Syndromes that Link the Endocrine System and Genitourinary Tract

Yasemin ÖZLÜK, İşın KILIÇASLAN
Department of Pathology, Istanbul Faculty of Medicine, Istanbul University, İSTANBUL, TURKEY

ABSTRACT

The endocrine system and genitourinary tract unite in various syndromes. Genitourinary malignancies may cause paraneoplastic endocrine syndromes by secreting hormonal substances. These entities include Cushing’s syndrome, hypercalcemia, hyperglycemia, polycythemia, hypertension, and inappropriate ADH or HCG production. The most important syndromic scenarios that links these two systems are hereditary renal cancer syndromes with specific genotype/phenotype correlation. There are also some very rare entities in which endocrine and genitourinary systems are involved such as Carney complex, congenital adrenal hyperplasia and Beckwith-Wiedemann syndrome. We will review all the syndromes regarding manifestations present in endocrine and genitourinary organs.

Key Words: Urogenital system, Endocrine system, SDH, TMEM127, VHL

INTRODUCTION

In this article, we focus on the relationship between the endocrine system and genitourinary tract. This review includes two main topics. First, the paraneoplastic “endocrine” syndromes that occur in the presence of a genitourinary malignancy, and second, hereditary syndromes including both the endocrine system and genitourinary tract. These are discussed based on findings related to both systems.

PARANEOPLASTIC ENDOCRINE SYNDROMES RELATED TO THE GENITOURINARY TRACT

Paraneoplastic syndromes occur in 15-20% of all cancer patients. Paraneoplastic syndromes include endocrinologic, dermatologic, hematologic, neurologic, and osteoarticular manifestations (1). Paraneoplastic endocrine syndrome (PNES) is characterized by complex signs and symptoms related to ectopic production of a hormone or hormonal substance by tumor cells that do not normally produce these molecules. It is important to recognize these syndromes because they may lead to early recognition of cancer, the hormonal substances produced may be used as tumor markers, and they are a major cause of morbidity and mortality. PNES is usually managed by treating the underlying disease.

PNES caused by genitourinary cancers are rare, but have been reported. Renal cell carcinoma (RCC) is the most frequent urologic malignancy that causes paraneoplastic syndrome, followed by prostate cancer (1). PNES related to genitourinary malignancies are summarized in Table I. Most PNES are associated with neuroendocrine tumors, whereas humoral hypercalcemia of malignancy is primarily linked with squamous cell carcinoma (2). Paraneoplastic syndromes are uncommon among bladder and testicular cancers.

A. Paraneoplastic Hypercortisolism (Cushing’s Syndrome)

Inappropriate synthesis of corticotropin-releasing hormone (CRH) or adrenocorticotropic hormone (ACTH) determines overproduction of gluco- and/or mineralocorticoids from adrenal cortex. It may occur either by secretion of CRH or ACTH. The most common cause is the ectopic production of pro-opiomelanocortin (POMC), which is the precursor polypeptide of ACTH. Ectopic ACTH directly stimulates the adrenal cortex, which causes cortical hyperplasia and hypercortisolism. Clinically, patients with paraneoplastic hypercortisolism present with symptoms of Cushing’s syndrome (CS), such as central obesity, round face, excess sweating, red stria, muscle weakness, osteoporosis, hyperpigmentation, hypertension, insulin resistance, and psychologic disturbances. Muscle weakness caused by marked hypokalemia may be the first and dominant clinical symptom (3).

Among genitourinary malignancies, prostate, bladder, and renal cell carcinoma may cause CS (3-8). Prostate carcinomas are of interest in which normal levels of
testosterone are detected despite medical or surgically castration (9). Although cortisol levels are elevated both in CS due to paraneoplastic hypercortisolism and pituitary gland tumor, paraneoplastic hypercortisolism shows very high ACTH levels and does not respond to dexamethasone suppression test (2).

