The summer skin check

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BACKGROUND
Skin cancer is a major public health problem in Australia and represents a substantial health cost. General practitioners provide the majority of care to patients with skin cancer, so becoming familiar with the clinical features and management of these tumours is important.

OBJECTIVE
To provide a pictorial essay on the common types of benign and malignant skin lesions encountered in general practice, and briefly describe key clinical features, differential diagnosis and management options.

DISCUSSION
Examination for skin cancer should be considered in the general practice setting for all patients over the age of 40 years, particularly the elderly. A proper skin check requires a systematic approach and ideally an entire consultation should be set aside for this purpose. However, clinical examination conducted for other purposes provides an opportunity for screening and early detection of skin cancer.

History taking
The following points in history taking can help identify high risk patients:
• blistering sunburns
• occupational and recreational exposure to solar radiation
• family or personal history of melanoma or multiple atypical (dysplastic) naevi
• family or personal history of non-melanoma skin cancer (NMSC)
• immunosuppression – higher rates and earlier onset of skin cancer is present in transplant patients, therefore a high index of suspicion is necessary in this population
• exposure to arsenic is a known risk factor for the development of basal cell carcinoma (BCC) and Bowen disease
• a history of change or symptomatology in any lesion – skin cancers are changing lesions and the time course for this change is generally evident over a period of months.

Examination
The following factors are important in examination:
• good lighting, preferably natural light
• magnification – a dermatoscope can be helpful in diagnosis of both melanocytic and nonmelanocytic lesions
• positioning – patients should be undressed to their underwear, so a warm room or blanket for comfort is desirable. Having the patient lie on the examination couch can be useful so the soles of the feet are not missed
• look closely – gently stretching the skin can often make BCCs more obvious.
• palpate – dermatofibromas for example have a characteristic feel on palpation
• concentrate on high risk areas – the highest rates for NMSC are found on the face, especially the nasolabial fold, eyelid and the nonmucosal skin of the lip followed by the ears, nose and cheek. In men, the neck, back and shoulders, and in women, the neck, shoulders and outer arms are also sites of predilection
• scars should be checked for evidence of recurrence of previously excised lesions - BCCs may develop within longstanding scars and ulcers
• lymph nodes should be assessed for the early detection of metastatic disease if there is a history of melanoma or squamous cell carcinomas (SCCs).

Management
Biopsy of a lesion (punch or shave) is advisable when a firm clinical diagnosis cannot be made or when the treatment choice may be dictated by the tumour type or pattern of growth. Multiple 2 mm punch biopsies may be necessary to define poorly demarcated tumours. Tailor each management decision to the particular lesion in the individual patient. In general, simple surgical excision with primary closure is the treatment of choice for most skin cancers, although some anatomical areas require particular care (Table 1) and specialist referral may be desirable for technically difficult excisions. Prevention should be emphasised (Table 2) and changes to look for in suspicious lesions discussed. Follow up is recommended for three reasons: detection of further primary tumours, detecting local persistence of previously treated tumours, and early detection of metastatic disease.
Seborrhoeic keratoses

Clinical features
Seborrhoeic keratoses (Figure 1, 2) are very common benign tumours usually found on the face and trunk. They develop gradually as slightly elevated, ovoid yellow-brown, well defined papules with a greasy appearance, later thickening to become brownish-black. The surface is irregular and shows minute pits.

Management
When flat and pigmented, seborrhoeic keratoses may require biopsy to differentiate them from lentigo maligna melanoma. These can be removed with a number of destructive techniques such as cryotherapy, curettage, electrosurgery or shave excision.

Multiple eruptive seborrhoeic keratoses are rare but may be a sign of internal malignancy (Leser-Trelat sign)

Dermatofibroma

Clinical features
Dermatofibroma (Figure 3) are benign tumours of fibroblasts that are firm, solitary dermal nodules usually found on the limbs of younger people. They have an ‘iceberg’ effect in that they feel larger than they look. The overlying epidermis is often lightly pigmented and dimples when the nodule is squeezed.

Management
No treatment is usually necessary, although biopsy may be performed if the diagnosis is uncertain.

Porokeratosis

Clinical features
Porokeratoses (Figure 4) are uncommon, distinctive, benign lesions due to clones of epidermal cells that show varying degrees of dysplasia. They develop as annular dry plaques surrounded by a raised fine keratotic wall. Porokeratoses may be isolated or multiple and most often appear on the limbs.

Management
None usually needed. If lesions are hyperkeratotic or multiple, cryotherapy may be performed.

