INTRODUCTION
Neurilemmoma is a rare tumor of neural crest cell origin. It is usually a benign slow-growing tumor, composed of Schwann cells, which normally produce the insulating myelin sheath covering the peripheral nerves. Schwann cells are very important in the regeneration of damaged peripheral nerves [1]. For reasons not yet understood, neurilemmomas are mostly benign and less than 1% become malignant, degenerating into neurofibrosarcoma. Most commonly they occur as solitary encapsulated subcutaneous tumors. More rarely, they are multiple or arise from points along the PNS, including cranial nerves, spinal roots, the brachial and lumbar-sacral plexus, or major peripheral nerves.

Most neurilemmomas occur in the head, neck, stomach [2] or limbs, with a few cases occurring in the retroperitoneal space. The cause of these neoplasms is unknown [3]. Neurilemmomas can arise from neurofibromatosis. Because these tumors can present in many locations, the clinical presentation can be varied [4]. Some may involve the spinal nerve roots and present with symptoms that mimic those of herniated disk disease of the spine [5]. Deeply situated tumors predominate in the posterior mediastinum and the retroperitoneum. Visceral neurilemmomas, especially of the adrenal gland, are extremely rare and are usually discovered incidentally. Adrenal neurilemmomas are thought to arise from Schwann cells associated with the phrenic nerve, the vagus nerve, and the sympathetic trunk that innervates the adrenal medulla [6].

CASE REPORT
In March 2006, a 41-year-old male patient was referred to the Ukrainian Scientific and Practical Centre for Endocrine Surgery and Organ and Tissue Transplantation as a case of a retroperitoneal tumor (incidentaloma) for further diagnostic evaluation and treatment. Physical examination then revealed a healthy-looking man 177 cm tall and 105 kg in weight, body mass index 33.5, with first stage obesity. No abnormalities were found. Thyroid sonography was normal. Abdominal sonography revealed an hyperechoic round mass (52 mm) in the upper pole of the left kidney.

Laboratory data revealed normal hematology and biochemistry. Endocrine studies showed normal levels of: urine 17-oxytocorticosterone – 3.9 mcmol/day, 17-oxytocorticosterone – 17.6 mcmol/day, 11 oxycorticosteroid – 0.16 mcmol/day. Catecholamine’s: adrenaline – 6.57 nmol/day, noradrenalin – 14.4 nmol/day, dopamine – 176.4 nmol/day, VMA – 4.95 mcmol/day. Computed tomography (CT) revealed a well-marginated heterogeneously enhanced capsulated mass 60x50 mm in the left adrenal region. The patient underwent laparoscopic left adrenalectomy. Pathologic finding of the left adrenal gland showed a picture of a benign neurilemmoma and hyperplasia of the adrenal cortex. The postoperative course was smooth and the patient was discharged after 2 weeks of hospital stay. During the follow-up period till 2014 endocrine studies and abdominal CT revealed no evidence of tumor recurrence. In 2009 thyroid sonography revealed non-toxic multinodular goiter and blood tests revealed hypothyroidism. The patient was prescribed thyroxine 100 mcg tablet daily. Routine monitoring of TSH was recommended. In 2014 the patient underwent a routine check-up at our hospital. Abdominal CT revealed microadenoma (13 mm, density -4 HU to +7 HU non contrast image) of the right adrenal gland and adrenal hyperplasia. Intra venous contrast imaging revealed during the venous phase (60 sec) density of +67 HU to +79 HU, wash out phase (15 min.) -2 HU to +11 HU. Laboratory data revealed normal hematology and biochemistry. Endocrine studies showed normal levels of: Urine metanephrines – 346.9 mcg/day, cortisol (overnight dexamethasone test) – 0.6 mcg/dl, aldosterone – 17.3 ng/l, rennin 4.83 ng/l, TSH – 6.21 µIU/mL, ARR – 3.58, plasma potassium – 4.6 mmol/l.

The patient was recommended to have a routine checkup (abdominal CT and blood tests) after 1 year as the laboratory findings were in the normal range and the patient was normotensive without any complaints. The dose of thyroxine was corrected to 125 mcg/daily.

DISCUSSION
Neurilemmoma is a benign, slow-growing, encaps-
sulated neoplasm in which the principal component arises from neural crest cells and comprises differentiated Schwann cells in a poorly collagenized stroma. Neurilemmomas were first described by Vercay in 1908, with further sub-classification into two distinct histologic patterns performed by Antonini in 1920 [7]. On computed tomography, a neurilemmoma appears as a well-demarcated, round or oval mass that may be homogeneous. However, other cases in the literature have shown prominent cystic degeneration and calcifications [8].

A 2002 National Institutes of Health (NIH), science statement, on the management of adrenal incidentalomas concludes that lesions greater than 6 cm should be excised, those less than 4 cm with imaging characteristics that appear benign generally not be resected, and those between 4 cm and 6 cm can be either closely observed or resected. However, non-secreting tumors with of size >4 cm can be excised with an added benefit of definitive diagnosis [9].

The differential diagnosis of adrenal incidentaloma can be extensive and ranges from cortical lesions such as adenoma and carcinoma to medullary lesions such as neuroblastoma, ganglioneuroma, pheochromocytoma, neurilemmoma and neurofibroma. Rare lesions like leiomyosarcoma and malignant peripheral nerve sheath tumors can also occur.

Histologically, the growth patterns in neurilemmomas include Antoni type A neurilemoma and type B neurilemoma [1]. Immunohistochemistry of adrenal neurilemmoma shows strong and diffuse staining for S-100. They also display pericellular reactivity for collagen IV, laminin, and show absence of reactivity for keratin, desmin, actin, muscle-related antigens, HMB-45, Melan-A, chromogranin, synaptophysin, and CD34 [10].

The management of neurilemmoma requires complete surgical excision. Laparoscopic adrenalectomy is safe and feasible for diagnosis and treatment of benign adrenal neurolemoma [11].

In conclusion, total excision of benign neurolemoma is associated with favorable outcome in patients, but as our case depicts regular follow up is needed, as our patient presented with a hormonally inactive micro adenoma and adrenal hyperplasia of the right adrenal gland 8 years post left adrenalectomy.

REFERENCES
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Neurilemmoma (Schwannoma) of the adrenal gland
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It’s a benign tumor composed of Schwann cells. It occurs as a solitary encapsulated subcutaneous tumor of spinal roots, cranial nerves, rarely extending to the internal organs. It generally presents itself as a small tumor (2-3 cm in diameter), in rare cases the tumor can weigh up to 2.5 kg. Case report is about a male patient (YOB 1965) who was diagnosed in 2006 with a hormonally inactive retroperitoneal tumor (60x50 mm). Pathologic finding of the left adrenal gland showed a picture of a begin neurilemmoma and hyperplasia of the adrenal cortex. In 2014 during routine checkup abdominal CT revealed microadenoma and hyperplasia of right adrenal gland.

Key words: Neurilemmoma, adrenal gland, treatment.