PRRT as an alternative method of treatment in patient with glucagonoma syndrome: A case report


1. Department of Endocrinology, Diabetology and Internal Medicine, Medical University of Białystok, Poland
2. Department of Endocrinology and Nuclear Medicine, Military Institute of Medicine, Warsaw, Poland
3. Department of Radiology, Medical University of Białystok, Poland
4. Students’ Research Group in the Department of Endocrinology, Diabetology and Internal Medicine, Medical University of Białystok, Poland

A - Conception and study design, B - Data collection, C - Data analysis, D - Writing the paper, E - Review article, F - Approval of the final version of the article

ABSTRACT

Introduction: Glucagonoma is a rare pancreatic neuroendocrine tumor derived from alpha-cells of the islet of Langerhans. Due to oversecretion of glucagon it is associated with a characteristic paraneoplastic phenomenon, called glucagonoma syndrome, which consists of necrolytic migratory erythema (NME), weight loss, diabetes mellitus, diarrhea, normochromic normocytic anemia, deep vein thrombosis or pulmonary embolism and neuropsychiatric disturbances. Treatment modalities include surgical removal of tumor, somatostatin analogs and peptide receptor radionuclide therapy (PRRT).

Case report: We present a case of 61-year-old woman diagnosed with glucagonoma in April 2012. Initially, body-caudal pancreatectomy and resection of regional lymph nodes were performed. Five months after surgery, a PET-CT scan detected pathological mass with expression of somatostatin receptors in pancreatic body and metastases to regional lymph nodes. What is more, since April 2014 the patient had complained about persistent pruritus of the entire body. At present, due to the nonsurgical pancreatic mass and metastases she is treated with somatostatin analogs and PRRT. During this therapy the pruritus had decreased and currently there is no sign of cutaneous disease. Moreover, reduction of tumor size was obtained.

Conclusions: PRRT may reduce tumor size and by reducing bothersome symptoms substantially improve the quality of life in patients with SSTR-positive tumors.

Key words: Glucagonoma, tumor, peptide receptor radionuclide therapy

DOI: 10.5604/01.3001.0010.1922

*Corresponding author:
Anna Popławska-Kita
Department of Endocrinology, Diabetology and Internal Medicine
Medical University of Białystok, Skłodowskiej-Curie 24A
15-276 Białystok, Poland
Tel.: +48 604425324; Fax: +48 857461909

Received: 09.06.2016
Accepted: 19.06.2016
Progress in Health Sciences
Vol. 6(1) 2016 pp 209-214
© Medical University of Białystok, Poland
INTRODUCTION

Glucagonoma is a rare, slowly growing, frequently malignant pancreatic neuroendocrine tumor (NET) arising from alpha-cells of the islets of the Langerhans. In 1942, Becker, Kahn and Rothman first noticed an association between characteristic skin disorder and pancreatic tumor [1]. Then in 1966 McGavran et al. [1] discovered the origin of this disorder thanks to radioimmunoassay technique for glucagon and named this combination of symptoms as a glucagonoma syndrome [2]. Since its first description [1], around 300 cases of glucagonoma and glucagonoma syndrome have been described. The incidence of glucagonoma is estimated at 1 in 20 million [3]. The tumor is usually bigger than 5 cm [4] and the most common location of this lesion is the tail of the pancreas [3,5,6].

Glucagonoma, due to oversecretion of glucagon is associated with a characteristic paraneoplastic phenomenon, called glucagonoma syndrome, which consists of specific rash known as necrolytic migratory erythema (NME), mild diabetes mellitus (DM), weight loss, normochromic normocytic anemia, glossitis, stomatitis, hypoaminoacidemia, deep vein thrombosis and depression [7].

Diagnosis of this disorder is based on clinical features, laboratory tests, such as elevated glucagon serum level and visualization of the pancreatic tumor. Treatment modalities include surgical removal of tumor, somatostatin analogs and peptide receptor radionuclide therapy (PRRT) [5,7].

