Russell's diencephalic syndrome of early childhood

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SUMMARY A case of Russell's diencephalic syndrome of early childhood is presented and the world literature reviewed. Assays of serum growth hormone (HGH) and fat mobilizing substance (FMS) were performed and gave very abnormal results. The role of these two substances is discussed and their value in the diagnosis of this syndrome is emphasized.

Within the last two decades interest has been focused on the diencephalon and its pathology, and it is now known that discrete organic lesions of the hypothalamus and adjacent structures may produce characteristic symptoms. Thus, it has been known for very many years that posteriorly placed tumours of the hypothalamus may be associated with precocious puberty, somnolence, obesity, diabetes insipidus, dysthermia, and various emotional disturbances. Some conditions, however, such as leprechaunism (Donohue's syndrome), cerebral gigantism (Soto's syndrome), generalized lipodystrophy with gigantism, and possibly progeria (Hutchinson-Gilford syndrome) may be due to a functional disorder of the diencephalon rather than to a discrete organic lesion within one or more of its component structures. It was not, however, until 1951 that Russell described an apparently specific syndrome of early childhood resulting from a neoplastic lesion of the anterior hypothalamus, and not until 1957 that Lorimer Dods became the second author to report further examples of the syndrome. Russell would appear to be the only author to report cases from the United Kingdom.

Russell's syndrome is probably much more common than the 50 cases in the literature would suggest and the purpose of this paper is to describe a further case from the United Kingdom with the results of relevant investigations and treatment.

CASE REPORT

S.H., a girl aged 23 months, was the first child of elderly parents (maternal age, 43 years; paternal age, 45 years). She presented with a three month history of rapid, progressive weight loss and sleep reversal. The mother had been a ward sister on a well-known neurological unit. The first 20 months of life had been uncomplicated but after this time there was a dramatic change (Figs 1 and 2). In spite of an apparently normal dietary intake she rapidly lost weight (Fig. 3) and at the same time became hyperactive by night with long periods of sleep by day. While in the ward she appeared affectionate, without aggression, but hyperkinetic. She talked and sang to herself incessantly and demonstrated slight ataxia on
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turning and reaching out for objects. Clinical examination revealed, apart from gross marasmus, nystagmus and numerous café-au-lait patches over her upper trunk. Her body temperature was consistently about 35.6°C (96°F) and never exceeded this. There was no polydipsia or polyuria and her appetite was usually normal. Her height and weight, from measurements kept by the parents, both followed the 97th percentile, but the weight dropped to well below the 3rd percentile in just over three months. During this period of time her height did not leave the 97th percentile.

INVESTIGATIONS The results of the following investigations were within normal limits: Haemoglobin, sedimentation rate, urea, plasma electrolytes, cholesterol, calcium, phosphorus, alkaline phosphatase, plasma proteins, random and fasting blood glucose, D-xylose absorption test, serum lead, protein bound iodine, and electroencephalograph. Urine specific gravity was consistently over 1.010.

**Abnormal results** The cerebrospinal fluid contained protein 100 mg/100 ml. A pneumoencephalogram showed a large and mainly left-sided anterior hypothalamic tumour (Figs 4 and 5) indenting the third ventricle.

The cisterna lamina terminalis was displaced forwards and the interpeduncular system was obliterated. The tumour also encroached on the upper part of the prepontine cisterns and displaced the lower margin of the left frontal horn upwards and forwards. The aqueduct was rather more upright than normal and the midbrain was displaced backwards.

Serum growth hormone (HGH) was assayed using a double antibody radioimmunoassay technique as described by Jackson, Grant, and Clayton (1968); HGH production was stimulated with oral Bovril (20 g/1.5 m² given in 160 ml warm water) and estimated in venous blood samples taken from an indwelling intravenous cannula. The results obtained were comparable with those expected in an adult subject with acromegaly (Table 1). A similar assay was performed eight months after radiotherapy.

Urinary fat mobilizing substance (FMS) was assayed by the method of Pawan (1969). Plasma levels of ketones (as acetone) and free fatty acids (FFA) were measured in mice injected with prepared...
FIG. 4. Pneumoencephalogram to show distortion of third ventricle.