The rare giant forms may be premalignant

Solar keratoses

Clinical features
Solar keratoses (Figure 5) are common lesions associated with dysplasia, particularly within basal keratinocytes. They represent a potential precursor of SCC, although only a small percentage evolve into invasive SCCs.
Clinical features
Solar keratoses are found on the chronically sun exposed sites of the head and neck, dorsa of the hands, and the forearms, and present as erythematous macule with superimposed hyperkeratosis. Solar keratoses are generally multiple and may be confluent.

Differential diagnosis
Bowen disease, SCC, solar lentigo or lentigo maligna if pigmented.

Management options
- No treatment
- Cryotherapy
- 5-flurouracil
- Imiquimod
- Curettage
- Laser resurfacing.

Thickening and tenderness on lateral palpation are indicators of possible transformation into invasive SCC

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Differential diagnosis
- SCC
- Atypical fibroxanthoma
- Merkel cell carcinoma
- Amelanotic melanoma.

Management options
- Partial biopsy will almost always be reported as a well differentiated SCC as pathology requires the entire architecture to suggest the possibility of keratoacanthoma. If in doubt, treat as a SCC
- Curettage +/- cryotherapy
- Surgical excision
- Superficial radiotherapy
- Laser ablation.

Due to extension of keratinocyte atypia, down hair follicles recurrence after treatment with superficial modalities (eg. cryotherapy) can occur, particularly on the face

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Bowen disease
Bowen disease (intraepidermal SCC) (Figure 6) is associated with full thickness epidermal dysplasia with follicular involvement and is classified as an intraepidermal SCC. Despite this, they tend to have a prolonged in-situ phase that may last many years.

Clinical features
Longstanding, slowly enlarging lesions that are generally asymptomatic with a predilection for the lower legs (especially in females) but may occur at any site. Lesions are a sharply defined, round to oval, erythematous hyperkeratotic plaque. The degree of hyperkeratosis may vary. The rate of transformation to invasive SCC has not been established, but appears to be low.

Keratoacanthoma
Keratoacanthoma (Figure 7) are lesions that can be thought of as a well differentiated forms of SCC characterised by spontaneous resolution.

Clinical features
Keratoacanthoma are most common on the chronically exposed sites of the head, neck, hands, and forearms. They rapidly enlarge over a period of 4–8 weeks to form a tender, erythematous nodule with a central keratotic plug. Resolution takes 6–12 weeks, sometimes leaving a scar. A minority will persist as SCC.

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Squamous cell carcinoma
Squamous cell carcinomas (Figure 8, 9) are characterised by lobular proliferation of keratinocytes with intradermal invasion and prominent keratin production. The majority of SCCs are thought to arise from solar keratoses.

Figure 6. Bowen disease
Figure 7. Keratoacanthoma
Figure 8. Squamous cell carcinoma
Figure 9. Squamous cell carcinoma
Clinical features
Squamous cell carcinomas are characterised by tender erythematous papules or nodules, with a variable amount of hyperkeratosis enlarging over a period of months, usually with associated tenderness. Bleeding and ulceration may develop and the risk of metastasis increases with tumour thickness, clinical immunosuppression and poorly differentiated tumours.

Differential diagnosis
- Hyperkeratotic solar keratosis or Bowen disease – clinical diagnosis can be difficult in the early stages
- Nodular basal cell carcinoma
- Amelanotic melanoma
- Atypical fibroxanthoma – particularly if on the scalp.

Management options
- Surgical excision
- Superficial radiotherapy.

Beware tumours located on lips, ears or scalp as they have a higher risk of early metastasis usually to local lymph nodes.

Basal cell carcinomas
Basal cell carcinomas (BCCs) are tumours derived from basal keratinocytes. Their onset is insidious. In months or years they extend peripherally as well as in depth and are capable of destroying cartilage and bone. Metastases rarely develop from BCCs, the main risk being recurrence after incomplete removal. There are three common growth patterns of BCC (superficial multifocal, nodular and morphoeic) that have a distinctive clinical presentation. Any of these may also show ulceration or pigmentation.

Superficial multifocal BCC
Clinical features
Superficial BCCs generally occur on the trunk and limbs, are bright pink and shiny, and usually well defined erythematous macular lesion (Figure 10). Stretching the skin will highlight a thread-like pearly rim or islands of pearliness throughout the lesion. There is progressive enlargement over months to years but ulceration and bleeding is uncommon. With time, areas of nodular or even sclerosing growth pattern may supervene. Recurrence is common.

Differential diagnosis
- Bowen disease
- Solar keratosis
- Psoriasis
- Eczema.

Management options
- Cryotherapy
- Imiquimod
- 5-flurouracil
- Photodynamic therapy
- Surgical excision
- Curettage and diathermy
- Superficial radiotherapy.

Nodular BCC
Clinical features
Nodular BCCs (Figure 11) are more often found on the head and neck. They are a shiny, translucent (pearly), telangiectatic papule or nodule. Recurrent ulceration is frequent and pigmentation may suggest melanoma.

