generally have among the poorest outcomes.

Summary of important points
• Rare tumours don’t get diagnosed clinically. They are invariably diagnosed as a ‘surprise’ on the histology report.
• Consider biopsy of the atypical or recalcitrant actinic keratoses before proceeding to another ‘freeze’.
• Send every specimen to the histologist, not the bin.

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or lipoma can sometimes lead to such a ‘surprise’.4

• Consult with colleagues and consider involving multidisciplinary experts in management of that surprise unusual tumour. Management predominantly involves surgery but can involve adjunctive radiotherapy or chemotherapy (Table 1).  
• Unusual tumours are often diagnosed late and many have poor prognoses.

- Sometimes a small biopsy may not provide the answer and complete local excision is required for histologic diagnosis.3
- Check whether there are management trials for possible enrolment of your patient with an unusual tumour.

Conflict of interest: none.

References

Table 1. Some rare skin malignancies and management considerations (tumours are ordered from the most to least fatal)

<table>
<thead>
<tr>
<th>Tumour</th>
<th>5 year survival</th>
<th>Clinical characteristics</th>
<th>Surgery involved</th>
<th>Chemotherapy involved</th>
<th>Radiation involved</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cutaneous angiosarcoma (AS)</td>
<td>15%</td>
<td>Mostly on face, scalp, or breast. Local recurrence and metastatic spread frequent, especially to lung. Can occur in postradiation scar</td>
<td>Very wide local excision</td>
<td>Often</td>
<td>Often</td>
</tr>
<tr>
<td>Merkel cell carcinoma (MCC)</td>
<td>40–68%</td>
<td>Local recurrence common even after very wide surgery. Mostly on head and neck. Spontaneous regression can occur</td>
<td>Mohs or very wide surgery</td>
<td>Occasionally</td>
<td>Often</td>
</tr>
<tr>
<td>Sebaceous carcinoma (SC)</td>
<td>60–70%</td>
<td>Often subcutaneous, often on eyelids or scalp. Can be associated with visceral tumours</td>
<td>Very wide local excision</td>
<td>No</td>
<td>Often</td>
</tr>
<tr>
<td>Dermatofibrosarcoma protuberans (DFSP)</td>
<td>93+%</td>
<td>Local recurrence/destruction in 50–75% of cases; can be fatal. &lt;5% metastasis. Can look like a morphoeic BCC. Often extends well beyond apparent borders. Typically in young/middle aged with predilection for pectoral and pelvic regions</td>
<td>Mohs surgery ideal</td>
<td>Limited</td>
<td>Some</td>
</tr>
<tr>
<td>Digital papillary adenocarcinoma</td>
<td>95%</td>
<td>Occurs on the digits. Often looks encapsulated and hence less aggressive than the reality. Metastasises frequently to lungs</td>
<td>Amputate digit</td>
<td>Some</td>
<td>Often</td>
</tr>
<tr>
<td>Microcystic adnexal carcinoma (MAC)</td>
<td>99%</td>
<td>Behaves like aggressive BCC. High local recurrence risk. 90% are on head and neck</td>
<td>Mohs surgery ideal</td>
<td>Some</td>
<td>Some</td>
</tr>
<tr>
<td>Eccrine adenocarcinoma</td>
<td>99+%</td>
<td>Behaves like aggressive BCC. Many around eye. Many subtypes. High local recurrence risk</td>
<td>4 mm margin excision</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Atypical fibroxanthoma (AFX)</td>
<td>99+%</td>
<td>Low grade sarcoma. Behaves like a BCC. Metastasis very rare. Most occur on elderly, sun damaged head and neck skin. Can occur in old radiation scars</td>
<td>4 mm margin excision</td>
<td>No</td>
<td>No</td>
</tr>
</tbody>
</table>
| Kaposi sarcoma (KS)              | Death from other cause | Three subtypes:  
  • elderly of Jewish/Mediterranean descent  
  • immunosuppressed (eg. postrenal transplant)  
  • HIV/AIDS related | No                                         | Often                 | Usual              |