FIG. 5. Pneumoencephalogram to show distortion of ventricular system by anterior hypothalamic tumour.
TABLE 1
SERUM GROWTH HORMONE (HGH) ASSAY

<table>
<thead>
<tr>
<th>Time (min)</th>
<th>Normal values (µU/ml.)</th>
<th>Before treatment (µU/ml.)</th>
<th>Three months after radiotherapy (µU/ml.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fasting</td>
<td>0-10</td>
<td>93.5</td>
<td>48.8</td>
</tr>
<tr>
<td>30</td>
<td></td>
<td>&gt; 200</td>
<td>68.8</td>
</tr>
<tr>
<td>60</td>
<td>maximum rise</td>
<td>&gt; 200</td>
<td>71.6</td>
</tr>
<tr>
<td>90</td>
<td>30-40</td>
<td>&gt; 165</td>
<td>54.8</td>
</tr>
<tr>
<td>120</td>
<td></td>
<td>&gt; 143</td>
<td>40.0</td>
</tr>
<tr>
<td>150</td>
<td></td>
<td>84.5</td>
<td>27.4</td>
</tr>
</tbody>
</table>

urine (see Kekwick and Pawan, 1967) and compared with results obtained from a control series of mice injected with saline. From the increase in plasma ketones the assay revealed a very great deal (74% increase) of FMS activity. Unfortunately, the assay was performed three months after radiotherapy and it would have been interesting to have measured FMS activity before treatment.

Ketones (as acetone) | Free fatty acids
Control mice 2.21 mg/100 ml. | 558 µE/l.
Test mice 3.85 mg/100 ml. (74% inc.) | 599 µE/l. (7% inc.)

At the time of this assay the patient was on a normal diet and eating well. In Pawan’s series of 72 normal subjects there was no percentage increase of ketones or free fatty acids in mouse plasma, and a similar finding was reported in six subjects after pituitary ablation.

TREATMENT AND PROGRESS Once the diagnosis of a large anterior hypothalamic tumour was confirmed radiotherapy was given as surgery was considered impossible. A total of 4,650 r were delivered by cobalt γ-ray therapy to the tumour over 19 sessions (44 days). After this course of radiation a slight improvement was noticed in the succeeding months. Three months after treatment she had gained a little weight and was much less hyperkinetic. There was, however, a sinister decrease in the child’s visual acuity and it is likely that there is considerable involvement of the optic chiasma. Formal perimetry was not attempted.

DISCUSSION

The case described conforms to the original description by Russell (1951) of the diencephalic syndrome which bears his name. In reviewing the world literature there is found to be a great similarity between the described cases and the definition of a specific syndrome must be confirmed (Table 2).

Children with Russell’s diencephalic syndrome commonly present in the first year of life and probably never after the age of 5 years. The youngest reported cases were those of Dods (1957) and Smith, Weinburg, and McAlister (1964) who presented at the age of 2 months.

In virtually all the cases profound emaciation with hyperactivity, apparent well-being, and euphoria were prominent and, apart from nystagmus, there was a gross paucity of neurological signs. In no case was there evidence of raised intracranial pressure, but extension of the tumour to involve the optic discs was present in about one fifth of the cases. Paradoxical obesity and increased height occurred in a case described by Fishman and Peake (1970) and was thought to be secondary to extension of the tumour to involve hypothalamic centres regulating food intake. Similar changes occurred in one of the cases described by Gamstorp, Kjellman, and Palmgren (1967) after partial surgical removal of an anterior hypothalamic tumour.

Apart from gross, progressive subcutaneous wasting, these children exhibit a curious euphoria and lack of aggression which Spiegel, Wycis, Marks, and Lee (1947) attribute to interruption of the frontothalamic-hypothalamic pathway or even to discrete involvement, by the tumour, of the dorsomedial thalamic nuclei.

