Adrenal Ganglioneuroma Presenting With Adrenal Insufficiency After Unilateral Adrenalectomy

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Abstract

Objective: To present a case of Adrenal Ganglioneuroma presenting with adrenal insufficiency after unilateral adrenalectomy.

Methods: Adrenal insufficiency is rare after unilateral adrenalectomy for non-functioning adrenal tumours (1). Subclinical Cushing's syndrome, which is defined as mild and subtle elevations of cortisol level in the absence of clinical signs of Cushing's syndrome with adrenal incidentalomas, is more frequent than previously thought (2). If unrecognized prior to adrenal surgery this could lead to adrenocortical insufficiency due to hypothalamic-pituitary-adrenal axis suppression from the subtle cortisol elevations with life threatening implications (2).

Results: A case of apparent subclinical Cushing's syndrome in a patient with adrenal ganglioneuroma manifesting as adrenocortical insufficiency in the immediate post operative period needing treatment with corticosteroids.

Conclusion: Adrenal ganglioneuroma is a rare cause of subclinical Cushing's syndrome. If subclinical Cushing's syndrome is unrecognized prior to adrenalectomy it may lead to adrenal insufficiency in the immediate post operative period. Dynamic testing for subtle cortisol excess should be done in all patients before adrenalectomy.

INTRODUCTION

Adrenal insufficiency is rare after unilateral adrenalectomy for non-functioning adrenal tumours (1). Incidental adrenal neoplasms, called incidentalomas mostly tend to be benign and asymptomatic (2). Subclinical Cushing’s syndrome (SCS) with subtle cortisol elevations and abnormalities of the hypothalamic-pituitary-adrenal (HPA) axis without clinical signs is more frequent than previously thought (2). The incidence of SCS is approximately 5-10 % of the adrenal incidentalomas (2). Dynamic testing is recommended for evaluating autonomous cortisol production by an adrenal incidentaloma (3). The overnight 1 Mg dexamethasone suppression test (ODST) is recommended as the initial screening test (3). In those patients where surgery is done for adrenal incidentalomas based on size considerations (>4Cms) or suspicion of malignancy then subclinical Cushing’s syndrome should be considered and evaluated due to the risk of adrenal insufficiency perioperatively (3).

Glucocorticoid therapy in the perioperative period and post operative assessment of HPA axis recovery are indicated in patients with SCS (3). We would like to present one such case where the pre-operative cortisol levels were normal, but post operatively the cortisol levels were low with a subnormal Synacthen test confirming adrenal insufficiency.
CASE PRESENTATION

A 43 years old female patient was referred to the Endocrinology department at Al Zahra Hospital, Sharjah for a low cortisol level after she had the removal of a left adrenal adenoma laparoscopically. On post-op day 2 the patient had an AM cortisol in the hospital which was found to be low at 1.9 micrograms per dl (mcg/dl). She was asymptomatic. Her serum sodium was found to be slightly low at 133 mmol/l (Normal range 135-153), potassium was normal at 3.8 mmol/l (Normal range 3.5-5.3). She was in no acute distress and her vital signs were normal. A Synacthen (Cosyntropin, Tetracosactrin) test was ordered which confirmed a low baseline cortisol of 1.8 mcg/dl half an hour after injecting Synacthen it was 10.2 mcg/dl and 1 hour post Synacthen was 15.8 mcg/dl, a subnormal response to Synacthen.

An ACTH (intact) level done prior to giving Synacthen was low at 1.3 picograms/ml(pg/ml) (Normal range 10-60 pg/ml). The patient denied receiving any steroids of any kind either orally or parenterally or intranasally prior to surgery. She did not have any stigmata of Cushing’s syndrome. The patient is known to have type 1 diabetes as well as primary hypothyroidism over several years with reasonable though slightly suboptimal glycemic control (Hba1C of 7.3). She has not noticed any increase in insulin requirements over the past several years. She did not have any stigmata of Cushing’s syndrome. She has no history of hypertension. She did not receive any Etomidate by the anesthesiologist as well. Pre operative hormonal workup done by the urologist showed a 11 AM cortisol level of 14.4 mcg/dl and plasma catecholamines were normal. Serum electrolytes prior to surgery were normal. Pre-operatively she had a CT scan of the abdomen which showed a well defined mass lesion of the left adrenal gland 6.5 X 4.0 Cms without significant enhancement with contrast.

