CLINICAL STUDY

Long-acting somatostatin analogues are an effective treatment for type 1 gastric carcinoid tumours

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Abstract

Background: Gastric carcinoid tumours type 1 (GCA1) originate from hyperplastic enterochromaffin-like (ECL) cells secondary to hypergastrinaemia. Treatment with somatostatin analogues (SSA) might impede ECL-cell hyperplasia by suppressing gastrin secretion and/or by a direct anti-proliferative effect on ECL cells. We conducted a multicentre prospective study to assess the effects of long-acting SSA on hypergastrinaemia and ECL-cell proliferation in patients with GCA1.

Methods: We studied 15 patients with GCA1 treated with monthly long-acting release octreotide (LAR) (20–30 mg; n = 14) or Lanreotide 90 mg (n = 1) for at least 6 months. Patients had serum gastrin and chromogranin A measurements performed and biopsies taken from both tumours and surrounding mucosa before, and every 6–12 months following treatment. Sections were immunostained for neuroendocrine markers. The cell proliferation index Ki-67, intensity of staining before and after treatment and the degree of gastric wall invasion were also assessed.

Results: All patients tolerated treatment well (mean follow-up of 18 months). In 11 patients (73%), a complete disappearance of the tumours at 1 year of treatment was observed on endoscopy, while in three patients (20%), the tumours decreased significantly in number and size. Gastrin levels normalized in 25% of patients, and were reduced by more than 80% in the remaining 75%.

Conclusions: Treatment with SSAs in GCA1 leads to a substantial tumour load reduction, with a concomitant decrease of serum gastrin levels. Our data indicate an important anti-proliferative effect of SSA on ECL cells, providing clinical benefit and obviating, at least temporarily, the need for invasive therapies for GCA1.

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Introduction

Gastric carcinoids (GCs) are endocrine tumours of the gastric mucosa that originate from enterochromaffin-like (ECL) cells (1). These tumours are divided into three distinct types: the majority (75%) is type 1 GCA1, associated with chronic atrophic gastritis; type 2 (GCA2: 5–10%) is associated with the Zollinger–Ellison syndrome and occur almost exclusively in the context of multiple endocrine neoplasia type 1. Both type 1 and type 2 GCs are well-differentiated tumours related to hypergastrinaemia, and have an overall excellent prognosis. By contrast, type 3 GCs (15–25%) are not related to hypergastrinaemia and follow an aggressive course (1–4).

Almost all type 1 GCs (ECLomas) are non-functioning tumours, typically discovered during upper gastrointestinal (GI) track endoscopy performed for either non-specific symptoms (nausea, vomiting and abdominal pain or dyspepsia) (5), and/or during investigation of anaemia (6). Very occasionally, (less than 1%), patients may complain of flushing and present with the 'atypical carcinoid syndrome' (7, 8). Type 1 GCs occur more frequently in women in the fifth and seventh decades (9, 10). However, with more extensive use of endoscopy, particularly in patients with other autoimmune diseases, the age at diagnosis may be younger (11, 12). In addition, in cases of pernicious anaemia (PA)-induced atrophic gastritis, areas of intestinal metaplasia within the gastric mucosa can develop increasing further the risk for malignancy (13).

Traditionally, GCA1s are endoscopically or surgically removed, depending on the number, appearance and size of the tumours (14, 15). Antrectomy, with surgical excision of the majority of the G cells, is thought to facilitate regression of these tumours by removing the source of excessive gastrin secretion; however, the long-term benefits of antrectomy still remain uncertain
Although proton pump inhibitors (PPIs) are effective in reducing hypergastrinaemia-induced gastric acid hypersecretion in GCA2 (17), they do not affect ECL-cell hyperplasia, therefore their role in GCA1 is limited (11). Moreover, in selected cases, significant reduction of hypergastrinaemia did not prevent development of ECL carcinoid (18), suggesting that, in addition to hypergastrinaemia, other pathogenic or genetic factors may be involved.

