A CASE OF PULMONARY CRYPTOCOCCOSIS

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The patient, a married woman of 50, with no extensive travel outside East Anglia, had an attack of influenza in January, 1951, which left her with a cough which persisted up to the time of operation in August, 1951. There was only a trace of sputum and no haemoptysis or pain, and apart from losing 1 st. in weight over the previous six months and a slightly unhealthy appearance, she was otherwise normal and fit. She was referred to the chest physician (Dr. C. Downes), and a radiograph taken on June 7, 1951, revealed, as the only abnormality, an opacity occupying a position in the right middle and anterior portion of the right lower lobe. She was bronchoscooped on June 13, and a rounded mass was seen just inside the orifice of the middle lobe bronchus. The bronchial aspirate contained numerous morphologically typical Cryptococci neoformans, and these were also present in the bronchial biopsy. Aerobic cultures of the aspirate yielded numerous culturally typical, yeast-like colonies growing preferentially on Sabouraud’s medium and at room temperature, and more reluctantly on blood agar and at 37° C. (Kurung, 1942). A guinea-pig inoculated with the aspirate died 24 days later with gastro-enteritis and yielded positive cryptococcal cultures from spleen and peritoneum. A mouse injected with a heavy subculture survived and was killed at seven weeks; while its only gross pathology was a few foci in the liver (histologically cryptococcal), all the organs yielded positive cryptococcal cultures (liver, spleen, kidney, lung, brain, and peritoneum).

The cryptococcus strain was insensitive to penicillin, streptomycin, aureomycin, sulphathiazole, sulphasalazine, and sulpha-mezathine. There were no cerebral symptoms and the lumbar C.S.F. was normal. A right pneumonectomy was successfully performed on June 26. The incised pneumonectomy specimen revealed a tumour-like mass occupying the middle lobe, which was firm, with a diffuse extension to the basal segments of the lower lobe, which were doughy. The upper lobe was unaffected and there was no enlargement of the regional glands. Histologically the “tumour” and extension consisted of a solid mass of cryptococci replacing the parenchyma and practically devoid of any supporting stroma; it was surrounded by a fibrous capsule in which appeared lymphocytes and giant cells.
Six days after operation the patient died suddenly from embolism of the left pulmonary artery. Before making the necropsy C.S.F. was removed by cisternal puncture; this fluid was blood-stained and loaded with cryptococci. At necropsy the brain showed merely some regions of indefinite degeneration (e.g., right third ventricle) and it required extensive histology to demonstrate the presence of cryptococci in the subarachnoid layers of the medulla, the only specific abnormality found. Other organs showed nothing significant. No cultures were prepared from the necropsy material.

**DISCUSSION**

Cryptococcosis is not endemic in Great Britain, only some half-dozen cases having been previously described (Carter and Young, 1950; Daniel, Schiller, and Vollum, 1949). In the lung the disease may be tumour-like ("toruloma"), bronchopneumonic, or miliary; it is usually progressive and ultimately initiates a fatal meningitis. This predilection for the brain and meninges (60%) and lung (20%) (Levin, 1937), taken in conjunction with symptomatology, necessitates differentiation from tuberculous meningitis, tuberculoma, or pulmonary tuberculosis (Hobby, 1949). Indeed, as
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cerebral involvement is usually preceded by some pulmonary symptoms, the lung is regarded as the usual portal of entry (Krainer, Small, Hewlitt, and Deness, 1946). The disease is highly lethal, affects people of all ages, and women about twice as frequently as men. Chemotherapy is disappointing (Hamilton and Thompson, 1946), and as the broad gelatinous capsule prevents antibody formation vaccine therapy also fails (Benham, 1935). Two successful pneumonectomies have been reported (Froio and Bailey, 1949; Dormer, Friedlander, Wiles, and Simson, 1945).

The source of the fungus is equivocal; in endemic areas a strain can be recovered from the faeces of certain animals prone to natural cryptococcal infections and from fermenting fruit substrates; the fruit strain is pathogenic to these animals, but both animal and fruit strains are serologically distinct from those which occur in man, and are non-pathogenic to man. In man two strains occur, skin saprophytes and true pathogens; these are morphologically identical and serologically related, and can only be distinguished by virulence tests in animals. Because of these complex facts autogenous infection is premised, i.e., that the skin saprophyte acquires pathogenicity (Conant, Martin, Smith, Baker, and Callaway, 1944). Despite an extensive literature (over 100 cases have been described since the first recorded case in 1894), the only known cases of direct infections in man are three premature infants who contracted fatal cryptococcal meningencephalitis from cryptococci present in the maternal genital tract (Neuhauser and Tucker, 1948).

**SUMMARY**

A case of pulmonary cryptococcosis is recorded.

**REFERENCES**