B. Paraneoplastic Hypercalcemia

Hypercalcemia of malignancy (HHM) is the most common PNES and seen in 30% of all patients with hypercalcemia. HHM is related to the production of parathyroid hormone-related peptide (PTHrP). PTHrP synthesis has been demonstrated in 20% of patients with cancer regarding breast, lung, and genitourinary tract (10-12). Hypercalcemia may be seen due to ectopic hormone production or localized osteolytic paracrine effect of metastatic tumor to bone or hyperparathyroidism, which may be present in patients with cancer. Ectopic hormone production (mostly PTHrP) with or without bone metastases is defined as PNES, but not hypercalcemia itself. HHM is the most common form of PNES among RCCs (13-20%) (13, 14) and squamous cell carcinomas of the bladder (15). RCC is also the most common tumor that causes hypercalcemia both in paraneoplastic and nonparaneoplastic mechanisms. Some 75% of RCCs that cause hypercalcemia are high stage and 50% have metastasize to bone, although the presence and degree of hypercalcemia have been shown not to be associated with grade or survival (16). Prostate cancer that causes hypercalcemia is extremely rare and is mostly related to disseminated bone metastasis (1). Although rare, penile squamous cell carcinoma may also culminate in hypercalcemia (17).

PTHrP has significant control over proliferation, differentiation, and death of many cell types. It binds to the PTH receptor and increases levels of serum Ca²⁺ with cAMP-mediated actions. The increase in serum Ca²⁺ is mediated by an increase in osteoclastic activity, reduced renal clearance of Ca, and decreased renal phosphorus reabsorption. The clinical findings of HHM are seen in various forms. Patients may present with nonspecific symptoms such as asthenia, headache, lack of appetite, vomiting, constipation, and polyuria-polydipsia. A more severe form of HHM with an acute confused or lethargic state or even coma may also be seen especially when Ca²⁺ levels are >12 mg/dl. Shock and death occur if Ca²⁺ levels exceed 18 mg/dl (2). The measurement of PTHrP can also be helpful in the differential diagnosis.

Differential diagnosis include excessive vitamin D, sarcoidosis and bone metastasis in a patient with hypercalcemia, low levels of phosphorus and chloride accompanied by low PTH levels. Although PTHrP causes hypercalcemia, Ca²⁺ levels may be normal or even low in prostate carcinoma with neuroendocrine features and PTHrP secretion. This phenomenon is called ‘bone hunger syndrome’, which is metabolic derangement of metastatic prostate carcinoma. In this syndrome, Ca²⁺ is entrapped in bone related to excessive osteoblastic activity. This process results in hyperparathyroidism and increased osteoclastic activity in bones apart from at metastatic sites. Cleavage and inactivation of PTHrP by high levels of PSA are also involved in the development of this syndrome (18). Hypercalcemia is often the leading cause of death in patients with PTHrP producing tumors (1).

C. Paraneoplastic Hypertension

Hypertension related to RCC was first discovered in a patient whose hypertension was cured after nephrectomy (19). The incidence of hypertension in patients with RCC is reported as 14-35% (20). RCCs that cause hypertension typically show low-grade clear cell histology. The mechanisms of hypertension in RCC include renin secretion, ureteral obstruction, arteriovenous fistulae, renal artery stenosis, polycythemia, hypercalcemia, and cerebral metastasis (21). Renin is normally secreted by the juxtaglomerular...
apparatus of the nephron and its overexpression results in hypertension. Renin secretion is generally seen in juxtaglomerular tumors but RCCs may rarely show renin secretion. Hypokalemia and hyperaldosteronism are accompanied features of juxtaglomerular tumors but not RCCs with renin production (19, 21). There is no clear relationship between hypertension and prognosis among patients with RCC (22). Any etiologic factor that causes ischemia in the kidney increases renin secretion, which is the mechanism in ureteral obstruction, arteriovenous fistulae, and renal artery stenosis. There is no correlation with hypertension and renin level (1). Hypertension is reversible through surgical removal of the renal tumor in most cases (2, 19, 23).

D. Paraneoplastic Alterations in Glucose Metabolism
Abnormalities in glucose metabolism such as hypo- and hyperglycemia occur in RCCs. Insulin and glucagon were shown to be present in RCC tumor extracts (24). Rare cases of RCC leading to diabetes have been published (25, 26). The underlying mechanisms for this syndrome are unknown and no association with metastasis and previous history of diabetes also has been shown. It has been suggested that hormones secreted by RCCs antagonize the effect of insulin or promote glucogenesis and this could cause hyperglycemia (1).