Over the last few years, somatostatin analogues (SSAs) have been used in the treatment of patients with either GCA1 or GCA2 (11), based on their capability to inhibit gastrin release from the G cells and thus reduce the ECL-cell hyperplasia (19–22). Morphometric studies demonstrated that, while antrectomy specifically decreased the volume of ECL cells versus the total volume of endocrine cells, octreotide reduced the overall endocrine cell volume (19, 23). Although the number of treated patients is small, it has been suggested that SSA may exert important anti-proliferative effects either directly, by inhibiting ECL-cells proliferation, or indirectly through suppression of gastrin hypersecretion (24).

We conducted a multicentre prospective study in order to assess the effects of long-acting SSA on excessive gastrin secretion, and subsequent ECL-cell growth in patients with GCA1. We also studied the ability of these drugs to induce regression of macroscopically visible tumours and concomitant intestinal metaplasia in treated patients.

Materials and methods

Fifteen patients with GCs type 1 diagnosed and treated in three different centres for at least a 6-month period were studied. Information on clinical presentation, biochemical profile, radiological assessment, histopathological findings and extent of the disease (using the recently introduced tumour, lymph nodes, metastases (TNM) classification) (25) were recorded. In addition, the application of other therapeutic modalities including surgery, and long-term outcome of these tumours were also recorded. Patients were assessed before the initiation of treatment, and thereafter at 6-month intervals both clinically, biochemically, endoscopically and histopathologically according to a uniform protocol (see below). Fourteen patients were treated with octreotide LAR (Novartis) at a monthly dosage of 20 mg or 30 mg, depending on the symptomatic, biochemical and endoscopic response, while a patient received Somatuline Autogel (Ipsen, Paris, France) at a monthly dosage of 90 mg, due to intolerance to octreotide LAR.

Clinical assessment

In order to define symptomatic response, patients were questioned for the presence of specific symptoms such as abdominal pain, nausea, vomiting and dyspepsia (5).

In the absence of any symptoms, the reason for endoscopy was recorded particularly whether patients were investigated for anaemia either macrocytic or due to iron deficiency (5, 9). The presence of other autoimmune disorders in association with PA and the presence of carcinoid tumours or GI malignancies in other family members were also recorded.

Biochemical evaluation

All patients underwent a complete biochemical assessment. Pernicious anaemia was defined as low serum vitamin B12 levels (normal range 180–670 pmol/l) and at least one positive antibody titre against parietal cells, intrinsic factor and proton pump antigen. Serum gastrin and chromogranin A (CgA) were measured the morning before treatment with SSA after an overnight fast, and thereafter at regular intervals (3–6 months) during the study period. One patient was treated with PPIs preceding the beginning of the study due to suspected Barrett’s oesophagus; the same patient was also on aspirin therapy due to a transient ischaemic event (Table 2, patient number 6). When previously used, PPIs were discontinued at least 6 weeks before blood samples were taken. Serum gastrin and CgA were measured using commercially available radioimmunoassay kits: DiaSorin, Stillwater, MN 55082–0285, USA, using a normal reference range of 40–108 mU/l for gastrin, and CGA-RIACT, Cisbio International, Bagnols-Ceze, France, using a normal reference range of 19.4–98.1 ng/ml for CgA.

Radiological assessment

All patients underwent 111In-pentetreotide scintigraphy (Octreoscan) before the initiation of therapy. Ten patients also underwent computerized tomography (CT) of the abdomen to exclude the presence of distant metastases.

