Granulomatous hypophysitis caused by Rathke’s cleft cyst mimicking a growth hormone-secreting pituitary adenoma

Masato Hojo, Ryota Ishibashi, Hiroshi Arai, Susumu Miyamoto
Departments of Neurosurgery and Medicine and Clinical Science, Kyoto University Graduate School of Medicine, Kyoto 606-8507, Japan

ABSTRACT
We report a case of granulomatous hypophysitis caused by Rathke’s cleft cyst (RCC) mimicking a growth hormone (GH)-secreting pituitary adenoma. Neuroradiological and endocrinological evaluations showed abnormal findings consistent with acromegaly: Magnetic resonance imaging demonstrated a pituitary mass lesion, and GH and insulin-like growth factor I levels were markedly elevated, and GH levels were not suppressed in oral glucose tolerance test. Transsphenoidal surgery was performed, but no adenomatous tissue could be detected. Histological examination revealed RCC and concurrent granulomatous giant cell inflammatory reaction of the anterior hypophysis. To the authors’ knowledge, this is the first documented case of granulomatous hypophysitis caused by RCC mimicking a GH-secreting pituitary adenoma.

Key words: Acromegaly, granulomatous hypophysitis, growth hormone, insulin-like growth factor I, Rathke’s cleft cyst

Introduction
Granulomatous hypophysitis is a rare inflammatory disorder of the pituitary gland; it is pathologically characterized by granulomatous reaction with epithelioid histiocytes, lymphocytes, plasma cells, and multinucleated giant cells. This pituitary lesion is distinct from systemic granulomatous disorders such as sarcoidosis, tuberculosis, and syphilis, and has two possible different etiologies: Autoimmune disorder and secondary inflammation. Although Rathke’s cleft cyst (RCC) is one of the causes of pituitary inflammation, granulomatous hypophysitis caused by RCC is a very rare pituitary disorder. Moreover, the predominant endocrinological feature of granulomatous hypophysitis is hypopituitarism, and no case with growth hormone (GH) hypersecretion has been previously reported. Here, we report an extremely rare case of granulomatous hypophysitis caused by RCC, which endocrinologically and neuroradiologically mimicked a GH-secreting pituitary adenoma.

Case Report
A 22-year-old man presented with a 2-month history of gradual onset of headache, excessive sweating, fatigue, and gonadal dysfunction. The characteristic facial features of acromegaly were not apparent. Magnetic resonance imaging (MRI) demonstrated a pituitary mass lesion, and GH and insulin-like growth factor I levels were markedly elevated, and GH levels were not suppressed in oral glucose tolerance test. Transsphenoidal surgery was performed, but no adenomatous tissue could be detected. Histological examination revealed RCC and concurrent granulomatous giant cell inflammatory reaction of the anterior hypophysis. To the authors’ knowledge, this is the first documented case of granulomatous hypophysitis caused by RCC mimicking a GH-secreting pituitary adenoma.

Address for correspondence:
Dr. Masato Hojo, Department of Neurosurgery, Kyoto University Graduate School of Medicine, 54 Kawahara-cho, Shogoin, Sakyo-ku, Kyoto 606-8507, Japan.
E-mail: mhojo@kuhp.kyoto-u.ac.jp

evaluations showed that the GH level was increased: 5.42 µg/L (normal value, 0.04-0.34 µg/L), and the insulin-like growth factor I (IGF-I) level was also elevated: 134.9 nmol/L (normal value, 13.9-52.1 nmol/L). In oral glucose tolerance test (OGTT), GH levels were not suppressed [Table 1]. These findings were consistent with acromegaly. Mild adrenal dysfunction and diabetes insipidus were proved, and replacement therapy was started.

Two months after the first visit, transsphenoidal surgery was performed. The operation was performed via a sublabial transsphenoidal approach. After dural opening, the anterior pituitary gland was encountered. When this anterior gland was incised, white creamy content poured out [Figure 2a]. The cyst wall and the anterior lobe were incised and sent for histological examination. The biopsy specimens (3 mm in size) were obtained in different three sites. Although we inspected the lesion meticulously and carefully, we could not detect any adenomatous tissue.

Histological examination revealed an epithelial cyst wall consistent with RCC [Figure 2b]. No neoplasm could be detected. The pituitary tissue was infiltrated by mature lymphocytes, plasma cells, and neutrophils. The surrounding tissue was infiltrated by plasma cells, neutrophils, and multinucleated giant cells consistent with granulomatous giant cell inflammatory reaction of the anterior pituitary gland [Figure 2c]. Immunohistochemistry showed that GH-positive cells were increased in limited areas, and that positive staining for GH was also observed in the extracellular space [Figure 2d]. This finding suggested the possibility that somatotrophs might be stimulated and destructed due to inflammation and that GH might be released resulting in hypersecretion. In immunohistochemical studies, the cyst wall did not contain endocrine cells scattered between ciliated and mucin-producing cells. We concluded, therefore, that this pituitary lesion was granulomatous hypophysitis caused by RCC, but not pituitary adenoma, and GH hypersecretion was induced by pituitary inflammation.