E. Paraneoplastic Polycythemia (Inappropriate Erythropoietin (EPO) Secretion)
Polycythemia in RCCs accounts for 1-8% of cases (27). It is believed to be associated with inappropriate secretion of EPO. Although EPO production is documented in 66% of RCCs, erythrocytosis is only seen in 8% (27). The other renal tumor that also shows increased EPO secretion is nephroblastoma (28, 29). EPO expression in tumor cells has been demonstrated in protein and mRNA levels (30-32). EPO production is induced through either hypoxia inducible factor (HIF) or von Hippel Lindau (VHL) mutations without HIF activation (33, 34). The VHL gene is the most-frequently involved gene in the pathogenesis of RCC both in sporadic and familial forms (35). Clear cell RCC is thought to originate from proximal tubular epithelium. In contrast with tumor cells, normal proximal tubular epithelial cells were not shown to express EPO even in hypoxic conditions (31). This finding suggested that VHL mutations may play a role in tumor cells that transform into EPO-secreting cells. EPO is more likely involved in local tumor progression rather than metastasis (36). There are conflicting results in terms of the effect of EPO and EPO-receptor expression on the prognosis of RCC (37-39).

F. Paraneoplastic Inappropriate Human Chorionic Gonadotropin (HCG) Secretion
Elevated levels of HCG have been demonstrated in urine and/or serum samples of genitourinary carcinomas including renal, prostate, and bladder (40). Clinical findings for elevated HCG in males are gynecomastia and decreased libido. There is no direct evidence that tumor cells produce HCG.

G. Paraneoplastic Inappropriate Antidiuretic Hormone (ADH) Secretion
The syndrome of inappropriate ADH (SIADH) is defined as serum hyponatremia, hypo-osmolality and elevated urinary sodium concentrations with normal renal and adrenal homeostasis (41). Although the plasma level of ADH is increased, it is not crucial for the diagnosis of SIADH (2). SIADH may be seen in various clinical situations: malignancies, pulmonary and neurologic diseases, endocrine disorders, and as a drug adverse effect. Lung small cell carcinomas are the most frequent malignant tumor in which SIADH is seen (1). Prostate cancer rarely presents with SIADH and most cases are poorly differentiated adenocarcinomas (42). Primary small cell carcinoma of the prostate is very rare and its SIADH association is even rarer (43, 44). Besides high levels of ADH, positive immunoreactivity for anti-ADH antibody has been documented (44). Bladder small cell carcinoma showing SIADH is extremely rare (45, 46). No other SIADH-related genitourinary malignancies are reported in the literature.
This syndrome is characterized by multiple benign and malignant neoplasms in various organs such as kidney, eye, brain, spine, pancreas, and the adrenal. The neoplasia spectrum includes retinal and craniospinal hemangioblastomas; renal, pancreatic and hepatic cysts; adrenal, hepatic, and pulmonary hemangiomas; and solid tumors. Solid tumors include clear cell RCCs, pheochromocytomas, paragangliomas, pancreatic neuroendocrine tumors, endolymphatic sac tumors, epididymal and broad ligament papillary cystadenoma (49). VHL is not rare and accounts for 1 per 36,000 live births and the penetrance is 95% at 65 years of age (49).

The diagnosis of VHL depends on the presence of a family history. A single VHL-associated neoplasm is enough to diagnose VHL if a family history is present. However, cases with de novo mutations do not show a family history. In this setting, at least 2 VHL-associated neoplasms are required for diagnosis (49, 50). Genetic counseling should be offered if a suspicion of VHL disease is present. The clinical classification of VHL is based on genotype-phenotype correlation. Type 2 VHL differs from Type 1 with the occurrence of pheochromocytoma and is further subdivided according to the risk of RCC (2A: low risk, 2B: high risk, and 2C without RCC) (50).

The protein product of VHL gene, pVHL, regulates HIF-1 and HIF-2, which are transcription factors that regulate hypoxia-responsive genes including vascular endothelial growth factor (VEGF), platelet-derived growth factor (PDGFb), transforming growth factor (TGFa) and erythropoietin (51). The VHL gene is inactivated when both alleles are involved (Knudson’s two-hit hypothesis) (52). Inactivation of the VHL gene results in upregulation of HIF-1&2, which promote angiogenesis and cell proliferation.