Endoscopic and histopathological assessment

All patients underwent upper GI endoscopy and endoscopic ultrasonography (EUS). Upper GI endoscopy with multiple biopsies in the antrum, body and fundus was performed and the ‘dominant’ lesions were removed if possible. EUS was performed to evaluate the extension of the lesions and exclude invasion of the muscularis wall, regional lymph node and/or visible metastases. Biopsies from both the tumours, if not completely resected, and surrounding mucosa in the antrum, body and fundus of the stomach were performed every 6 months during treatment. Sections were immunostained for chromogranin, neuron specific enolase (NSE), synaptophysin (SYP) and the Ki-67 proliferative index using the mindbomb homolog (Drosophila) (MB-I) antibody. The diagnosis of carcinoid tumours was confirmed morphologically during endoscopy together with a positive immunocytochemical staining for NSE, SYP and/or chromogranin.
Evaluation of response to treatment

In all cases, the disease response was defined using established World Health Organization (WHO) criteria, as follows: 1) complete response (complete regression of all clinical, radiological and hormonal evidence of the tumour); 2) partial response (a 50% or greater reduction in all measurable tumour, clinical symptoms and hormonal levels, with no appearance of new lesions); 3) stable disease (less than 50% reduction or no greater than 25% increase in tumour size, clinical symptoms and hormonal measurements) and 4) tumour progression (appearance of new lesions, or an increase of 25% or more in tumour size, and clinical/hormonal deterioration) (12).

Patients were considered in remission if symptoms disappeared, gastrin and CgA levels were substantially reduced or returned to normal range and if there was no evidence of residual disease following treatment with SSAs. When a lesion was removed endoscopically prior to the initiation of therapy with SSAs, response to treatment was evaluated in the remaining lesions. The study was approved by the local institutional ethical committees and informed consent was obtained from all patients.

Statistical analysis

Results were expressed as mean±s.d. Differences in variables were analyzed by the χ² test and Student’s t-test using the SigmaStat 2.03 computerized program (Systat Software Inc., Point Richmond, CA, USA). P < 0.05 was considered significant.

Results

The clinical characteristics of all patients included in the study are shown in Table 1. The cohort included 2 men and 13 women with a mean age of 53.9 years. The mean duration of follow-up was 18 months (range 6–48 months). Other autoimmune diseases (such as Hashimoto’s thyroiditis, Graves’ disease, Sjögren’s syndrome or Addison’s disease) were diagnosed in ten patients (67%). To’s thyroiditis, Graves’ disease, Sjo¨gren’s syndrome or Addison’s disease) were diagnosed in ten patients (67%).

In two patients (13%) there was a positive first-degree relative family history of GC tumours. In one patient (Table 2, patient 3), there were no other adverse effects associated with SSA treatment.

Follow-up assessment (after treatment with SSA)

All patients tolerated treatment well and none discontinued treatment during a follow-up period of 17.8±7.47 months (mean±s.d.). Apart from a slight perturbation in the control of pre-treatment diabetes mellitus in one patient (Table 2, patient 3), there were no other adverse effects associated with SSA treatment.

Thirteen patients (87%) had symptoms attributed to atrophic gastritis (such as abdominal pain, nausea, vomiting or dyspepsia) that improved in all following treatment with SSA (using the described WHO criteria) (12). Serum gastrin was measured in all patients and decreased progressively in all except one patient from 898±418 mI/l pre-treatment, reaching 304±278 mI/l at last visit (normal range 40–108 mI/l, P < 0.005; Fig. 3A). In one patient, despite an initial positive response to treatment (in terms of a decrease in serum gastrin levels and tumour regression at endoscopy), an escape phenomenon was observed, as higher gastrin levels were found at the last examination besides...
Table 2. Clinical and histopathological characteristics of the study patients before and during somatostatin analogues (SSAs) treatment.

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In patient 15, after an initial good response to treatment, the tumour progressed, in association with elevation of gastrin and chromogranin A levels in the serum. PA, pernicious anaemia; Nr. of lesions, single lesion seen on endoscopy, multiples ≥ 2 lesions seen on endoscopy; ECLH, ECL-cell hyperplasia; SomA, Somatuline Autogel; SanLAR, Sandostatin LAR.
treatment with SS analogues (Table 2, patient 15). The levels of serum CgA were consistently evaluated in five patients (34%) and decreased significantly in all except one, from 370 ± 183 ng/ml to 148 ± 69.3 ng/ml (mean ± s.d.; normal range 19.4–98.1 ng/ml, P < 0.005; Fig. 3B).

