INTRODUCTION

Paraneoplastic Cushing syndrome represents 5-10% of all Cushing syndrome and has a severe prognosis due to severe metabolic imbalance, denutrition, associated infections and progression of tumoral underlying pathology. The death is a rule in more than 50% of cases. Some medication used to treat it – Metirapone – is not available in Romania. Ketoconazole was recently approved by CHMP for treatment of paraneoplastic Cushing’s only in November 2014.

CLINICAL CASE

A 67 years old woman presented on the 21st of November with mental confusion, progressive weight loss, severe edema and kypokalemia, without typical features of Cushing or hyperpigmentation. Patient’s behavior changed in the last 5 months, she was nasty with her daughter, bickering, while diabetes and hypertension aggravated in the last 3 months.

The electrolytic imbalance was severe at admission K 1.65 mmol/l, in spite of multiple attempts to correct it with 150 mmol/day KCl on peripheral i.v. line, 40 mmol/day of KCl orally and 200 mg/day of Spironolactone, treatment used initially in “C. I. Parhon” National Institute of Endocrinology. Patient was transferred in the I. C. U. of “Dr. Carol Davila” Central Military Emergency Hospital for the weekend, in order to obtain a better control using a central intravenous catheter.

Laboratory work and imaging

One month prior to admittance patient had hypercortisolism, with normal hepatic citolytic enzymes and normal TSH, free T4 and calcitonin values. Investigations at admittance revealed paraneoplastic Cushing’s with ACTH 82.5 pg/ml, cortisol levels more than 63 mcg/dl, UFC (urinary free cortisol) 2,866 mcg/24 h (21-111); DHEA-sulfate 230.5 mcg/24 h; 2 mg Dexametasone suppression test showed unsuppressed cortisol 59.17 mcg/dl, ACTH 123.8 pg/ml.

Patient also associated empty sella syndrome with thyrotrophic and gonadotrophic insufficiency, normal prolactin and IGF-1, normal mineralocorticoid hormones, cathecolamines, serotonin and 5HIAA, slightly enlarged cromogranine A – 148 ng/ml (upper limit 125 ng/ml).

Patient had also left breast tumor, Helicobacter pylori gastritis, polinodular goiter, denutrition and hepatic dysfunction.

Imaging techniques: 99m Tc Tektrotyd scintigraphy
showed uptake at 10 minute in left breast and jejunal loop; upper and lower endoscopy, echo endoscopy – revealed no tumors; bronchoscopy – no visible tumor (after procedure, patient suffered a syncope that lasted 2 minutes); thyroid ultrasound – found nodule of 15/18/22 mm located in the lower part of left lobe – 15/18/22 mm, with low peripheral vascularisation, uptakes iodine at CT scan and has a peripheral calcification.

**Figure 1.** Abdominal CT showed minimally enlarged right adrenal gland, with nodule 0.88/0.88 cm; enlarged left adrenal gland, with nodule of 1.03/1.24 cm.

**Figure 2.** Thoracic CT revealed pulmonary tumor located in Fowler segment of left superior lung lobe.

**Treatment**

We initiated treatment with Ketoconazole 400 mg, 1 day, and then 600 mg, for 2 days, but with inadequate correction of alkalosis and hypokalemia – pH was 7.54-7.59, BE 5.7-9.8 mmol/l, K 3.16 mmol/l. The third day patient became septic (probably MRSA Staphylococcus) due to central catheter and interstitial pneumonia – fibrinogen 660 mg/dl, with high liver enzymes – AST 87-160 UI/l, ALT 95-103 UI/l, GGT 348-365 UI/l, total bilirubine 2.44 mg/dl, leucocytes 13,400/mm³, granulocytes 8,500/mm³. Cortisol levels were 26.3-29.2 mcg/dl and Ketoconazole was increased to 1,200 mg/day, also associating Tavanic 500 mg initially, then Tigecycline 100 mg/day. The high values of ALT and AST were due to sepsis and did not increase after doubling Ketoconazole dosage. After 1 day of high dose Ketoconazole, K was 4.7 mmol/l, with introduction of Mifepristone 200 mg/day. The seventh day after Mifepristone was introduced, cortisol levels were 18.7 mcg/dl (4.2-38.4), allowing surgery.