Renal tumors in VHL are bilateral, multiple, and accompanied by numerous cystic background lesions (49, 53) (Figure 1A-D). Multiple renal tumors and small microscopic foci (tumorlets) are morphologically typical of clear cell RCCs, and renal cysts are lined by flat epithelial cells with clear cytoplasmics. The cyst epithelium may show stratification, tufting and papillary structures. These cystic lesions showing epithelial changes listed above, are likely to represent early neoplastic transformation (54). Although diagnostic terminology for these cysts has not been standardized, descriptive terms such as “renal cyst with epithelial proliferation” or “renal epithelial cyst with atypia” have been suggested (54). These cystic lesions may mimic clear cell-tubulopapillary RCC in which multilocular masses are confluent, unlike individual cysts of VHL.

<table>
<thead>
<tr>
<th>Hereditary syndrome</th>
<th>Involved gene</th>
<th>Genitourinary manifestations</th>
<th>Endocrine manifestations</th>
</tr>
</thead>
<tbody>
<tr>
<td>von Hippel–Lindau syndrome</td>
<td>VHL</td>
<td>Multiple bilateral CCRCC</td>
<td>PHEO/PGL</td>
</tr>
<tr>
<td>Birt-Hogg-Dubé syndrome</td>
<td>BHD</td>
<td>Oncocytic renal tumors (oncocytoma, chromophobe RCC, hybrid oncocytic tumors with clear cells)</td>
<td>Medullary thyroid carcinoma</td>
</tr>
<tr>
<td>Hereditary leiomyomatosis RCC</td>
<td>FH</td>
<td>RCC</td>
<td>Adrenal cortical proliferation and PHEO/PGL</td>
</tr>
<tr>
<td>Familial PGL syndromes</td>
<td>SDH (SDH-B,C,D)</td>
<td>SDH mutant renal neoplasia (CCRCC, papillary RCC, chromophobe RCC, oncocytoma)</td>
<td>PHEO/PGL</td>
</tr>
<tr>
<td>Hyperparathyroidism-Jaw syndrome</td>
<td>CDC73</td>
<td>Mixed epithelial stromal tumor Adult nephroblastoma Papillary RCC Multiple renal cysts</td>
<td>Hyperparathyroidism (parathyroid adenoma/carcinoma)</td>
</tr>
<tr>
<td>PTEN hamartoma tumor syndrome (Cowden syndrome)</td>
<td>PTEN</td>
<td>Malformations Papillary RCC Chromophobe RCC Prostate carcinoma</td>
<td>Non-medullary thyroid carcinoma Hashimoto thyroiditis</td>
</tr>
</tbody>
</table>

The association of sporadic bilateral renal masses with clear cell-tubulopapillary RCC morphology with VHL is the subject of debate (55).

The common endocrine manifestations in VHL include pancreatic neuroendocrine tumors (NETs) and pheochromocytoma/paragangliomas (PHEO/PGL).

Pancreatic NETs account for almost 17% of patients with VHL (56, 57). Like in syndromic form of neoplastic diseases, there are precursor lesions for pancreatic NETs in VHL including nesidioblastosis, islet dysplasia (<0.5 mm), pancreatic neuroendocrine (endocrine) microadenomas/adenomatosis (0.5-5 mm) and peliosis of nontumorous islets (58). The tumor should be at least 0.5 cm to describe it as a NET. Histologic grading is performed in accordance with the 3-tiered World Health Organization (2010) system regarding the Ki-67 (MIB-1) proliferation index and mitotic activity (59). Morphologically, besides showing characteristic neuroendocrine morphology, these NETs also contain clear cells or multivacuolated lipid-rich cells (58). Immunohistochemistry reveals positivity for neuroendocrine markers as well as pancreatic and gastrointestinal hormones (60, 61). Inhibin was also shown to be positive in clear cell pancreatic NETs in VHL (60). Clear cell change and inhibin immunoexpression in pancreatic NETs are suggested to be related to HIF1a increase in VHL. VHL is more likely present in cases with multifocal clear cell NETs that express inhibin (60). Pancreatic NETs are malignant, independent of their size or grade (59). Features of local invasiveness, which may lead to life threatening complications, are present in 60% of pancreatic NETs in VHL (56). Pancreatic cysts and serous cystadenomas may also be present in 17-56% of the patients with VHL (57, 61). Pancreatic NETs are multiple well-demarcated, brown

Figure 1: von Hippel Lindau syndrome with multiple cysts and cystic renal cell carcinoma. A) Simple cyst, B) Cyst with papillary proliferations, C) Atypical epithelial cells lining the cyst D) Cystic clear cell renal cell carcinoma (H&E).
to yellow masses most commonly on a cystic background throughout the pancreas (62). Pancreatic cysts are generally asymptomatic and do not need treatment. The mean age at which neuroendocrine tumors develop is 35 years, whereas for pancreatic NETs is 37 years (53).