In all patients, gastroscopic examination showed a reduction in the size and number of carcinoid tumours at 6 months of treatment. In 11 patients (73%), a complete disappearance of the tumours at 1 year of treatment was observed (Figs 1 and 2). In three patients (20%; Table 2, patients 8, 9 and 13), the tumours decreased significantly in number (from ‘multiple’ to ‘single’) and in size (from 7.33 ± 3.06 mm to 1.33 ± 0.57 mm, mean ± s.d.), without complete disappearance; in this small subgroup of patients, an important decrease in the number of cells positively staining for neuroendocrine markers was observed, in parallel with inhibition of Ki-67 proliferation index, which decreased from 2.33 ± 1.15 to 1.67 ± 0.57 (mean ± s.d.). In one patient (Table 2, patient 15), following initial response, the original tumour progressed, in association with elevation of serum gastrin and CgA levels, and he was referred for antrectomy and surgical excision of the tumour. Intestinal metaplasia regressed and disappeared in the two patients observed before treatment.

All patients are still treated with SSA in an effort to determine the optimal duration of treatment in terms of gastrin and CgA suppression and tumour regression (Fig. 3).

**Discussion**

Despite the relatively ‘benign’ and indolent biological behaviour of GCA1 tumours, some may produce non-specific symptoms, whereas ~8–23% may metastasize to regional lymph nodes and rarely to the liver albeit with a low overall mortality (5). Although biotherapy with SSA is not currently recommended according to the European Neuroendocrine Tumours Society (ENETS) guidelines, it has been occasionally used in reference centres (26). The combination of octreotide and α-interferon has been found to be of value in a patient with metastatic disease to the liver (5), whereas several, mainly small-sized studies, have documented a beneficial effect of treatment with SSA (26–29). The results of our study confirm these findings and show that treatment with long-acting SSA given at monthly intervals for a period of at least 6 months produces a significant suppression in gastrin and CgA levels, albeit without achieving complete normalization. Despite this apparently incomplete antisecretory effect, a significant anti-tumour effect is obtained, as shown from the regression of ECL-cell hyperplasia and tumour disappearance in the great majority of patients. This finding suggests a direct effect of these drugs on tumour cell proliferation. However, as the reported 5-year survival
of GCA1 patients is good particularly when followed regularly with endoscopy (30), and rarely spontaneous tumour regression may occur (31), the results of our study, and other similar studies should be considered on an individual patient basis.

In patients with GCA1 (type 1 ECLomas), annual surveillance is thought to be sufficient for patients harbouring tumours less than 10 mm (32). In the case of larger tumours, endoscopic resection is recommended for up to six polyps not involving the muscularis, whereas in the remaining patients, local surgical tumour resection should be performed (30). However, the validity of such an approach is based on the findings of a single study (30). In addition, no strict guidelines are available for patients with tumours greater than 1 cm in size, when the risk of malignancy may be increased, and for those whose tumours recur despite repeated excisions (32). Antral resection, which is thought to diminish repeated and chronic gastrin stimulation of ECL cells, is effective in 80% of type 1 tumours (33–35). Antrectomy and local resection is also performed in the presence of deep gastric parietal wall invasion and positive margins following endoscopic removal (36). Partial or total gastrectomy with lymph node dissection is employed in the presence of malignancy or recurrence besides local surgical resection (36).