Growth hormone (HGH) assay has been made in only two patients other than that presented here, but the exact levels of HGH in one of these (Smith et al., 1964) is not recorded. In both cases the levels were markedly raised. In the present case the levels recorded, both in the fasting state and after stimulation with Bovril

TABLE 2
REPORTED FINDINGS IN 29 CASES OF RUSSELL’S DIENCEPHALIC SYNDROME

1. Profound cachexia 29/29
2. Sex incidence 18/117
3. Euphoria mentioned in 16/29
4. Hyperactivity mentioned in 20/29
5. Nystagmus mentioned in 16/29
6. Optic atrophy 9/29
7. Pallor without anaemia mentioned in 14/29
8. Anterior hypothalamic tumour 29/29
9. Age at presentation 2-32 months
(Jackson et al., 1968) were greatly raised and equivalent to those expected in adults with acromegaly. These high levels were not associated with any increase in height although sustained high levels of growth hormone should increase linear growth (Daughaday, 1968). Three months after treatment with radiotherapy a second HGH assay was performed and showed a substantial drop in HGH levels when compared with the first assay. The levels were, however, still above those expected in normal subjects. The effect of excess HGH production in this syndrome is not fully understood but it is probable that its described lipolytic action (Zierler and Rabinowitz, 1963) is, in part, responsible for dispersing subcutaneous fat, which Poznanski and Manson (1963) showed to be completely absent. Similar loss of subcutaneous fat is seen in other conditions of presumed diencephalic origin—namely, leprechaunism (Donohue and Uchida, 1954; Salmon and Webb, 1963), progeria (Schinz, Baensch, Friedl, and Uehlinger, 1952; Thomson and Forfar, 1950), and generalized lipodystrophy (Berardinelli, 1954; Seip and Trygstad, 1963; Fairney, Lewis, and Cottom, 1969). It is of interest to recall that Villee, Nichols, and Talbot (1969) showed that growth in progeria was also unresponsive to HGH. In leprechaunism the only report of HGH assay is an isolated measurement reported by Dekaban (1965) which is at the upper level of the normal fasting range (1–10 μIU/ml). The total lack of subcutaneous fat in Russell's diencephalic syndrome may be partly attributable to increased production of HGH, but the lipid mobilizing fraction (LM) isolated by Zarafonitis, Seifler, Baeder, and Kalas (1959) from the pituitary and the fat mobilizing polypeptide (FMS) isolated from urine by Chalmers, Pawan, and Kekwick (1960) may also play their part. These last two substances may be quite unrelated to HGH but Pawan (1971) suggests that FMS may in fact prove to be a degradation product of HGH. FMS activity was estimated in the present case and found to be significantly raised. The estimation was made three months after radiotherapy had produced clinical improvement and a slight gain in weight. At this time HGH assay (Table 1) revealed a considerable drop from the pretreatment levels. It would be of the greatest interest to assay FMS before treatment in this syndrome.

In 1960 Murray found that specific stimulation of the anterior hypothalamus in dogs produced fluctuating levels of blood lipids. Since all pituitary hormones are thought to be under hypothalamic control, it is possible that lipid depletion may result from the action, direct or otherwise, of a specific humoral substance produced by the presence of the anterior hypothalamic tumour in this syndrome.

With the exception of one case seen by Dods (1957) where a clinical picture resembling Russell's syndrome resulted from a midline cerebellar astrocytoma, all the lesions were accurately located in the anterior hypothalamus and extended upwards to displace the floor of the third ventricle. Extension forwards to involve the optic tracts was common. The tumours were commonly astrocytomas but oligodendrogliomas, spongiosblastomas, and ependymomas have all been recorded. In one case (Braun and Forney, 1959) a perforated duodenal ulcer with massive haemorrhage complicated the main presentation, although we should recall that the association of peptic ulceration with intracranial neoplasm was noticed as long ago as 1850 by Rokitansky and again in 1932 by Cushing.

The presentation of Russell's syndrome is so stereotyped that diagnosis should not be difficult. The profound cachexia with hyperactivity and a happy affectionate child should be contrasted with the cachectic picture of chronic starvation, severe malabsorption, gastroenteritis, neoplasia in general, anorexia nervosa, thyrotoxicosis, and chronic diabetes.

The absence of neurological signs is important, although nystagmus may be present. Air studies and growth hormone assay should confirm the diagnosis. Treatment is at the best palliative and neurosurgery may be impossible. Fishman and Peake (1970) record a six year survival after radiotherapy in one case, but few patients survive one year after the initial diagnosis.

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