Adrenal cysts-a report of four cases including one with bilateral involvement

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Abstract

Adrenal cystic lesions are rare entities. Four types of cystic lesions of the adrenal gland have been described in literature namely, epithelial cysts, endothelial cysts, pseudocysts and parasitic cysts. Tumors of the adrenal gland, both benign and malignant may also show cystic change and have to be considered in the differential diagnosis of cystic lesions of the adrenal gland. Lymphangioendotheliomas, though common at other sites, are unusual in the adrenal glands and bilateral involvement is extremely rare. We present four cases of adrenal cysts, two cases of lymphangioendothelial cysts and two cases of pseudocysts from 108 adrenalectomy specimens received over 5 years. Two of these had hypertension which resolved after surgery. All of our cases did well postoperatively. Prognosis of these benign cysts is excellent and the patients are disease-free after surgery.

Keywords: Adrenal cyst, Lymphangioendothelial, Pseudocyst

Introduction

Adrenal cystic lesions are rare with an incidence of 0.06% in general population. [1, 2]. Various types of cystic lesions of the adrenal gland have been described and have been classified into epithelial cysts, endothelial cysts, pseudocysts and parasitic cysts [1]. Tumors of the adrenal gland, both benign and malignant can also show cystic change and have to be considered in the differential diagnosis of cystic lesions of the adrenal gland [2]. Lymphangioendotheliomas though common at other sites, are unusual in the adrenal glands and bilateral involvement is extremely rare [1,3]. We present four cases of adrenal cysts, one of which was bilateral and three unilateral. Two of the cases including one with bilateral involvement, were classified as lymphangioendothelial cysts and two were pseudocysts.

Case-1: A 48 year old female presented with complains of pain in abdomen since one month, which was dull aching and located in the periumbilical region. Her basal cortisol was 12.8 µg/dl, ACTH and catecolamine levels were normal. CT scan showed bilateral, non- enhancing, cystic adrenal masses with foci of calcification. (Fig1a) A right adrenalectomy was done and we received the adrenal gland entirely as a multiloculated cystic mass measuring 7 x 4 x 3 cm and weighing 30 gms. (Fig 2a) These cysts varied in size from 2 mm to 8 mm. Clear fluid was seen in the cysts. Normal yellowish brown adrenal tissue was seen between the cysts. Microscopy showed adrenal tissue replaced by multiple cysts of varying sizes. The lining epithelium was focally present at places but mostly denuded and the cysts contained eosinophilic fluid material. Focal calcification was also seen in the cyst wall. (Fig 3a, 3b) Cyst lining wherever present

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was positive for CD34 (Fig 3c), CD 31 and negative for cytokeratin. A diagnosis of bilateral lymphangioendothelial cyst was made. Her postoperative course was uneventful and she was discharged on the fourth postoperative day.

Figure-1: CT scan pictures

1a: Case 1.-CT scan showed bilateral non enhancing cystic adrenal masses with foci of calcification
1b: Case 2- CT scan of abdomen revealed right side, single, huge, multiloculated hypodense cystic lesion measuring 12x 10 x10 cm
1c: Case 4-CT scan showed multiple gall stones and a cyst in the right adrenal measuring 8x 5x 8 cm in size

Case-2: A 60 year old male presented with abdominal pain and vomiting since 2 years. He had a history of hypertension since 1 year. Serum cortisol, ACTH, and catecholamine levels were within normal limits. CT scan of abdomen revealed a right-sided single huge, multiloculated hypodense cystic lesion measuring 12x 10 x10 cm. (Fig 1b) A right adrenalectomy was done and the specimen measured 10x 9.5 x 8.5 cm, weighing 70 gms and was composed of multiloculated cysts varying in size from 7 mm to 1 cm. Microscopy revealed adrenal tissue with multiloculated cysts lined by flattened endothelial cells. The cysts contained eosinophilic material. Foci of calcification and fibrosis were also seen. (Fig 3d, 3e) The CD 34 and CD 31 stain was strongly positive in the lining of the cyst wall. (Fig3f) Cytokeratin was negative. A diagnosis of Lymphangioendothelial cyst of the adrenal gland was made. The hypertension resolved after surgery.