Case report

In March 2012, a 61-year-old female was admitted to Department of Gastroenterology and Internal Medicine of Medical University of Białystok due to intense abdominal pain, which was radiating to vertebral column. She also complained about vomiting, flatulence and bad general condition. The anamnesis revealed 4-year history of similar (but slighter) episodes of pain and the weight loss of about 20 kg over the past one year. Additionally, the patient had suffered from DM type 2 since 2010 and arterial hypertension for about 10 years. Glycemia level was well-controlled by metformin 500 mg twice a day. Upon physical examination there was pain in the upper and the middle abdomen, there was no visceromegaly or palpable masses. The patient had high blood pressure (180/95 mmHg). Admission laboratory tests demonstrated mild leukocytosis, elevated C-reactive protein concentration and pyuria. In the investigation, abdominal ultrasound, CT and upper gastrointestinal endoscopic ultrasonography were performed, which disclosed a hypoechogenic mass with several small calcifications and irregular margins on the pancreas topography of approximately 55 x 40 mm in size. The tumor was extending into splenic artery and vein. The tail of the pancreas was invisible, probably atrophied, the pancreatic head was normal. There were enlarged celiac lymph nodes of 7-8 mm and one hypoechogenic lymph node of 11.5 mm above pancreatic head (suspicion of metastasis) (Fig.1).

Figure 1. Hypoechogenic mass with several small calcifications and irregular margins on the pancreas topography of approximately 55 x 40 mm in size. The tumor was extending into splenic artery and vein. The tail of the pancreas was invisible, probably atrophied, the pancreatic head was normal. There were enlarged celiac lymph nodes of 7-8 mm and one hypoechogenic lymph node of 11.5 mm above pancreatic head (suspicion of metastasis).

In November 2012, CT scan showed oval, marginal enhancing pancreatic mass of 26 x 21 x 23 mm, which could be cancer remains or recurrence. There were also slightly enlarged, non-homogenous enhancing paraaortic lymph node and hypodense nodules nearby the lesser curvature of the stomach.

Figure 2. In November 2012, CT scan showed oval, marginal enhancing pancreatic mass of 26 x 21 x 23 mm, which could be cancer remains or recurrence. There were also slightly enlarged, non-homogenous enhancing paraaortic lymph node and hypodense nodules nearby the lesser curvature of the stomach.
Transesophageal biopsy was performed and pathological examination showed inflammatory cells and cells suspicious for malignancy.

Transesophageal biopsy was performed and pathological examination showed inflammatory cells and cells suspicious for malignancy. Chromogranin A concentration was 379.4 ng/ml (normal range 27-94 ng/ml).

In April 2012, body-caudal pancreatomy and resection of regional lymph nodes were performed. The tumor was spreading to the celiac trunk and to the inferior mesenteric vein. The histopathologic report revealed a neuroendocrine carcinoma with expression of glucagon, chromogranin A, synaptophysin and Ki 67 - 2%. 5 months after surgery a PET-CT scan detected pathological mass (23 x 14 mm in size) with expression of somatostatin receptors (SSTR) in pancreas and metastases to regional lymph nodes. There was also a lesion of 26.6 x 22.5 mm without expression of SSTR. Estimated chromogranin A concentration was in the normal range (36.3 ng/ml).

In November 2012, a contrast-enhanced CT scan showed oval, marginal enhancing pancreatic mass of 26 x 21 x 23 mm, which could be cancer remains or recurrence. There were also slightly enlarged, non-homogenous enhancing paraaortic lymph node (11 mm) and hypodense nodules (6-11 mm) nearby the lesser curvature of the stomach. (Fig.2). At that point, chromogranin A concentration increased up to 157.0 ng/ml; 123.2 ng/ml.

Due to the nonsurgical, SSTR-positive tumor somatostatin analog has been introduced to patient (Sandostatin LAR 30 mg) since December 2013 (on average once a month). During the treatment chromogranin A concentration stayed in the normal range (36.3 ng/ml).

In March 2014, the abdominal CT scan identified 20 x 18 mm oval mass with marginal calcification without evidence of enhancement. Nearby pancreatic head there was a hyperdense mass of 5 mm. In lesser curvature of the stomach area there was a tumor of 9 x 7 mm in size (Fig.3).

The patient had complained about persistant pruritus of the entire body since April 2014. Physical examination revealed nonspecific, migratory erythematous skin lesions, which involved face, chest, abdomen and limbs. She was treated with various unguents and creams without relief of the symptoms. Somatostatin injections slightly relieved symptoms for a short time. Chromogranin A concentration was still in the normal range (37.8 ng/ml).