Pheochromocytomas are seen in 10-20% of patients with VHL (63) and the mean age at presentation is 20 years (64). They are multiple and bilateral (65). Although pheochromocytomas are more common in VHL, extraadrenal paragangliomas may also be seen parasym pathetic in glomus jugulare, carotid body, and periaortic tissue (66). PHEO/ PGLs in VHL are less likely to present with paroxysmal type of hypertension because they show increased levels of dopamine and norepinephrine (65).

The outcome of VHL depends on NETs whose metastatic risk was reported as 11-17% (56). Recent development of techniques like nephron-sparing surgery and radiofrequency ablation used for the treatment of RCCs improved the survival related to the renal masses (67).

A2. Birt-Hogg-Dubé (BHD) Syndrome

BHD is an autosomal dominant syndrome characterized by the occurrence of fibrofolliculomas, lung cysts and spontaneous pneumothorax, and renal cancer (68-70). BHD is seen in approximately 1 per 200 000 individuals. Intestinal polyposis (71, 72) and medullary thyroid carcinoma (70) in patients with BHD has also been reported. However, the general consensus is that BHD is not associated with intestinal polyps or colonic cancer (73). BHD is overlooked because of its mild variable presentation. The gene for BHD was identified in 2001 in affected families and the gene product was named folliculin (FLCN) (74). Although its function has been controversial, a relation to cilio-pathy and involvement in cell polarity, regulation of cell-cell adhesion, and negative regulation of rRNA synthesis was suggested recently (75-77).

Renal tumors seen in BHD are papillary RCC, clear cell RCC, chromophobe RCC, and oncocytoma (78). The underlying renal parenchyma typically shows “renal oncocyto sis”, which is defined as the presence of diffuse oncocy tic changes including dominant oncocy tic neoplasm, smaller oncocytic nodules, infiltrating oncocy tic cells, cortical cysts lined by oncocy tic cells, and oncocy tic change in nonneoplastic renal tubules (54). An unusual form of a low-grade oncocy tic renal tumor with features of both chromophobe RCC and oncocytoma, called “hybrid oncocy tic tumor”, is the most common and characteristic tumor for BHD. The most common feature of this tumor is the presence of individual tumor cells with clear cytoplasm without prominent nuclear membrane irregularity. If a patient has a hybrid oncocy tic tumor, BHD should be considered (79). The diagnosis for BHD requires at least one major or two of the minor criteria previously defined by Menko et al. to be fulfilled (80). The major criteria are as follows: 1- adult onset at least 5 fibrofolliculomas, one of which is histologically confirmed, 2- FLCN mutation. Minor criteria are: 1- multiple lung cysts, 2- renal cancer (early onset, aged <50 years, multifocal/bilateral, hybrid oncocy tic morphology), 3- a first degree relative with BHD.

A3. Hereditary Leiomyomatosis RCC (HLRCC) Syndrome

HLRCC is an autosomal dominant syndrome associated with a germline mutation in fumarate hydratase (FH) gene characterized by cutaneous and uterine leiomyomas in association with RCC (81). The biallelic inactivation of the FH gene required for the development of HLRCC acts as a tumor suppressor gene (82). Although hereditary renal cancers are mostly multiple and bilateral, RCCs in HLRCC are unilateral and solitary (54). RCCs that occur in HLRCC are frequently high stage and fatal (83). These RCCs strongly resemble type 2 papillary RCCs. Although papillary features are frequently prominent, tubulopapillary, tubular, solid and cystic elements, collecting duct carcinoma-like areas with infiltrating tubules, nests or individual cells in a desmoplasmatic stroma are also present (84, 85). Characteristic morphologic features of tumor cells in HLRCC are the distinct nuclei with prominent eosinophilic nucleoli surrounded by clear halo-like CMV inclusions (85). The striking feature in this syndrome is that tumor cells of leiomyomas also contain these inclusion-like eosinophilic nucleoli. Cysts lined by epithelial cells that share the same cellular features with HLRCC are reported as precancerous lesions (86). The morphologic spectrum for leiomyomas seen in HLRCC were described as: fascicular growth pattern, staghorn vasculature, cytoplasmic fibrillarity, and characteristic eosinophilic nucleolus surrounded by a perinuclear halo (87-90).

