Use of the somatostatin analogue octreotide to localise and manage somatostatin-producing tumours

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Abstract

Background—Somatostatin receptor scintigraphy (SRS) and octreotide therapy have both changed the management of gastroenteropancreatic endocrine tumours, but very few data are available on the use of SRS and octreotide to visualise and treat somatostatinomas.

Method—The results of SRS and octreotide treatment in three somatostatinoma patients were examined.

Results—SRS was able to detect extensive hepatic involvement in patient 1, one hepatic and one pancreatic lesion in patient 2, and one hepatic lesion in patient 3. Octreotide therapy (0.5 mg/day subcutaneously) was effective in decreasing plasma levels of somatostatin in all three patients. Symptoms (diabetes and diarrhoea) were greatly improved in the two patients with “somatostatinoma syndrome”.

Conclusion—The study shows that somatostatinoma, like most other gastroenteropancreatic endocrine tumours, possesses functioning somatostatin receptors.

Keywords: somatostatin; octreotide; somatostatin receptor scintigraphy; gastroenteropancreatic endocrine tumour; somatostatinoma

Somatostatinoma is a rare malignant somatostatin (SS)-producing tumour; fewer than 100 cases have been described accounting for less than 1% of all gut and pancreatic endocrine tumours. Some 75% of somatostatinomas are localised in the pancreas, and most non-pancreatic tumours are found in the duodenum and jejunum. Overall, more than 85% of the cases have displayed metastases, located mainly in the liver and lymph nodes. The first cases reported were found by accident, and the diagnosis of somatostatinoma is generally based on postoperative histological examination. Preoperative diagnosis of a somatostatinoma remains difficult because the so called “somatostatinoma syndrome” (diabetes mellitus, gallbladder disease, diarrhoea, and weight loss) actually produces non-specific symptoms. Furthermore, for unknown reasons, the syndrome occurs only in the pancreas; the duodenal location of primary tumour is apparently associated with “local discomfort” or is completely symptom-free. The histological characteristics of pancreatic somatostatinomas are indistinguishable from those of other pancreatic endocrine tumours. The diagnosis is therefore established by immunocytochemical examination of the resected tissue with typical endocrine tumour features and a predominant or pure population of “D” cells. Often a significantly increased concentration of SS-like immunoreactivity in the plasma and/or tumoral extracts further supports the diagnosis of somatostatinoma in many studies.

Over the last few years, octreotide, a synthetic SS analogue, has drawn a lot of attention mainly because of its antisecretory and antiproliferative effects on SS receptor bearing tumours. Recently a modified analogue of octreotide, labelled with indium 111 (Octreoscan), has been used to localise SS receptor positive tumours in vivo. To our knowledge, very few data are available on octreotide treatment of somatostatinoma and/or tumour lesion detection with Octreoscan, although it is known that the “D” cells possess SS receptors. In the present study we report on three patients with SS-producing tumour (two pancreatic, one duodenal), treated long term with octreotide after evaluation of SS receptor status using SS receptor scintigraphy (SRS).

Patients

Fifty two consecutive patients referred to our unit with a diagnosis of a gastroenteropancreatic tumour were evaluated for somatostatinomas. In three of the patients, a somatostatinoma was diagnosed on the basis of high basal circulating levels of SS, increased “D” cell number in tumour tissue specimens, and, in two of them, the presence of somatostatinoma syndrome (table 1).

Table 1 Clinical and biochemical data of the three patients with somatostatinoma

<table>
<thead>
<tr>
<th>Patient</th>
<th>Age (y)</th>
<th>Sex</th>
<th>Gallstone disease</th>
<th>Diabetes type</th>
<th>Diarrhoea</th>
<th>Weight loss</th>
<th>SS (pg/ml)</th>
<th>NSE (g/l)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patient 1</td>
<td>75</td>
<td>F</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>67.5</td>
<td>22</td>
</tr>
<tr>
<td>Patient 2</td>
<td>61</td>
<td>F</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>65</td>
<td>24</td>
</tr>
<tr>
<td>Patient 3</td>
<td>60</td>
<td>M</td>
<td>Yes (sludge)</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>200</td>
<td>32</td>
</tr>
</tbody>
</table>

SS, somatostatin; NSE, neuron specific enolase. Normal values: SS <25 pg/ml; NSE <12 g/l.
PATIENT 1
A 75 year old woman was admitted to the hospital in poor general condition with type II diabetes, diarrhoea, and mild weight loss (9 kg over the preceding six months). She reported that gallbladder stones had been diagnosed within the previous year. On admission, stool weight was 605 g per day with an elevated amount of fat (21 g/day). Basal plasma levels of SS were increased as well as serum NSE (table 1). Abdominal ultrasound showed a mass in the tail of the pancreas (diameter 4 cm) and another nodule in the liver. All these findings were further confirmed by CT scan (table 2). Histological examination of a liver biopsy specimen showed typical structures of a pancreatic endocrine tumour consistent with a pancreatic primary. Immunocytochemical staining showed diffuse positivity for SS and NSE. SRS confirmed the number and location of the tumoral lesions visualised by CT scan (table 2). The patient refused surgery, and treatment with octreotide (0.5 mg/day subcutaneously) was initiated.