In September 2014, a 63-year-old patient was admitted to Department of Endocrinology, Diabetology and Internal Medicine of Medical University of Bialystok to prepare for peptide receptor radionuclide therapy (PRRT). The patient’s insulin demand dropped (down to 8 units per day), so insulin therapy was interrupted and glimepiride 2 mg once a day was administered. Based on thyroid ultrasound and laboratory findings, nontoxic multinodular goiter was diagnosed. Ultrasound guided fine-needle aspiration biopsy results indicated benign lesions. Due to pruritus an antihistamine, a tranquilizer and an alpha lipoic acid were prescribed. Although the patient’s adherence was good, there had been no significant improvement.

Figure 3. In March 2014, CT scan identified 20 x 18 mm oval mass with marginal calcification without evidence of enhancement. Nearby pancreatic head there was a hyperdense mass of 5 mm. In lesser curvature of the stomach area there was a tumor of 9 x 7 mm in size.

In March 2014, the abdominal CT scan identified 20 x 18 mm oval mass with marginal calcification without evidence of enhancement. Nearby pancreatic head there was a hyperdense mass of 5 mm. In lesser curvature of the stomach area there was a tumor of 9 x 7 mm in size (Fig.3).

The patient had complained about persistant pruritus of the entire body since April 2014. Physical examination revealed nonspecific, migratory erythematous skin lesions, which involved face, chest, abdomen and limbs. She was treated with various unguents and creams without relief of the symptoms. Somatostatin injections slightly relieved symptoms for a short time. Chromogranin A concentration was still in the normal range (37.8 ng/ml).

In September 2014, a 63-year-old patient was admitted to Department of Endocrinology, Diabetology and Internal Medicine of Medical University of Bialystok to prepare for peptide receptor radionuclide therapy (PRRT). The patient’s insulin demand dropped (down to 8 units per day), so insulin therapy was interrupted and glimepiride 2 mg once a day was administered. Based on thyroid ultrasound and laboratory findings, nontoxic multinodular goiter was diagnosed. Ultrasound guided fine-needle aspiration biopsy results indicated benign lesions. Due to pruritus an antihistamine, a tranquilizer and an alpha lipoic acid were prescribed. Although the patient’s adherence was good, there had been no significant improvement.

Figure 4. January 2015. 18FDG PET-CT - mild 18F-FDG uptake by the mass in the pancreatic body’s area.

In October 2014, the 18FDG PET-CT showed mild 18F-FDG uptake by the mass in the pancreatic body’s area (19 x 13 mm - smaller than in 2012). In November 2014, the 99mTc-Tektreotide whole body scintigraphy revealed two lesions (27 mm and 17 mm) with expression of somatostatin receptors on the pancreatic body.
topography. In December 2014, the CT scan showed two masses: the diameter of the first was 26 mm (previously - 24 mm), of the second - 17 mm (previously - 19 mm). The pancreatic head was polycyclic and enhanced in non-homogenous way.

In January 2015, the patient was admitted to Department of Endocrinology and Isotope Therapy of Military Institute of Medicine in Warsaw for peptide receptor radionuclide therapy. The patient was treated, under the cover of nephroprotection, with intravenous 200mCi 177Lu-DOTA-TATE (3,7 GBq) (Fig.4). This treatment was repeated in April and July 2015. After PRRT the chromogranin A concentration stayed in the normal range (69.2 ng/ml).

In July 2015, the patient was admitted to Department of Endocrinology, Diabetology and Internal Medicine due to bad glycemic control (glucose concentration from 36 to 280 mg%). Laboratory measurements demonstrated elevated LDL-cholesterol level. HbA1c was in recommended range (7.0 %). Abdominal ultrasound detected steatohepatitis. Hypoglycemic therapy was modified. Apart from glimepiride 2 mg once a day, metformin 850 mg three times a day was added.

Figure 5 November 2015. SPECT-CT of the abdomen revealed Lu-177 uptake by two points (diameters: 24 mm and 23 mm) in pancreas area.

In November 2015, the Lu-177 whole body scintigraphy and SPECT-CT of the abdomen revealed Lu-177 uptake by two points (diameters: 24 mm and 23 mm) in pancreas area (Fig.5). Peptide receptor radionuclide therapy has been modified and the patient received 90Y/177Lu-DOTA-TATE (1.85 GBq + 1.85 GBq). During PRRT the pruritus had become progressively lesser. Currently there was no sign of cutaneous disease.

In February 2016, the abdominal PET-CT scan showed a pancreatic body mass of 18.9 mm (Fig.6). Minimal reduction of tumor size and stabilization of proliferative disease were obtained.

Figure 6. In February 2016, the abdominal PET-CT scan - a pancreatic body mass of 18.9 mm.