The rationale of using SSA in patients with GCA1 resides in the notion that the lack of gastrin’s stimulatory effect on ECL cells may lead to regression of hyperplastic and/or dysplastic lesions (1). This is also the case when patients with GCA1 undergo an antrectomy thus removing the source of excessive gastrin secretion. However, such an approach is not always successful, as cases of recurrence of GCA1 after antrectomy have described (37, 38). This could be attributed to established and evolving changes in the genome of remaining ECL cells that undergo autonomous proliferation besides lack of gastrin stimulation. The use of SSA, apart from a reduction in gastrin hypersecretion, may also exert an anti-proliferative effect in the hyperplastic or dysplastic ECL cells. This approach is not associated with the complications of a relatively extensive surgery, such as antrectomy, and offers the opportunity of a medically induced reduction of the size and/or the number of the ECLomas. In addition, most ECLomas are preceded (or accompanied) by linear or micronodular hyperplasia or dysplasia of ECL cells (39). This condition is associated with a 26-fold increase in the risk of developing ECLomas in patients with chronic gastritis (40). As shown from the results of our study, SSA can also affect these changes therefore substantially reducing the risk of further lesion development. Intestinal metaplasia, which may also arise in the context of long-lasting chronic atrophic gastritis and may precede the development of gastric adenocarcinoma (41–43), was totally suppressed by treatment with SSA in the present study.

Information regarding the use of SSA in the treatment of GCA1 is rather scanty. Following the disappearance of GCA2 after s.c. octreotide administration (29), the long-acting SSA octreotide-LAR induced a permanent regression and disappearance of multiple GCA1s with normalization of elevated serum gastrin in a patient treated for 9 months (28). Treatment with monthly injections of octreotide-LAR for 1 year was also assessed in five patients with hypergastrinaemia-induced GCs (27). At the end of the study, although gastrin levels did not totally normalize, a substantial reduction in the density of ECL cells and tumour load, accompanied with a concomitant reduction in CgA levels was noticed. These findings indicate a possible direct anti-proliferative effect of the SSA.
treatment. Recently, nine patients with more than five type I gastric endocrine tumours each less than 1 cm in size, without invasion of the muscularis propria and a Ki-67 index lower than 3%, were treated with long-acting SSA; in all patients, the tumours disappeared after 12 months of therapy, in parallel with significant reduction of CgA and gastrin levels (26). However, it is still unclear why some patients fail to achieve normalization of serum gastrin and why a minority besides an initial biochemical and morphological response develop a rebound increase in gastrin levels, using the same dosage of the given drugs. This heterogeneous response of GCs to the administration of SSA suggests that the effects of these drugs probably include a combination of both a direct effect on the tumour and an indirect effect, on the gastrin-induced cellular hyperplasia.

Despite the complete disappearance of the tumours in the majority of patients included in the present study, in a small subgroup of three patients the tumours did not totally regress. This different tumour behaviour was not found to be related to higher levels of gastrin or CgA. Still, in this small subgroup of limited responsive patients, in terms of partial reduction in the number and size of the tumours, and in their index of proliferation, treatment with SSA may induce an important tumour down staging (i.e. large tumours can become smaller and therefore removed endoscopically rather than proceeding to surgery).

The administration of SSA as a chronic therapeutic option in GCA1 patients could be limited by undesirable side-effects (44, 45), or by the high cost of these drugs; however, side-effects are mostly temporary, usually observed in a minority of patients, as demonstrated in our study. In order to address these potential limitations of treatment with SSA, a prospective study, including a control group (i.e. patients without treatment with SSA), is required. Such a study could be important as it could justify the use of SSA in particular groups of patients, such as older patients who may have other comorbidities and in whom surgical therapy, when indicated, may not be easily applicable and without significant risks, or patients that cannot be followed up regularly.

Based on the results of our study, SSA could be used as first-line treatment for hypergastrinaemia-associated GCs, replacing the conventional endoscopic or surgical excisions, particularly in patients with multiple or relapsing tumours. In these patients, treatment with SSA could be continued as long as gastrin/CgA levels are suppressed, in parallel with decrease/disappearance of the tumours observed on regular endoscopy. Moreover, surgical procedures should most probably be performed only in patients unresponsive to medical treatment.

Declaration of interest
The authors declare that there is no conflict of interest that would prejudice the impartiality of this scientific work.

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