PATIENT 2
A 60 year old man was referred to us after having undergone pancreatic duodenectomy (Whipple procedure) to remove a 4.6 × 3.3 cm head of pancreas mass and a parietal nodule of the second portion of the duodenum. The histological appearance of the tumour was typical of a duodenal somatostatinoma, showing an endocrine tumour with numerous glandular structures often with a psammoma body in the lumen (fig 1). The tumour, originating from the duodenal mucosa, was extensively infiltrating the underlying muscular wall and the contiguous pancreatic region. On immunocytochemistry, virtually all tumour cells exhibited intense immunostaining for SS (fig 1).

On admission, the patient was in good general condition and reported no specific symptoms. Plasma levels of SS and NSE were considerably elevated (table 1). Abdominal ultrasound showed gallbladder sludge with microstones, and a CT scan showed no tumoral lesions (table 2). SRS showed an area of intense uptake in the left lobe of the liver (table 2). Octreotide treatment (0.5 mg/day subcutaneously) was started.

PATIENT 3
A 61 year old woman was hospitalised as the result of the sudden development of type II diabetes, diarrhoea, and moderate weight loss (3.3 kg over the preceding six months). She had been diagnosed with diabetes mellitus for more than one year. On admission, the patient was in good general condition, and reported no specific symptoms. Plasma levels of SS and NSE were improved after six months of octreotide treatment and remained improved at the one year evaluation. Octreotide therapy was effective in progressively decreasing plasma levels of SS in all three patients by 40–80% after one year.

Discussion
This study describes three patients with a gastroenteropancreatic endocrine tumour according to all diagnostic criteria (clinical, biochemical, and histological) for defining an
SS-producing tumour. Furthermore, this study suggests that somatostatinoma tumour tissue possesses functioning SS receptors. This suggestion is supported by the ability of \(^{111}\)In-Octreoscan scintigraphy to detect somatostatinoma tumour lesions in all three patients studied and by the effectiveness of octreotide in reducing circulating levels of SS and controlling related symptoms both in primary and/or metastatic somatostatinoma patients.

Previous studies, using in vitro and in vivo binding techniques, have shown the presence of SS receptors in 80% of the gastroenteropancreatic endocrine tumours.\(^{10}\) Few data are available on evaluation in vivo of SS receptors in somatostatinoma tumour tissue.\(^{11}\) In one case, treatment with octreotide did not reduce comparatively high circulating SS levels possibly because of receptor tachyphylaxis, as suggested by the authors.\(^{8}\)

In the present study, an octreotide-like compound was able to bind and visualise the SS receptors expressed by somatostatinoma tumour tissues in patients with abnormally high plasma levels of SS, and therefore the high SS levels cannot be the cause of the false negative results. There are various possible explanations for these data. Firstly, the circulating SS levels may not be high enough to interfere with binding of the SS analogue; moreover octreotide may have a higher affinity for tumoral SS receptors than the circulating SS. Secondly, the tumour may produce forms of SS that are not biologically active—for instance, prohormones—and enzymic degradation may affect natural SS but not synthetic SS analogues.\(^{12}\) Furthermore, octreotide treatment was effective in reducing plasma levels of SS and related symptoms in the two patients with the syndrome, whereas, in patient 3, SRS failed to visualise two of the three immunohistochemically proven hepatic somatostatinoma lesions, suggesting the presence of a different SS receptor subtype pattern on some (the ones that gave negative results on scintigraphy) metastatic lesions.\(^{13}\)

It has also been reported that injection of tolbutamide or calcium/pentagastrin is able to stimulate SS secretion only in somatostatinoma patients with normal basal levels of SS and not in normal controls.\(^{14}\) We did not try any stimulation tests as our patients had elevated basal SS levels.

Although the reasons for the false negative SRS results for patient 3 are not known, as previously reported, differences in blood supply\(^{15}\) and locally high concentrations of endogenous SS with alterations in receptor function have been reported to play an important role in SRS false negative gastroenteropancreatic endocrine tumour visualisation. Our data indicate that SRS, a non-invasive diagnostic procedure, is able successfully to localise primary and metastatic somatostatinoma tumours. Our results suggest that \(^{111}\)In-pentetreotide-SPECT, as shown for other gastroenteropancreatic endocrine tumours,\(^{7}\) may be a useful early diagnostic technique for accurately staging tumour extension in patients with somatostatinoma. Moreover, because of its ability to visualise SS receptors in vivo, SRS is a useful tool for selecting those patients with somatostatinoma syndrome who will benefit from octreotide treatment.

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