FAMILIAL AND MULTIPLE PHAEOCHROMOCYTOMAS

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HYMAN and Mencher (1943) first reported the occurrence of phaeochromocytomas in more than one member of a single family. A familial incidence of these tumours has since been observed in at least seventeen apparently unrelated families (see Carman and Brashear, 1960 for earlier references; Smits and Huizinga, 1961; Cushman, 1962; Grunstein and Finkelstein, 1962; Tisherman, Gregg and Danowski, 1962; Hagen and Barrows, 1963). Approximately 9% of all patients with phaeochromocytomas harbour the tumours in both adrenal glands (Graham, 1951; Barbeau, Marc-Aurele, Brouillet, Yitke, Leboeuf, Cartier, Mignault and Genest, 1958; Hume, 1960), but the incidence of bilateral adrenal tumours rises to 24% in children and to about 50% in familial cases (Hume, 1960). We report here the occurrence of bilateral adrenal phaeochromocytomas in one patient and of a single adrenal phaeochrome tumour in his sister.

Methods

Urine was collected for 24 hours into a bottle containing 10 ml. of concentrated hydrochloric acid. The quantity of adrenaline and noradrenaline in urine and tumour tissue was estimated using the rat blood pressure preparation and the extraction procedures described by Euler & Hellner (1951) and Hamilton, Litchfield, Peart & Sowry (1953). Vanilmandelic acid (VMA) was estimated by the method of Sandler and Ruthven (1961).

Case Reports

Case No. 1: B. H. Male, aged 44 years. The earlier history of this male patient was reported in previous papers from this hospital (case 3, Barnett, Blacket, Depoorter, Sanderson and Wilson, 1950; case 2, Hamilton, Litchfield, Peart and Sowry, 1953). As a child, he experienced attacks of bradycardia and pallor lasting 2-3 minutes and when he was eleven years old, an electrocardiogram revealed a short P-R interval and the changes associated with left bundle branch block (case 7, Wolff, Parkinson and White, 1930). These symptoms subsided, but in 1941, when aged 23, he experienced attacks of epigastric discomfort followed by a slow thumping sensation in the chest, pallor and sometimes a throbbing headache. These symptoms varied in duration from about 2 to 15 minutes and occurred up to twelve times daily. When he was symptom-free the blood pressure varied from 120/86 to 136/90 mm. Hg.; during the attacks it increased to a maximum of 254/154. A plain X-ray of the abdomen in 1949 (Dr. E. Rohan Williams) demonstrated a round mass above the upper pole of the right kidney and in November of the same year, one of us (A.D.W.) removed the right suprarenal gland containing an encapsulated tumour. The tumour weighed 200 g.; brownish granules appeared within the cells after treatment with bichromate and histological examination confirmed the diagnosis of phaeochromocytoma (Dr. R. H. Heptinstall). Pharmacological investigations were not performed at this stage. The patient then remained symptom-free until October, 1957 when he experienced attacks of palpitation and frequent frontal headaches. These symptoms occurred less often during the next few years, but in July 1962 he again experienced attacks of slow, regular forceful palpitation associated with throbbing headache, pallor and a sense of “abdominal emptiness”. These lasted about five minutes, and occurred 2-3 times daily. A spontaneous attack was not observed, but his blood pressure varied between 100/60 and 170/100. Intravenous injection of histamine (20 mcg.) precipitated a typical paroxysm in which the blood pressure increased from 160/100 to 250/150 and ventricular ectopics appeared on the electrocardiogram.

Investigations: The urine was free of sugar and protein. During a twenty-four hour period without an attack, the urine contained 380 µg noradrenaline, 145 µg. adrenaline and 14.8 mg. V.M.A. Pre-sacral retro-peritoneal air insufflation followed by tomodiagnosis showed a spherical mass anterior to the upper pole of the left kidney (Dr. David Sutton). At operation on July 12th, 1962 (A.D.W.) the tumour was found surrounded by a thin rind of adrenal cortex which could not be saved. The tumour was 4 cm. in diameter, weighed 65 g., contained 6.1 mg. noradrenaline and 1.0 mg. adrenaline per g. wet weight.

Since the opposite adrenal gland had been removed previously, the patient was given cortisone by intramuscular injection on the day before, and hydrocortisone intravenously on the day of the operation. Two hours after the operation, the systolic blood pressure suddenly fell to 50 mm. Hg., and this was rapidly corrected by a brief noradrenaline infusion. The subsequent course was uneventful and he has remained symptom-free while continuing replacement therapy with cortisone (37.5 mg. daily) and 9 α-fluorohydrocortisone (0.1 mg. daily). In September, 1964 his blood pressure was 140/80 mm. Hg.
Case no. 2: E.H. Female, aged 48 years. This unmarried sister of case 1 was well until 1960 when she first noticed attacks of weakness, tremor, occipital headache and nausea without vomiting. Slight bradycardia and blurring of near vision occurred in some attacks. She had experienced many minor attacks lasting 10-15 minutes and seven severe paroxysms which persisted for about an hour. Small attacks were sometimes provoked by twisting the trunk to the right.

Examination showed a rather thin, alert woman whose blood pressure was 145/90 and pulse rate regular at 70 per min. Spontaneous attacks did not occur while the patient was in hospital but intravenous injection of histamine (20 mcg) precipitated a typical paroxysm during which the pulse became irregular at a rate of 130 per min, and the blood pressure increased from 145/90 to 225/150.