Figure-2: Gross

2a: Case 1- Multiloculated cystic adrenal mass
2b: Case 4 - Large uniloculated cyst with smooth inner surface

Case-3: A 51 year old female with pain in the right hypochondrium and breathlessness since 3-4 months. CT scan of the abdomen showed a well defined 6.7 x 6.5 x 6 cm cystic mass in the right adrenal gland. A clinical diagnosis of adrenal cyst or hydatid cyst was available. We received a cystic structure measuring 8 x 5 x 3 cm with adrenal tissue identified at
the periphery. The cyst was uniloculated and contained hemorrhagic necrotic material. Microscopy revealed a cystic structure composed of fibro-collagenous tissue. The cyst had no lining seen despite extensive sections studied. The cyst was however lined by hemorrhagic, necrotic material and showed extensive calcification in the wall. A diagnosis of pseudocyst of the adrenal gland was made.

![Figure-3: Microscopy](image)

3a: Case 1- Cysts contained eosinophilic fluid material  
3b: Case 1- Focal calcification was seen in the cyst wall  
3c: Case1-Cyst lining positive for CD34  
3d: Case 2- Cysts lined by flattened endothelial cells & contained eosinophilic material  
3e: Case 2- Adrenal tissue seen in the wall  
3f: Case 2-Cyst lining positive for CD34  
3g: Case 4-Cyst wall made up of dense fibrous tissue  
3h: Case 4- Calcification and adrenocortical clear cells were seen

**Case-4:** A 50 year old female presented with pain in abdomen since 3 months. CT scan showed multiple gall stones and a cyst in the right adrenal gland measuring 8x 8 x 5 cm in size. (Fig 1c) Her basal cortisol was raised. ACTH and catecolamine levels were within normal limits. She was hypertensive since one year. We received an adrenalectomy and cholecystectomy specimens. The adrenalectomy specimen was composed of a large cyst measuring 7.5 cm in diameter and showed adrenal tissue in the wall. The inner surface was smooth and yellow in color. (Fig 2b) Microscopy revealed the cyst wall made up of dense fibrous tissue with calcification. Adrenocortical clear cells were seen focally in the cyst wall. No lining epithelium to the cyst wall was seen despite extensive sections studied. (Fig 3g, 3h) A diagnosis of pseudocyst of adrenal gland was made. The hypertension resolved after surgery.

**Discussion**

We received total 108 adrenalectomy specimens over a five 5 year period at our institute. Amongst these, four cases of adrenal cysts were received and one of them showed bilateral involvement. Cystic lesions of the adrenal gland though uncommon, should be considered in the differential diagnosis of abdominal mass [4].

Ninety five percent of lymphangiomas are located in the neck or axilla and the other 5 % are located in abdominal cavity including mesentery of small intestine, omentum, mesocolon and retroperitonium [5]. Adrenal lymphangiomas are relatively rare with an incidence of 0.06% in general population. Lymphangiomas are seen at all ages with peak in 3rd to 6th decade of life. All our cases are in the 4th to 5th decade of life. Three of our patients were female and one male. Adrenal cysts are often discovered incidentally but large cysts can present with abdominal pain, hemorrhage, rupture, mass lesion, arterial hypertension especially in functional cysts etc. [6] All four of our cases presented with abdominal pain of varying duration. Hypertension was seen in 2 of our patients and cortisol level was raised in one of them and her hypertension resolved after surgery.
The pathogenesis of the cystic lymphangioendotheliomas is not well known. It is believed that these lesions occur from continued growth of ectopic or malformed lymphatic tissue or represent a hyperplastic reaction to inflammation or a lymphatic hamartomas or blockage of draining lymphatics with proximal dilatation of lymphatic system [5]. Imaging can pick up the cystic nature of these cysts as well as calcification in the cyst wall.

Summary of the four cases.

