A pearly papule with telangiectasia

Alvin Lim, James Muir

CASE
A man aged 52 years with a history of non-melanoma skin cancer presented for a skin examination. He had a history of chronic sun exposure since childhood. He had no history of melanoma. He had no other comorbidities and did not take any regular medications. On examination, he had extensive sun-damaged skin on his face. On the left lateral aspect of the chin there was a firm, non-tender, skin-coloured papule (Figure 1). He was unaware of this lesion.

QUESTION 1
What is the differential diagnosis for a skin-coloured facial papule?

QUESTION 2
What features would you look for on dermoscopy to differentiate between causes of skin-coloured facial papules?

ANSWER 1
The clinical differential diagnosis for this lesion includes:
- basal cell carcinoma (BCC)
- amelanotic melanoma
- intradermal naevus
- angiofibroma (fibrous papule of the face)
- adnexal tumours
- sebaceous hyperplasia.

ANSWER 2
Dermoscopy reveals diagnostic features invisible to the naked eye or loupe magnification. BCC, intradermal naevi, sebaceous hyperplasia and amelanotic malignant melanoma typically show distinctive dermoscopic features that allow differentiation. Table 1 outlines the distinguishing features for the differential diagnosis for a non-pigmented facial papule.

CASE CONTINUED
The lesion was examined with non-polarised contact dermoscopy (Figure 2). On dermoscopy, there was a pearly papule measuring 2 mm with branching blood vessels (arborising telangiectasia). The papule was abutting a hair follicle. Of note, there was background sun-damaged skin characterised by erythema, pink-red pseudonetwork (annular pigmentation that resembles a network, surrounding white areas corresponding to follicular openings), unfocused blood vessels and white halo structures around yellow (keratin-filled) follicular openings.

QUESTION 3
What is the presumptive diagnosis at this stage?

QUESTION 4
What is the next step in management?

ANSWER 3
BCC is the presumptive diagnosis in a high-risk patient with multiple non-melanoma skin cancers and chronic sun exposure. It is commonly diagnosed by clinical and dermoscopic examination. Clinically, BCC appears shiny and pearly (translucent), which can be made more obvious by stretching the surrounding skin.

On dermoscopy, a pinkish hue and arborising telangiectasia are common
Angiofibroma, also called fibrous papule, is a benign tumour caused by proliferation of fibroblasts, collagen and dilated blood vessels in the dermis.12 Usually <5 mm in size, it appears commonly on the face as a shiny, smooth skin-coloured or pink papule. Dermoscopically, it may have cobblestone, globular or structureless pigmentation. Comma vessels or arborising vessels may be observed, but they lack other BCC-specific features.25 It is usually present from adolescence without history of change.

Sebaceous hyperplasia appears clinically as a pearly papule on the forehead and cheeks of elderly individuals. Dermoscopically, it is distinguished from BCC by multiple white-yellow globules and branching vessels that do not cross the centre (a follicular opening).15

**Table 1. Differential diagnoses for a non-pigmented facial papule**

| Diagnosis                  | Characteristics                                                                 |
|----------------------------|---------------------------------------------------------------------------------|
| Basal cell carcinoma (BCC) | BCC is the most common skin malignancy. It is associated with chronic sun exposure and most commonly occurs on the head and neck. Predisposing factors include fair skin, male sex, age >50 years, prior history of skin cancer, exposure to ionising radiation or arsenic, scars and immunosuppression.4 Dermoscopic features include arborising telangiectasia, focal ulceration, white structureless areas and white perpendicular lines.2 |
| Amelanotic melanoma (AM)   | AM may mimic BCC clinically. Lacking pigment, AM may appear as a nodule that is skin-coloured, pink, or red. Dermoscopically, the vascular pattern may help distinguish AM from benign, nonpigmented lesions, particularly the presence of linear irregular vessels and milky red globules.10,11 |
| Intradermal naevus          | Intradermal naevus is a benign melanocytic proliferation that appears as a nodule that is skin-coloured, pink or light brown. Dermoscopically, it may have cobblestone, globular or structureless pigmentation. Comma vessels or arborising vessels may be observed, but they lack other BCC-specific features.25 It is usually present from adolescence without history of change. |
| Sebaceous hyperplasia       | Sebaceous hyperplasia appears clinically as a pearly papule on the forehead and cheeks of elderly individuals. Dermoscopically, it is distinguished from BCC by multiple white-yellow globules and branching vessels that do not cross the centre (a follicular opening).15 |

**ANSWER 4**

The diagnosis of BCC should be confirmed histologically. Consideration should be given to the size, shape and site of the lesion when deciding on biopsy technique.3 Where possible, for clinically suspected non-melanoma skin cancer, complete excision of the lesion is recommended as per the Australian Cancer Council Guidelines in order to facilitate assessment of its nature and extent.7 For small lesions, an excisional biopsy with an appropriate margin serves as both a diagnostic test and definitive treatment. If this is not possible, a partial biopsy from the centre of the lesion is preferred, with a view to definitive management following histological examination.5

**CASE CONTINUED**

The lesion was excised with an ellipse with 2 mm margins with the intent of achieving definitive treatment. The specimen was sent for histology. On histopathology, no malignancy was found on initial examination. Further examination of multiple superficial and deep sections did not reveal evidence of malignancy. The lesion was diagnosed as a benign fibrous papule with surrounding solar elastosis and telangiectasia.

**QUESTION 5**

What are the consequences of an incorrect diagnosis? What are the risks for excisional biopsy as a diagnostic and treatment modality?

**QUESTION 6**

How accurate is clinical and dermoscopic detection of skin cancer?

**ANSWER 5**

An incorrect diagnosis of skin cancer using visual inspection and dermoscopy may lead to unnecessary excision, inappropriate treatment (eg non-surgical treatments) and patient anxiety.8 The risk of excisional biopsy includes subjecting the patient to an invasive procedure and its associated consequences – scar and wound management, risk of infection, cosmetic disfigurement, functional impairment and expense. The size of the resulting skin defect should be considered when deciding whether this method is appropriate. If surgery will produce a significant scar, then consideration should be given to an initial biopsy.
ANSWER 6
Dermoscopy increases the accuracy of skin cancer detection when added to visual inspection. A meta-analysis reported the sensitivity and specificity of dermoscopy for detecting BCC to be 91.2% and 95%, respectively. Adding dermoscopy to visual inspection with the naked eye increased both sensitivity (from 66.9% to 85%) and specificity (from 97.2% to 98.2%). Although various dermoscopic algorithms have been published to aid melanoma diagnosis, there is no clear evidence that one is superior. A 2016 Cochrane review indicated no improvement in using a dermoscopic checklist for melanoma diagnosis versus clinician experience. A comparative study of diagnostic error in order to consider limitations, and it is important for clinicians to be aware of the possibility of diagnostic error in order to consider the most appropriate management.

Key points
• Facial papules have a broad range of potential benign and malignant diagnoses.
• A pearly papule with telangiectasia on the face of a patient with risk factors for skin cancer should raise suspicion for malignancy.
• Clinical diagnosis of skin cancer has limitations, and it is important for clinicians to be aware of the possibility of diagnostic error in order to consider the most appropriate management.

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Correspondence ajgp@racgp.org.au