The Basal Cell Naevus Syndrome: A Case in the Falkland Islands

Maj R Pilcher*
BSc, FDSRCPS, RADC
Senior Specialist in Oral Surgery
Princess Alexandra Hospital, Wroughton, Swindon, Wilts. SN4 0QJ

SUMMARY: A case of basal cell naevus syndrome is presented. The jaw cysts were proved histopathologically to be odontogenic keratocysts. The multiple naevoid lesions on the skin were identified as basal cell carcinoma. Intracranial calcification is illustrated to a degree not previously reported in a case with this syndrome. The importance of identifying this syndrome is discussed and its presentation in the Falkland Islands is reported.

Introduction
The basal cell naevus (Gorlin-Goltz) syndrome was first described by Jarisch in 1894 (1): the paper by Gorlin and Goltz was published in 1960 (2) with a review by Gorlin in 1987 (3). Two Egyptian skeletons of the Early Dynastic period have been shown to have bony anomalies compatible with this diagnosis (4).

The syndrome shows autosomal dominant inheritance with no sex predilection, variable expressivity and high penetrance in up to 50% of cases (5).

Gorlin and Goltz described a case with multiple jaw cysts, multiple basal cell naevi on the back and face, and bifid ribs. Since then other reports have added other anomalies associated with the syndrome which are summarised as follows:

(a) Cutaneous anomalies: including basal cell naevi, benign dermal cysts and tumours, palmar pitting, palmar and plantar keratosis and dermal calcinosis. The basal cell naevi may appear at birth or be first noted in infancy. Occasionally they may begin at about puberty and gradually increase in size and number over a period of years. There may be few or several hundred and there is a tendency for bilateral distribution, but this need not be symmetrical. The size of the individual lesion varies from that of a pinhead to approximately 5 mm in diameter. Pedunculated and papular lesions are common on the neck, eyelids and axillae. The colour is that of the surrounding skin although they may be hyperpigmented. The individual lesions are elevated, firm, symptomless, and painless to touch. Their surface is smooth and occasionally may be covered by minute blood vessels. Their distribution varies the commonest areas being the trunk and extremities. Although aggressive behaviour may occur resulting in gross destruction the majority behave in a benign fashion. A lesion which has ulcerated is a sign of aggressive change. They may closely resemble the appearance of neurofibromatosis. The following features may precede the appearance of the skin lesions.

(b) Dental and osseous anomalies including multiple jaw cysts, mild mandibular prognathism, cleft lip and palate, rib anomalies (often bifid), vertebral anomalies and brachymetacarpalism. The jaw cysts are the most common characteristic of the syndrome. They may be simple occurring close to the apex of a tooth, dentigerous containing a tooth or primordial (odontogenic keratocyst). They may develop by the 7th year of age and are usually located in the premolar-molar area resulting in displacement of the child’s teeth. Mandibular cysts are more common than maxillary cysts and often secondary infection prompts oral surgical attention. Ballooning and weakening of the bone may cause deformity and pathological fractures.

(c) Ophthalmologic abnormalities including hypertelorism with wide nasal bridge, congenital blindness and internal strabismus.

(d) Neurological anomalies including mental retardation, dural calcification, agenesis of the corpus callosum, congenital hydrocephalus and an increased incidence of medulloblastomas.

(e) Sexual abnormalities including hypogonadism in males and ovarian tumours in females (6).

A relationship to pseudohypoparathyroidism has been suggested due to the lack of response to parathormone and the shortened fourth metacarpals found in some patients (7).

Histologically the basal cell naevi are identical to basal cell carcinoma. Clinically they differ by their multiplicity, their early age at appearance and capability for rapid growth and central erosion.

The odontogenic keratocysts of the jaw show a high incidence of recurrence following removal and are at increased risk of association with malignancy such as fibrosarcoma (8-10), ameloblastoma (11), squamous cell carcinoma (12-13), leiomyoma (14-15) and spindle cell carcinoma (16). These cases illustrate the potentially highly destructive nature of the keratocyst being capable of destroying a major part of the facial skeleton with involvement of the orbit and contents.

Case Report
A 56 year old Caucasian male farmworker of the Falkland Islands presented complaining of a painful
Clinical findings were a characteristic facies with bossing of frontal and parietal bones, well developed supraorbital ridges, heavy eyebrows, hypertelorism, a broad nasal root, hypoplasia of the maxilla and mild prognathism (Fig 1). Naevi were present on both left and right eyelids (Fig 2) and he had an epidermoid cyst on his left forearm.