|                      | Case 1 | Case 2 | Case 3 | Case 4 |
|----------------------|--------|--------|--------|--------|
| **History**          | 48 year, female | 60 year male, hypertensive & diabetic since 1 year | 51 year female | 50 year female, known diabetic and hypertensive |
| **Clinical presentation** | dull ache in periumbilical region since 1 month, no abnormality on abdominal palpation | acute abdominal pain, vomiting since 2 years, tenderness | pain in the right hypochondrium and breathlessness since 3-4 months, no abnormality | diffuse pain in abdomen since 3 months, tenderness |
| **Laboratory investigations** | Basal cortisol-normal, ACTH-normal, Catecolamines-normal | Basal cortisol-normal, ACTH-normal, Catecolamines-normal | Not available | Basal cortisol-raised, ACTH-normal, Catecolamines-normal |
| **Imaging-CT scan**  | multiple cysts of both adrenal glands with peripheral rim of calcification | Right sided single huge, multiloculated hypodense cystic lesion measuring 12x10x10cm | a well defined 6.7 x 6.5 x 6 cm cystic mass in the right adrenal gland | Right sided single uniloculated lesion in the suprarenal region 8.5 x 8 cm |
| **Gross Pathology**  | weight: 30 g, size: 7x4x3 cm, multiple cysts, size range 0.2-0.8cm cysts, contained whitish clear fluid, adrenal tissue identified in the septae of the cyst | weight 70 gm, size:10x 8.5 x 9.5 cm, multiloculated cyst, locule size ranged from 7.3 to 7 to 1x1 cm, inner surface was smooth with yellowish areas, adrenal gland tissue seen in the wall of cyst | size:8 x 5 x 3 cm, cyst which was uniloculated, contained hemorrhagic necrotic material, adrenal tissue identified in the cyst wall | size: 7.5 cm diameter, uniloculated cyst. Inner surface was smooth with yellow areas |
| **Histopathology**   | multiple cysts lined by flattened endothelial cells, septae show muscle fibers, lymphoid aggregates & calcification, eosinophilic material in their lumina, adrenal cortical cells in the septae of cyst. | multioculated cystic structure with cysts lined by flattened endothelium, wall of cyst show focal fibrosis, calcification and lymphocytes, eosinophilic material in their lumina, adrenal cortical cells in the septae of cyst seen | a cystic structure composed of fibrocollagenous tissue lined by hemorrhagic necrotic material, absence of lining epithelium seen and extensive calcification, adrenal cortical cells in the wall | cyst lined by thick fibrous tissue, absence of lining epithelium Wall shows calcification, adrenal cortical tissue within the wall |
| **Immunohistochemistry** | CD 34 positive, CK negative | CD 34 positive, CK negative | - | - |
| **Diagnosis**        | Bilateral lymphangioendothelial cyst of the adrenal glands | Lymphangioendothelial cyst of the adrenal gland | Pseudocyst of the adrenal gland | Pseudocyst of the adrenal gland |

Adrenal cysts are classified into 4 types, endothelial cysts [45%], pseudocysts 39%, epithelial cysts (9%) and parasitic cysts (7%). Endothelial cysts can be divided into two types, angiomatous and lymphangiomatous [4] True vascular endothelial cysts (hemangiomas) are rare entities [4].
Adrenal cystic lymphangioendotheliomas are thin walled multiloculated cystic lesions with clear to brown colored fluid. These cysts are lined by flattened endothelial cells on microscopy. Immunohistochemistry using antibody to CD 31, CD 34 and factor VIII confirm the endothelial lining of the cyst. [1] D2-40 is a specific marker for the lymphatic origin of the cyst wall and co expression of CD 34, CD31, D2-40 and factor VIII related antigen confirms its lymphatic origin. Both our cases of multiloculated cysts of adrenal showed positivity for CD 31, CD 34 and negativity for cytokeratin. D2-40 was not available at our institute. On the basis of light microscopy and CD 31 and CD 34 positivity and cytokeratin negativity; our multiloculated cystic lesions were labeled as lymphangioendothelial cysts.

Psedocysts are the commonest type of adrenal cysts, and are unilocular thick walled cysts and are devoid of epithelium and they can be calcified or even ossified. They can result from hemorrhage into the adrenal gland due to trauma, toxins, infectious process and tumor [5].

Epithelial cysts of adrenal are cysts lined by epithelium and cyst wall lining is positive for cytokeratin. They could represent cystic adenomas [4].

All our patients had surgical removal of the adrenal glands done, and postoperative period was uneventful. Two patients, one with lymphangioendothelial and one with pseudocyst had hypertension which resolved post surgery. Current treatment recommendations state that small asymptomatic lesions be observed while large complicated cysts be excised.[6] Since cystic lesions of adrenal are a heterogenous group of lesion and can even include malignancies with cystic degeneration, a thorough histopathological examination and immunohistochemistry is essential for diagnosis. Prognosis of these non-malignant cysts is excellent and the patients are disease- free after surgery.

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