Investigations. There was no sugar or protein in the urine and the blood sugar, haemoglobin, plasma electrolytes, chest X-ray and intravenous pyelogram were normal. During a twenty-four hour period without an attack, the urine contained 100 µg noradrenaline, 50 µg of adrenaline and 12.6 mg. of VMA. These results are discussed in a separate communication (Brown, Ruthven and Sandler, 1966). 25 µg. of noradrenaline and 45 µg. of adrenaline were excreted in four hours after the attack precipitated by histamine. Pre-sacral retroperitoneal pneumography followed by tomography demonstrated a spherical mass anterior to the upper pole of the left kidney. (Dr. E. Rohan Williams). At operation in March, 1962 the tumour was removed without the rest of the gland to which it was attached by a thin bridge of tissue. The tumour weighed 30 g., and contained 20 µg. noradrenaline and 50 µg. adrenaline per g., wet weight. The post-operative course was uneventful and 19 days later a twenty-four hour urine specimen contained 60 µg. noradrenaline, 20 µg. adrenaline and 4.5 µg. VMA. (i.e., all within the normal range). Attacks have not occurred since the operation.

Discussion

In 1957, the world literature contained records of 626 patients with phaeochromocytoma (Barbeau, 1957). The present report brings the total number of familial patients with phaeochromocytomas to at least fifty and the affected kindred to eighteen. A family history of these tumours, therefore, occurs in 7-8% of the published series, but the true incidence is probably lower since familial cases are more likely to be recorded. The incidence of phaeochromocytomas is increased in patients with neurofibromatosis and possibly in cases with other hereditary neocutaneous syndromes (Glushien, Mansuy and Littman, 1953; Chapman, Kemp and Taliaferro, 1959), but evidence of these diseases were not found in our cases or in their relatives. Smits and Huizinga (1961) found four patients with phaeochromocytomas and ten other cases with a history of characteristic attacks, in a single Dutch family. In seven of the seventeen published family pedigrees, phaeochromocytomas occurred in a parent and sibling and there was a strong clinical suspicion of these tumours in the father or mother of five other cases. The mother of the two cases reported here has persistent hypertension but she denied any paroxysmal symptoms. Phaeochromocytomas may present in this way (Thorn, Hindle and Sandmeyer, 1944; Goldenberg, Aranow, Smith and Faber, 1950; de Graeff and Horak, 1963), but the diagnosis seems unlikely in this patient since the excretion of catechol amines was normal (Table 1). Although the urinary catechol amines may be normal between (Pitkanen, 1956; Thompson and Trounce, 1958) or during (Litchfield and Peart, 1956) paroxysmal hypertensive attacks, as far as we are aware increased excretion has always been found in patients with a phaeochromocytoma and persistent hypertension. The father of this family has carcinoma of the prostate gland. He denied symptoms suggestive of phaeochromocytoma and his blood pressure was 170/100 mm. Hg. Although carcinoma of the thyroid gland is more common in patients with phaeochromocytoma (Sipple, 1961), we are not aware of a similar relationship between adrenal and prostatic tumours. The two children of B.H. are symptom-free and the excretion of catechol amines and VMA was normal in both cases (Table 1). However, the average age at the time of diagnosis in familial cases is twenty-six years. The surprisingly high incidence of familial phaeochromocytomas has led to the suggestion that this is a hereditary trait with a dominant autosomal mode of inheritance (Carman and Brashear, 1960; Smits and Huizinga, 1961; Cushman, 1962). If this is valid, incomplete penetrance of the gene might explain the absence of tumours from some generations, while truly sporadic cases could be caused by mutation. In this connection, it is noteworthy that neurofibromatosis, which is often associated with a phaeochromocytoma, appears to arise by mutation more frequently than most hereditary traits (Crowe, Schull and Neel, 1956).

Although bilateral adrenal phaeochromocytomas occur in approximately 50% of patients with a family history of this disease (Hume, 1960), the long symptom-free period after the first operation in patient B.H. suggests that the second tumour was a more recent development. However, the frequency of attacks varies and a spontaneous remission
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TABLE 1.
URINARY EXCRETION OF NORADRENALINE, ADRENALINE AND VMA

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<tr>
<th></th>
<th>Excretion in Urine/24 Hours</th>
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<tr>
<td></td>
<td>Noradrenaline µg</td>
</tr>
<tr>
<td>Case B.H.</td>
<td>Before Operation</td>
</tr>
<tr>
<td>Wife</td>
<td>380</td>
</tr>
<tr>
<td>Son</td>
<td>33</td>
</tr>
<tr>
<td>Daughter</td>
<td>35</td>
</tr>
<tr>
<td>Father</td>
<td>12</td>
</tr>
<tr>
<td>Mother</td>
<td>18</td>
</tr>
<tr>
<td>Sister</td>
<td>30</td>
</tr>
<tr>
<td>(Case E.H.)</td>
<td>Before Operation</td>
</tr>
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<td></td>
<td>15</td>
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<thead>
<tr>
<th></th>
<th>Excretion in Urine/24 Hours</th>
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<tbody>
<tr>
<td></td>
<td>Noradrenaline µg</td>
</tr>
<tr>
<td></td>
<td>100</td>
</tr>
<tr>
<td></td>
<td>60</td>
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</tbody>
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* not estimated

0—not detected

Haemangiomata, Amer. J. med., 26, 883.


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Familial and multiple phaeochromocytomas.

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