Due to denutrition, pulmonary sepsis, lack of localization of tumor – lung/thyroid/ileum, recent syncope, severe brain atrophy with cognitive impairment, we decided to perform left adrenal gland resection.

The adrenal resection was difficult due to diffuse bleeding and lack of tissue elasticity.

Hepatic biopsy showed periportal fibrosis, but no necrosis of hepatocytes, probably due to toxic substances used at work; left adrenal was 7/3/1.5 cm in diameters, with focal hemorrhage. Immunohistochemistry – Ck7, Ck20, CEA, TF1, ER – negative, MELAN A positive – suggested diffuse hyperplasia of left adrenal gland.

**Evolution of patient**

One hour after left adrenalectomy – cortisol was 18.2 mcg/dl, ACTH 42.3 pg/ml, patient needed inotrop support with Noradrenaline, hydrocortisone 75 mg 1 day, 50 mg in the second day.

The third day cortisol desupressed to 51.25 mcg/dl, ACTH 43 pg/ml (3-66), K decreased again to 2.9 mmol/l, Hb was 8.4 g/dl. Ketoconazole 600 mg/day was started again. Patient had fever, delirium, pulmonary riles, so Meronem was initiated for 2 days, then Tigecycline 3 days, then 7 days of Klacid at home, also Calcium 1 g/day, 1,000 UI D3, 0.5 mcg of 1 alfa-calcidol/day, hepatic protection, vitamins, basal insulin.
Evolution after second surgery: 1 day after carcinoid resection ACTH was 5.95 pg/ml (3-66), cortisol 17.38 mcg/dl – labs were done at „C. I. Parhon” Institute.

12 days after carcinoid resection: ACTH 16.56 pg/ml (7.2-63); Cortisol 10.72 mcg/dl (6.2-19.4).

At 22 days: she lost 8 kg, ACTH 19.1 pg/ml, normal ALT and AST, GGT 236 UI/l, Mg 1.39 mg/dl, Ca 9.96 mg/dl.

At 47 days: she lost 15 kg, ACTH 16.5 pg/ml; cortisol 9 mcg/dl, Mg low even with supplementation.

At 78 days: GGT 259 UI/l (5-36), Mg 1.45 mg/dl, K 3.83 mmol/l, glucose 142 mg/dl, cortisol 20.87 mcg/dl, TSH 2 microU/ml.

At 3 months: basal cortisol 11.2 mcg/dl (6.2-19.4), basal ACTH 17.46 pg/ml (3-88), cortisol 24 EET – 3.24 mcg/dl, cortisol during suppression test with Dexamethasone 1 mg overnight – 0.48 mcg/dl, PTH 12.32 pg/ml (15-165), low 25OH–D 12.8 ng/ml (30-100), UFC 40.3 mcg/24 h (21-111).

We also performed control thoracic CT – revealed left breast tumor of 0.76/1.21 cm, right adrenal with stationary aspect; portal vein was enlarged 14.5 mm; patient performed FNAB of left thyroid nodule on 20 April 2015, showing benign adenoma.

Mild kypokalemia and hypomagnesemia, even with oral supplementation, sartan therapy and normal levels of cortisol and ACTH persisted after surgery, probably due to severe deficit of intracellular compartment, even at 3 months after carcinoid resection. Patient does not remember the 2 months prior to surgery, even if cognitive impairment is mild now.

**COMMENTS**

This case was difficult due to metabolic challenges, multiple associated pathology, lack of SSTR2 and SSTR5 receptors on lung carcinoid with negative scan, mild elevation of cromogranine A levels despite a typical bronchial carcinoid. Patient’s sister was operated for adrenal adenoma confirmed on histology exam, her daughter had papillary thyroid cancer, but no MEN association was proven in this.
family. Patient needed more than 30 days of hospital admittance in two different hospitals and five clinics in order to obtain a good clinical result. The vital risk was high due to sepsis, denutrition, metabolic and ionic imbalance, hepatic lesions, anesthesia, brain atrophy, relative adrenal insufficiency after surgery.

There are no guidelines that state the adequate cortisol levels to be reached before surgery, nor the duration of Ketoconazole wash-out to prevent adrenal insufficiency.

Recently the patient discovered ductal invasive left breast carcinoma, operated at 6 months after thoracic surgery and will soon start chemotherapy.

References:


