Primary melanocarcinoma of the lower respiratory tract

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Primary melanocarcinoma of the trachea and bronchi is extremely rare: in only four of the 16 patients previously reported was the possibility of an alternative primary site excluded by necropsy. We present the clinical and necropsy findings in a further case with some unusual histological features and a family history of melanocarcinoma.

Case report

A 70-year-old woman presented in May 1978 with a 12-month history of increasing exertional dyspnoea, a non-productive cough for three months but no history of haemoptysis, loss of weight, or dysphagia. She had smoked 10 cigarettes daily until 1958. There was no previous medical history of note but her daughter had died aged 31 years from disseminated melanocarcinoma after removal of a cutaneous primary from the calf. The patient's general condition was good, with minimal stridor but no finger clubbing or lymphadenopathy; reduced percussion note and breath sounds with associated wheezes were noted over the right midzone anteriorly. Chest radiography showed a dense mass in the right middle lobe, enlargement of the right hilum, and an abnormal para-tracheal shadow. Numbers of darkly pigmented cells were seen on sputum cytology. Fibreoptic bronchoscopy revealed extensive black discoulouration and swelling of the mucosa at the carina spreading down both main bronchi, the lumen being almost totally occluded on the right side. A biopsy showed diffuse infiltration of the bronchial mucosa by melanocarcinoma. No alternative primary was found on clinical examination and there was no history of spontaneous regression or surgical removal of a pigmented lesion. A divided four week course of radiotherapy was given with considerable symptomatic relief, but no improvement in either the radiological or bronchoscopic appearances. She died nine weeks after presentation having developed mediastinal emphysema and bronchopneumonia.

PATHOLOGICAL FINDINGS
At necropsy, the right middle lobe was found to be widely infiltrated by solid black neoplasm as were the carina, right main bronchus, middle lobe bronchus and its medial segmental branch, and the proximal 3 cm of the left main bronchus (fig 1). The hilar lymph nodes were massively replaced by the neoplasm which had also ulcerated through the oesophageal mucosa 2.5 cm below the level of the carina. Direct extension had occurred anteriorly to involve the fourth and fifth intercostal spaces, and metastases were seen in the right axillary and internal mammary lymph nodes. The liver and the ninth thoracic vertebra each contained a solitary 1.5 cm metastasis. The nasal cavities, meninges, and eyes (both of which were sectioned) showed no neoplasm.

Microscopy showed the neoplasm to be arranged in whorls and strands and to consist mainly of spindle-shaped melanocytes containing varying amounts of melanin pigment and with oval, often vesiculated, nuclei (fig 2). In some areas dendritic cells predominated, while in others multinucleated neoplastic giant cells were prominent: mitoses were scanty. Heavily pigmented melanophores were abundant throughout the neoplasm.

Discussion
Primary melanocarcinoma of the lower respiratory tract is extremely rare. Of the sixteen cases described in the literature only four1-4 entirely excluded an alternative primary neoplasm by necropsy examination. It is likely that these neoplasms arise from neuroectodermal melanoblasts which have emigrated to the respiratory mucosa during early embryonic development. The respiratory tract develops as a downgrowth from the foregut and hence it is rather surprising that melanocarcinomas are not seen more frequently in the bronchial tree, as is indeed the case for the oesophagus and nasopharynx. Although melanoblasts have been identified in the normal oesophageal epithelium,5 no such study has been performed on respiratory mucosa.

Most reported bronchial melanocarcinomas have arisen in the large central bronchi where they present as dark brown, usually polypoidal, growths. The patients present with a short history of cough, haemoptysis, or dyspnoea as their main symptoms. Bronchoscopy and biopsy are essential to establish the diagnosis but sputum cytology may also be of benefit, as in our case. Occasional patients have survived up to 11 years after lobectomy or pneumonectomy, but most pursue a rapid downhill course until their death within months of initial presentation.

In our case the neoplasm was rather unusual histologically in that it showed several features more...
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commonly associated with a malignant blue naevus—absence of junctional activity, the presence of dendritic and spindle-shaped cells, and a relative paucity of mitotic activity. We ultimately decided, nevertheless, that the most appropriate diagnosis was that of melanocarcinoma. Of further interest was the family history of this neoplasm. The occurrence in more than one member of a family of skin melanocarcinoma is well recognised: this is the first report of familial melanocarcinoma in which a bronchial primary has been demonstrated.

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References

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