There is no validated marker to determine patients with HLRCC. However, Bardella et al. (91) showed cytoplasmatic positivity for 2SC-modified proteins (2SCP) using immunohistochemistry and demonstrated a correct prediction for the mutation in FH gene. The immunopositivity of anti-2SC was explained by succination of cellular proteins due to the accumulation of high levels of fumarate, which form a stable chemical modification, S-(2-succino)-cysteine (2SC). The prospective use of this marker was suggested as a screening method for HLRCC. In a recent study by Chen et al. (84), this antibody was shown to be highly sensitive, and nuclear and cytoplasmic
staining was also specific for the diagnosis of HLRCC. In another study performed on uterine leiomyomas related to HLRCC, morphological reproducibility across cases and anti-2SC immunopositivity were demonstrated (90). This antibody is not commercially available and further validation studies are necessary. Identifying patients with HLRCC provides early detection of renal tumors before they become aggressive.

Adrenocortical hyperplasia was reported in a patient with HLRCC with a proven FH mutation (92). The patient had uterine and cutaneous leiomyomas but no renal tumor. ACTH-independent bilateral adrenocortical hyperplasia and clinical Cushing’s syndrome was demonstrated. In a retrospective review of 15 patients with adrenocortical hyperplasia and/or HLRCC only 3 patients, including the index case, were detected. However, FH mutation in the tumor DNA was not demonstrated in the other 2 patients. This case report suggests a relationship between adrenocortical hyperplasia and HLRCC. In another recent study by Shuch et al. (93) 7.8% of HLRCC cases showed uni- or bilateral adrenal nodular hyperplasia with histologically proven absence of adrenal malignancy.

### A4. Familial Paraganglioma and Pheochromocytoma (PGL/PHEO) Syndromes

Familial PGL syndromes belong to inherited renal syndromes and PHEO/PGL syndromes (94-96). Heritable pheochromocytoma syndromes include neurofibromatosis type I, VHL, multiple endocrine neoplasia type 2, familial PGL syndromes (types 1-4), and familial pheochromocytoma syndromes (TMEM127-gene-associated, MAX-gene-associated and SDHA-gene-associated) (97).

Familial PGL syndromes include adrenal/extraadrenal pheochromocytoma (with high frequency of head and neck PGL), RCC, thyroid carcinoma, and gastrointestinal stromal tumor (97). At least 30% of PHEO/PGL are hereditary (98). The gene involved in these 4 types of syndromes is succinate dehydrogenase (SDHB, SDHC, SDHD, SDHAF2) (Table III). SDH, also known as mitochondrial complex 2, is a mitochondrial respiratory enzyme that links the Krebs cycle to the electron transport chain. Double-hit inactivation almost always occurs in the presence of a germline mutation. SDHC and SDHD are anchoring components, whereas SDHA and SDHB are catalytic components. Any double-hit mutation in one of the components makes the entire molecule instable (99). In hypoxic conditions, HIFs provide cell adaptation and survival by inducing several genes on angiogenesis, energy metabolism, survival and growth. The tumorigenesis mechanisms involved in both SDH deficiency and VHL mutation are very similar to hypoxia and have been named as the pseudohypoxic response. Unlike VHL- and SDH-related tumors, on the other hand, the underlying mechanisms are refered to the activation of kinase receptor signaling pathway as in tumors associated with NF1, KIF1BB, TMEM127 and MAX-mutations (99).

SDHB immunohistochemistry is negative in tumor cells in all SDH gene-related tumors but positive in nontumoral cells in a granular (mitochondrial) pattern as an internal control. If SDHB is negative, then SDHA immunohistochemistry should be performed because SDHA and SDHB are both lost in cases with SDHA mutations, whereas SDHA expression is retained in the presence of SDHB, SDHC or SDHD mutations (100). SDH mutations account for almost 15% of all cases of PHEO/PGL (98). SDHB-driven PHEO/PGLs show very high malignant potential (99).