Radiographic findings included multilocular cystic lesions not only in the right body of his mandible (Fig 3) but also in the other three jaw quadrants (Fig 4). Intracranial calcification consisted of lamelliform calcification of the falk cerebri and the tentorium cerebelli, calcification of the wall of the superior sagittal sinus and calcification of the arachnoid granulations (Fig 5). Calcification of the sella turcica was also present.

Routine haematology and biochemical investigations were within normal limits including repeated serum calcium and phosphate levels.

Under endotracheal general anaesthesia the naevi were excised and the cysts enucleated. The left maxillary sinus was found to be largely destroyed. There was no involvement of the orbit.

Histopathological examination confirmed that the skin lesions were basal cell carcinoma (Fig 6). The jaw cysts in all four quadrants proved to be odontogenic keratocysts. They had a lumen containing epithelial debris and a lining of stratified squamous epithelium of about 6-8 layers thick which had a parakeratinised surface; there was a prominent basal cell layer and a subepithelial layer containing a chronic inflammatory infiltrate (Fig 7).

No other cases were identified in his blood relatives. The patient currently had no plans for marriage or to have offspring. Advice was given regarding avoiding sun exposure, wearing protective clothing and the use of sunscreens. Long term review of the patient and relevant relatives was arranged.
Discussion
Earlier diagnosis of this patient as having basal cell naevus syndrome had not been made despite having had multiple basal cell carcinoma excised on previous occasions. It is important in any patient presenting with multiple basal cell carcinoma to rule out the presence of this syndrome; this is because of the many associated problems which these patients may eventually face and the necessity to screen for other abnormalities which may be already present in the patient or relatives. Long term review must be arranged and genetic counselling organised.

Management of the basal cell carcinomata should be by excision. Mohs micrographic surgery (17) is indicated in the treatment of basal cell carcinoma arising in high risk areas particularly around the eyes and lips: the area macroscopically involved is removed after initial debulking or curettage, and divided for processing. The resected tissue and the surrounding skin are marked with ink and the edges of the tumour pieces are coded with dyes to aid orientation. A precise map showing the coded tissue segments in relation to the local anatomy is constructed. Tissue is subsequently processed taking horizontal rather than vertical sections allowing examination of 100% of the tumour base. Any remaining tumour can be clearly identified on the slide, the map and the patient enabling a second stage of excision to be directed specifically to the residual tumour leaving normal skin unscathed. When the basal cell carcinoma are too numerous to excise they must be watched closely and removed selectively when they threaten the patient by becoming large and ulcerated (18).

Radiotherapy for multiple lesions should be avoided as irradiation of the skin in this population leads to a striking increase in the number of basal cell carcinomata within the treated field (19-20). A further contraindication is that radiotherapy may promote malignant changes in jaw lesions which are in the radiation field (8 and 13).

Systemic retinoids have been shown to cause regression of the basal cell carcinomata and to decrease the rate of appearance of new lesions; however baseline activity quickly returns after discontinuation of therapy (21). Intralesional sustained release interferon has been used with some success (22).

Management of the odontogenic keratocysts is...
complicated by their tendency to recur. Suggested mechanisms by which they recur include the presence of satellite cells, increased mitotic activity of the cyst lining epithelial cells and increased collagenolytic and fibrinolytic activity of the cyst walls (23-27). Friability of the lining and difficulty in completely removing the cyst may also be a causal factor.

Browne (28) found there was no significant difference in the recurrence rate following treatment by three basic methods: marsupialisation, enucleation and primary closure, or enucleation and packing the wound open. He concluded that the rate or recurrence was related to the nature of the lesion and not to the method of treatment. Voorsmit (29) advocates the use of Carnoy’s solution in the cyst prior to enucleation. Ahlfors et al (30) advocate that the lesion be regarded as a benign cystic tumour rather than a developmental or other type of cyst and that it is more aggressive when associated with this syndrome. They suggest marginal resection including a rim of uninvolved bone, similar to the treatment in cases of unicystic ameloblastoma. Adherence of the cyst lining overlying mucosa may also require excision of this part of the mucosa. Mustaciuco et al (31) described enucleation of extensive keratocysts initially; rather than resection, thus maintaining the integrity of the mandible. The subsequent bony healing provided greater substance to the mandible allowing treatment of recurrence by block resection without involving the inferior border. McLoughlin et al (32) advocate aggressive surgery at an early stage of cyst development.

Long term review of the patient together with screening and long term review of relevant relatives plus genetic counselling will reduce both the morbidity and mortality attributable to the syndrome.

The presence of this syndrome in the Falkland Islands has not previously been reported.

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Fig 6. Basal cell carcinoma from lower left eyelid. H&E x 190.

Fig 7. Odontogenic keratocyst lining from left maxilla. H&E x 190.

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