DISCUSSION

Glucagonoma is a remarkably rare neuroendocrine tumor with estimated incidence 1 in 20 million [3] and is 1% of all pancreatic NETs [7]. Glucagonomas, due to oversecretion of glucagon are associated with a characteristic paraneoplastic phenomenon, called glucagonoma syndrome [8]. It’s characteristic features are: necrolytic migratory erythema (NME), weight loss, DM or glucose intolerance, mucosal abnormalities (such as glossitis, cheilitis, stomatitis), diarrhea [4]. Patients may also present normochromic normocytic anemia, hypoaminoacidemia, deep vein thrombosis or pulmonary embolism and neuropsychiatric disturbances (from which depression is most often) [5]. In the literature this syndrome may be also presented as 4D syndrome, which stands for dermatosis, diabetes, deep vein thrombosis and depression [7]. Our patient presented NME, weight loss and DM.

Although earlier studies suggest female predominance (3-4 : 1 female : male), a recent review suggests no gender predilection [9], with the median time of diagnosis in their sixth decade of life [5,3]. The diagnosis of glucagonoma should be based on elevated serum glucagon level and visualization of the pancreatic tumor [9]. Glucagonoma most often is found in the tail of the pancreas [5,3,6]. In our patient’s case there was a mass lesion in the pancreatic body. The tail of the pancreas was invisible in USG and CT, probably atrophied.

Glucagonoma is slow-growing and rather low-malignancy tumor [8], Ki-67 is routinely used as a static marker of proliferative activity. The fraction of Ki-67-positive tumor cells (the Ki-67 labeling index) is often correlated with the clinical
The best treatment strategy of glucagonoma is surgical removal of the tumor [11]. Surgical approach depends on location, size and pathological type of the tumor and it can be divided into local resection, pancreatoduodenectomy or pancreatic body and tail resection [11]. Unfortunately surgery is often impossible, due to expansive involvement of hepatic parenchyma or bilobar hepatic metastases [7]. In our patient case, due to infiltration into splenic artery and vein, the surgery was subtotal. In such cases non-surgical therapies play a great role in palliative treatment and relieving symptoms.

The base of that treatment are somatostatin analogs [12]. Somatostatin inhibits glucagon secretion, but due to 2-minute half-life it can’t be used in daily clinical practice. Therefore longer-acting somatostatin-analogs, such as octreotide and lanreotide have been developed [12]. They can be used if somatostatin receptors are present in histological examination or the Octreoscan is positive [7]. They allow to control NME in 50-90% cases and they also relieve other symptoms, like weight loss, abdominal pain and diarrhea [4]. In this case somatostatin injections slightly relieved skin symptoms for a short time. Moreover, the patient stopped complaining about abdominal pain and weight loss. Unfortunately, somatostatin analogs do not improve the course of diabetes [4]. Due to deterioration of DM, our patient required hospitalization and changing the treatment from insulin therapy and glimepiride to metformin treatment. Somatostatin may also have apoptotic and cytostatic effects [12]. It has been shown that octreotide can inhibit tumor growth in patients with midgut NETs [7,12]. When these analogs are used for more than 12 months, tachyphylaxis may occur [12]. Our patient has been treated on average once a month with 30mg of somatostatin since December 2013 and there weren’t any changes of the dose since the beginning of this therapy.

A relatively new therapeutic option is peptide receptor radionuclide therapy. Malignant cells of the tumor express a high number of somatostatin receptors [13]. By using appropriate vectors and linkers with hard beta-emitting yttrium or lutetium, it enables identification of the tumor and, what is more important, destruction of cells with somatostatin receptors [7]. Previous studies has shown that PRRT may substantially improve the quality of life in patients with SSTR-positive tumors [13]. The case of our patient seems to confirm that. So far she has received three doses of 200mCi 177Lu-DOTA-TATE and one dose of 90Y/177Lu-DOTA-TATE. Pruritus and skin lesions have decreased during this therapy and currently there is no sign of cutaneous disease. The patient’s quality of life improved.
CONCLUSIONS

Peptide receptor radionuclide therapy, by reducing bothersome symptoms, may substantially improve the quality of life in patients with SSTR-positive tumors. Moreover, due to selective concentration in tumor cells, it allows to obtain stabilization of proliferative disease.

Conflicts of Interest

The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the review reported. All authors declare any financial or other potential conflict of interest.

Funding

This research did not receive any specific grant from any funding agency in public, commercial or not-for-profit sector.

REFERENCES