The postoperative course was uneventful. On the next day after surgery, the IGF-I level had already improved: 37.6 nmol/L [Table 2]. Eight days after surgery, GH levels were suppressed to <2.0 µg/L in OGTT [Tables 1 and 2]. The visual field defect was resolved completely. Postoperative MRI revealed that the cyst was evacuated, but the pituitary stalk was thickened and heavily enhanced [Figure 3a]. Fifty days after surgery, GH levels were suppressed to <1.0 µg/L in OGTT [Tables 1 and 2]. 1-year after surgery, MRI demonstrated that the thickened and enhanced pituitary stalk was normalized [Figure 3b]. GH and IGF-I levels were also normal: 0.75 µg/L and 19.8 nmol/L, respectively. Hypopituitarism was gradually resolved except for diabetes insipidus. The patient is doing well on replacement therapy with only nasal vasopressin 6 years after surgery, and GH and IGF-I levels remain normal for these 6 years.

Table 1: GH levels in oral glucose tolerance test

<table>
<thead>
<tr>
<th>Time (min)</th>
<th>Preoperative</th>
<th>Postoperative day 8</th>
<th>Postoperative day 50</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>Glucose (mmol/L)</td>
<td>GH (µg/L)</td>
<td>Glucose (mmol/L)</td>
</tr>
<tr>
<td>0</td>
<td>4.72</td>
<td>8.47</td>
<td>3.77</td>
</tr>
<tr>
<td>30</td>
<td>6.72</td>
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<tr>
<td>90</td>
<td>4.55</td>
<td>4.11</td>
<td>5.94</td>
</tr>
<tr>
<td>120</td>
<td>5.99</td>
<td>3.14</td>
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</tr>
</tbody>
</table>

GH – Growth hormone. Nadir GH levels are indicated by boldface.
Discussion

It has been previously reported that RCC can be one of the causes of inflammatory reaction of the pituitary gland.[9,10] But granulomatous hypophysitis caused by RCC is a rare pituitary inflammatory disorder, and only four cases have been previously reported.[4,6-8] However, all four cases presented with hypopituitarism and were not associated with GH hypersecretion.[10,11] In contrast, our case is associated with GH hypersecretion and mimicked a GH-secreting pituitary adenoma. Since GH exhibits a prominent pulsatile secretion pattern, randomly drawn blood samples are not appropriate for the evaluation of GH-secretion pattern and measurement of IGF-I is necessary for the evaluation of GH hypersecretion.[15] Previously, only one case of lymphocytic hypophysitis with slightly elevated IGF-I level was reported.[16] In this case, the IGF-I level was slightly elevated: 62.5 nmol/L (normal value, 16.1-60.7 nmol/L).[16] In contrast, in our case, the IGF-I level was markedly elevated: 134.9 nmol/L (normal value, 13.9-52.1 nmol/L). To our knowledge, our case is the first report of granulomatous hypophysitis caused by RCC associated with severe GH hypersecretion.

In the present case, the elevated IGF-I level had already been normalized by the next day after surgery [Table 2]. Since it usually takes several months for the IGF-I level to be normalized after normalization of GH-secretion, it is unlikely that the IGF-I level was normalized by the direct effect of the surgery. According to this reason, it is highly unlikely that hidden GH producing pituitary adenoma had induced GH hypersecretion. Therefore, in the present case, it is considered highly likely that GH hypersecretion was induced by inflammation of the pituitary gland and normalization of GH-secretion was due to improvement of pituitary inflammation. We propose two possible mechanisms for GH hypersecretion in the present case. One possible mechanism is that destruction of somatotrophs by the inflammatory process may result in escape of hormone into the circulation. In the present case, immunohistochemical staining for GH demonstrated that positive staining for GH was observed in the extracellular space, suggesting the validity of our estimation. The other possible mechanism of GH hypersecretion is that cytokines secreted from lymphocytes may stimulate somatotrophs to overproduce GH. In vitro studies have shown that cytokines such as interleukin (IL)-6 and IL-1bera promote GH-secretion.[17,18] These in vitro data support our speculation on the mechanism of excessive GH production.

Financial support and sponsorship
Nil.

Conflicts of interest
There are no conflicts of interest.

Table 2: Clinical course, and changes of nadir GH levels in OGTT and IGF-I levels

<table>
<thead>
<tr>
<th></th>
<th>Nadir GH* (µg/L) (≤1.00 µg/L)</th>
<th>IGF-I (nmol/L) (13.9-52.1 nmol/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Postoperative day</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(days)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>−50</td>
<td>3.14</td>
<td>134.9</td>
</tr>
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<tr>
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<td>surgery)</td>
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<td>32.8</td>
</tr>
<tr>
<td>50</td>
<td>0.32</td>
<td>35.8</td>
</tr>
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</table>

*Nadir GH levels in OGTT. GH – Growth hormone; IGF-I – Insulin-like growth factor I; OGTT – Oral glucose tolerance test. Normal values are given in brackets.

References

Hojo, et al.: Granulomatous hypophysitis mimicking a GH-secreting pituitary adenoma