A diagnosis of secondary/tertiary adrenal insufficiency due to subclinical Cushing’s syndrome from the left adrenal tumor was made and she was started on hydrocortisone 25 mgs intravenously every 8 hours which was reduced over the next 48 hours and she was switched to Hydrocortisone 10 mgs BID and she was discharged home in a stable condition with proper instructions regarding stress dosing of hydrocortisone and was also given hydrocortisone 100 mgs/ml vial to be taken intramuscularly in case of an
emergency.

Histopathology of the large adrenal mass (Figures 1, 2, 3, 4, & 5) on the left side of 6.5 X 3.0 X 4.0 cms, showed a smooth encapsulated mass firm in consistency. Cut surface was fleshy pale grayish in color. Microscopic examination showed a benign tumor lesion made up of bundles of schwann cells including very few scattered and mature ganglion cells. The pathological diagnosis was Ganglioneuroma.

Two weeks post discharge the patient came back for follow up feeling good without any complaints. An 8 AM cortisol was done after holding the previous day’s evening dose as well as the morning dose of hydrocortisone and it came back at 12.4 mcg/dl. Presuming that the HPA axis is beginning to recover a repeat Synacthen test was done which showed a baseline cortisol of 10.7 mcg/dl and ACTH level of 33.3 pg/ml. Half an hour post Synacthen serum cortisol was 19.1 mcg/dl and 1 hour post was 22.3 mcg/dl indicating a normal response to Synacthen. The hydrocortisone dose was reduced to 10 Mgs OD. Two weeks later an 8 AM cortisol level was done after holding the AM dose of hydrocortisone and it was normal 12.3 mcg/dl and her electrolytes were normal. She was advised to completely discontinue the hydrocortisone and recommended stress doses of hydrocortisone for any major procedures in the next 6 months.

MACROSCOPIC EXAMINATION

Figure 1
Figure 1: Left adrenalectomy specimen made up of a large tumor mass measuring 6.5 x 3 x 4 Cms, potato like, having smooth capsulated surface.

Figure 2
Figure 2: Cut surface is fleshy, pale grayish in color

MICROSCOPIC EXAMINATION
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**Figure 3**
Figure 3: Benign tumor lesion made up of bundles of Schwann cells. Haematoxylin & Eosin stain (H&E). Magnification X 10 times.

**Figure 4**
Figure 4: Bundles of schwann cells. H&E stain. Magnification X 20 times.

**Figure 5**
Figure 5: Few scattered mature ganglion cells. H&E stain. Magnification X 40 times.

**CASE DISCUSSION**
This interesting patient was found to have an incidental left adrenal mass when she was worked up for left loin pains. Her pre-op hormonal workup for the adrenal mass revealed normal cortisol levels as well as plasma catecholamine levels. On day 2 of the post operative period she was found to have an AM cortisol level which was very low. A subsequent Synacthen test showed a subnormal response to tetracosactrin as well as a very low baseline ACTH (intact) level. She was asymptomatic. The only other laboratory abnormality was a slightly low serum sodium level which goes in favor of secondary adrenal insufficiency. She was treated with intravenous hydrocortisone without any mineralocorticoids. She was stable on the treatment. Within several weeks her
Hypothalamic pituitary adrenal axis improved and she was withdrawn from the steroid treatment.

In retrospect the explanation for the adrenal insufficiency was that she had subclinical Cushing’s syndrome from the left adrenal mass which caused HPA axis suppression. It is to be noted that her cortisol level prior to surgery was normal, emphasizing the need for dynamic testing for subclinical Cushing’s syndrome prior to surgery.

A literature review of reports of adrenal ganglioneuroma causing either pre-clinical or overt Cushing’s syndrome was made. There was one report of virilizing adrenal ganglioneuroma in a woman with subclinical Cushing’s syndrome by Diab et al(6) from the Cleveland clinic. There were reports of Cushing’s syndrome with adrenal neuroblastoma in an infant (Kenny et al)(7), Adrenocortical adenoma and ganglioneuroblastoma in a child( Dahms et al)(8) and ectopic Cushing’s syndrome due to an adrenal ganglioneuroma(Corcuff et al)(9). Adrenal ganglioneuroma is a rare cause of subclinical Cushing’s syndrome.

CONCLUSION

Adrenal insufficiency is rare after unilateral adrenalectomy for non-functioning adrenal tumors. Subclinical Cushing’s syndrome although rare could have enormous clinical implications to the patient if the diagnosis is missed as this could lead to catastrophic adrenal insufficiency in the immediate post op period. Dynamic testing of autonomous cortisol production for all adrenal masses is currently recommended pre-operatively. The current recommendations from AACE include perioperative glucocorticoid therapy and post operative assessment of HPA axis recovery in patients with SCS. Adrenal ganglioneuroma is a rare cause of subclinical Cushing’s syndrome.

References


