Non-functional adrenal gland ganglioneuroma: case report

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Abstract

Background: Adrenal ganglioneuromas in young adults are rare and ill-understood. We report an incidentally detected adrenal gland tumor diagnosed as ganglioneuroma (mature type).

Methods and result: Asian 33 years old male who presented with vomiting and epigastric pain for 2 months. USG and CT-abdomen revealed well-defined rounded hypo-dense non-enhancing homogenous mass of left adrenal gland. Histopathology examination revealed a well-encapsulated benign tumor of mature ganglion cells and Schwann-like cells arranged in fascicles, staining strongly with NSE and s-100 proteins, with adjacent unremarkable adrenal cortex and medulla.

Conclusion: Ganglioneuromas (GN) of adrenal gland in adults are very rare and can have serendipitous presentation. We describe such a non-functional adrenal GN in a young male.

Keywords: Adrenal gland ganglioneuroma, cholecystitis, ganglion cells, catecholamines, neuroblastoma, mediastinum

Introduction

Neuroblastic tumors arise from neural crest cells and encompass a spectrum ranging from neuroblastoma, an undifferentiated, malignant tumor to ganglioneuromas (GN), a well differentiated benign neoplasm, which consists of mature Schwann cells, ganglion cells and nerve fibers. GN rarely occurs in the adrenal gland, hence diagnosis is usually difficult [1]. GN do not secrete excess catecholamines or steroid hormones and are often clinically silent and asymptomatic even if they are large [2]. Detection of this tumor has increased with availability of imaging procedures like ultrasonography (USG)/computed tomography (CT)/magnetic resonance imaging (MRI) however histopathology still remains the gold standard for final diagnosis and ruling out a malignant tumor [3,4].

We describe a case of an asymptomatic, non-functional adrenal GN incidentally diagnosed in young adult.

Case presentation

A 33 years old man presented with complaints of vomiting and epigastric pain of 2 months duration. There was no flank pain, fever, hematuria, pyuria or weight loss. Past/ family history was non-contributory. On examination, he was well-built, afebrile, with pulse 96/min, blood pressure, 160/90 mmHg and body-mass index was 31.8 kg/m². Systemic examination was unremarkable. Abdomen was soft without organomegaly. Routine lab investigations were unremarkable with normal blood sugar, hematology, liver, and renal function profile. Endocrine tests, including cortisol, adrenocorticotropic hormone (ACTH) levels, 24 hours urinary catecholamines were within normal ranges. USG revealed unremarkable liver, spleen and pancreas; gall bladder (GB) was distended with 4 mm sized echogenic shadow partially occupying the lumen reported as calculus/sludge. Both kidneys were unremarkable. A hypo-echoic area with well-defined margins measuring 8.1x8 cm was seen over left supra-renal area displacing left kidney (LK) laterally. CT-abdomen revealed well-defined rounded hypo-dense non-enhancing homogenous mass devoid of necrosis/calcification in the region of left adrenal gland measuring 80x78 mm, indistinct from adrenal gland. Other abdominal, pelvic and urogenital organs were normal except for contracted GB with few tiny calculi in lumen.

MRI T2 weighted image showed well-defined encapsulated heterogeneously enhancing left adrenal mass measuring 77x72 mm with delayed contrast washout, causing mild compression of underlying LK upper pole without invading it (Figure 1). Right kidney and adrenal were normal, lymphadenopathy was absent. Clinical impression was benign left adrenal gland tumor.

Laparoscopic adrenalecstomy was performed to remove Non-adenomatous lesion and sent for histopathology examination. Gross examination revealed adrenal gland with tumor weighing 300 grams, measuring 10x9x7 cm, nodular, irregular, grey colored, congested and partially covered with fatty tissue. Cut section showed a well-encapsulated tumor with homogenous grey appearance (Figure 2). Microscopy examination revealed normal adrenal cortical and medullary tissue with adjacent well-encapsulated benign tumor of mature ganglion cells and...
Schwann cells arranged in fascicles (Figure 3), staining strongly with neuron-specific enolase and s-100 proteins (Figure 4). No immature cells/malignancy was noted. Final diagnosis of ganglioneuroma (mature type) of adrenal gland was made. After one year of follow-up, the patient is asymptomatic.

Discussion
Adrenal GN is uncommon tumor. To our knowledge there are totally less than 500 adrenal GN reported in the world literature. Tumors arising from neural crest cells including sympathetic ganglia and adrenal medulla comprise pheochromocytoma/paraganglioma, neuroblastomas, ganglioneuroblastoma and GN. These are “functional” i.e., catecholamine-producing or “non-functional”. GNs are supposed have low or no metabolic activity. The most common location of GN includes posterior mediastinum and retroperitoneum; only a small proportion of GNs are adrenal in origin. Adrenal GN is more frequent in children & 4th/5th decade of life unlike our patient who was in his 30s. The most common site of GN are retroperitonum (32-52%), mediastinal (39-43%), or cervical region (8-9%) and only rarely in adrenal gland [5,6].

Their serendipitous presentation makes diagnosis difficult and catches the treating physician by surprise. It is usually asymptomatic and hormonally silent. Though GN synthesizes catecholamines, it rarely causes hypertension and is incidentally detected on imaging studies for unrelated symptoms. However some patients do present with epigastric/abdominal pain with diarrhea, vomiting and hypertension as
was noted in our patient [1,7]. Incidentally our patient had associated gall stones and hence calculus cholecystitis was suspected. Long term studies in Chinese population showed 180 incidental adrenal tumors resected, out of which 17 (9.4%) were diagnosed as adrenal GN [3]. Increasing number of these tumors are being found incidentally by USG/CT/MRI especially in asymptomatic cases. Malignant transformation is rarely reported. It needs careful evaluation & surgical resection is the choice of therapy for such tumors [2,3,8].

Conclusion
To conclude, adrenal GN is a rare serendipitous tumor, which is extremely uncommon in adults. It can mimic chronic calculus cholecystitis and definite diagnosis can be established only after histological examination of the resected tumor.

List of abbreviations
GB: Gall bladder
GN: Ganglioneuroma
LK: Left kidney

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
RDP Major contributor in writing the manuscript, analysis and interpretation of patient data regarding the histopathological disease.
AVV Performed the histological examination and helped in preparation of the final manuscript.
HLT Clinical & post operative follow up of patient.

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