Differential diagnosis
- Nonpigmented intradermal naevi
- Neurilemmas
- SCC

Management options
- Surgical excision
- Superficial radiotherapy
- Intralesional interferon alfa
- Imiquimod
- Photodynamic therapy.

Morphoeic BCC
Histology
Morphoeic lesions have a sclerosing growth pattern with fibrosis surrounding areas of BCC (Figure 12).

Clinical features
Morphoeic lesions present mostly on the head and neck and can be clinically difficult to detect. They are often indurated to palpation, are frequently asymptomatic and tend to look like a pale scar. Morphoeic lesions are usually longstanding and tend to be deeply invasive.

Differential diagnosis
Major differential diagnosis is a scar and biopsy is necessary to establish the diagnosis.

Management options
- Consider referral for specialist care
- Moh’s micrographic surgery
- Superficial radiotherapy.

Morphoeic lesions need to be treated with respect as they can have extensive subclinical extension and perineural invasion particularly when located around the eyes, ears and nose. Recurrence is frequent due to incomplete excision.

Figure 10. Superficial BCC
Figure 11. Nodular BCC
Figure 12. Morphoeic BCC
Solar lentigo

Clinical features

Solar lentigo (Figure 13) represent a macular area of pigmentation appearing after either acute or chronic sun exposure. Histologically there is a linear increase of melanocytes at the dermoepidermal junction without cytologic atypia. Backs of the hands, face and shoulders are common sites. Over time they may evolve into seborrhoeic keratoses.

Management options

- Cryotherapy can lighten solar lentigo, but care should be taken in olive skinned patients as they may hyperpigment
- Laser.

Shave biopsy may be necessary to distinguish solar lentigo from lentigo maligna or superficial spreading melanoma

Atypical naevi

Clinical features

Atypical (dysplastic) naevi (Figure 15) are usually larger than ordinary naevi, predominate on the trunk and often showing a mixture of tan, dark brown and pink areas. The surface texture is often ‘pebbly’ and the border ‘smudgy’. The term atypical naevus syndrome refers to the occurrence of multiple (up to 80 or more) dysplastic naevi in an individual. Melanoma risk is increased in these patients with the risk increased further in melanoma prone families with atypical mole syndrome.

Management

Surveillance (baseline photography and regular 6–12 month follow up) enables early diagnosis of melanoma. This is much more cost effective in preventing life threatening melanoma than prophylactic excision of atypical naevi.

Due to the increased risk of melanoma in these patients regular follow up is necessary for life

Melanoma

Patients are often most concerned about pigmented lesions because of the possibility of melanoma and its capacity to be lethal. Table

Table 3. Margins of re-excision for melanoma after initial excisional biopsy

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Ink spot lentigo

Ink spot lentigo (Figure 14) are small (less than 5 mm) densely black macules, usually on sun exposed skin. They can be cause for concern because of their dark colour and irregular lateral margin. Melanocyte numbers and melanin production are increased in these lesions but there is no cellular atypia.

Although often conspicuous, ink spot lentigo are benign lesions

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Clinical practice: The summer skin check

Figure 17. Superficial spreading melanoma

Differential diagnosis

• Atypical (dysplastic) naevus
• Pigmented BCC
• Pigmented seborrhoeic keratosis
• Haemangioma.

Management

• Surgical excision.

Nodular melanoma

Nodular melanoma represent 15–35% of all melanomas and appear as a pigmented (or occasionally flesh coloured) amelanotic nodule with no antecedent in-situ phase (Figure 18). It is the most rapidly growing and aggressive type of melanoma.

Figure 18. Nodular melanoma

Differential diagnosis

• Pigmented BCC
• SCC or pyogenic granuloma if amelanotic.

Management

• Surgical excision.

Acral lentiginous melanoma

Acral lentiginous melanoma represent 5–10% of all melanomas and appear on the palmar, plantar and subungual skin (Figure 19). Although rare in caucasians, they are particularly common in both the Chinese and Japanese population.

Figure 19. Acral lentiginous melanoma

Differential diagnosis

• Melanocytic naevus
• Subungual haematoma
• Talon noir – a pigmented petechial area on the heel following minor trauma.

Management

• Surgical excision.

Partial biopsy of the suspected melanoma should be avoided as this will interfere with accurate histological staging

Acknowledgment

Thanks to Professor Robin Marks from St Vincent’s Hospital Melbourne and Associate Professor John Kelly, Victorian Melanoma Service, Alfred Hospital, Melbourne, for kindly allowing reproduction of some of the clinical images in this article.

Suggested reading


Conflict of interest: none declared.

References


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