Renal neoplasms have been reported in association with germline SDH mutations, SDHB related tumors being the

---

### Table III: Paraganglioma syndromes and related tumors

<table>
<thead>
<tr>
<th>PGL Syndrome</th>
<th>Gene</th>
<th>Related tumors of the endocrine and genitourinary systems</th>
</tr>
</thead>
<tbody>
<tr>
<td>PGL Type 1</td>
<td>SDHD</td>
<td>• PHEO/PGL (head&amp;neck &gt; adrenal &gt; abdominal extraadrenal &gt; thoracic)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• RCC (very rare)</td>
</tr>
<tr>
<td>PGL Type 2</td>
<td>SDHAF2</td>
<td>• Head and neck PGL</td>
</tr>
<tr>
<td>PGL Type 3</td>
<td>SDHC</td>
<td>• Head and neck PGL</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• RCC (clear cell, papillary type)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• PHEO (very rare)</td>
</tr>
<tr>
<td>PGL Type 4</td>
<td>SDHB</td>
<td>• PHEO/PGL (abdominal extraadrenal &gt; adrenal &gt; head&amp;neck &gt; thoracic)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• RCC</td>
</tr>
</tbody>
</table>

PGL: Paraganglioma; SDH: Succinate dehydrogenase; PHEO: Pheochromocytoma; RCC: Renal cell carcinoma.
most frequent. SDHC and SDHD-related renal tumors are very rare. SDHB-deficient RCCs, introduced as a provisional entity in the last Vancouver classification of renal tumors, are well-circumscribed tumors comprised of granular eosinophilic (oncocytic) cells that have intracytoplasmic flocculent eosinophilic inclusions accompanied by intratumoral mast cells (73, 101, 102). Although this unique morphology is distinctive, it is not present in all cases. The clinical clues are helpful as well as the typical morphology (if present) in SDHB immunohistochemical screening. These clues are bilaterality, multifocality, young-age onset, or a personal or familial history of renal carcinoma or PHEO/PGL or GIST (99). Germline SDHB mutated RCCs without syndrome-specific features account for 5% of inherited RCCs (103). Various histopathologic forms of renal tumors may occur including clear cell, papillary, and oncocytoma (104). Genotype-phenotype correlation was detected for SDHD mutations, whereas it could not be shown for SDHB. Missense mutations in SDHD gene were associated with a low risk for pheochromocytomas (104).

Germline TMEM127 mutation has been reported in hereditary RCCs (105) and PHEO (106, 107). TMEM127 plays a role in the modulation of endolysosomal function and its mutation activates mTOR pathway (105). The two-hit model of tumor suppressor gene inactivation is present among patients with TMEM127 mutation (108). In contrast to SDHB mutations, TMEM127 mutation-related pheochromocytomas do not exhibit norepinephrine or epinephrine (106, 107). Adrenal medullary hyperplasia was a reported endocrine manifestation among patients with TMEM127 mutation-related PHEOs (106). TMEM127 mutation was recently reported in association with co-occurrence of PHEO and renal carcinoma (109)(Figure 2). Although genotype-phenotype correlation is often present among hereditary RCCs, a possible correlation for TMEM127-related RCCs is under debate because of the limited number of cases.

A5. Hyperparathyroidism-Jaw Tumor Syndrome (HPT-JTS)

Hyperparathyroidism-jaw tumor syndrome (HPT-JTS) is a rare autosomal dominant disease related to a germline mutation in CDC73 (also known as HRPT2) tumor suppressor gene encoding parafibromin. There are also some other tumor suppressor genes (TROVE2 and GLRX2) that are also under investigation for HPT-JTS (110, 111). HPT-JTS is characterized by the presence of parathyroid neoplasms such as adenoma or carcinoma with associated hyperparathyroidism, fibroosseous lesions (ossifying fibroma) in the mandibula and maxilla, renal cysts, and renal and uterine neoplasms (112). Hyperparathyroidism is mostly associated with parathyroid hyperplasia or adenoma but parathyroid carcinoma is seen in 15% of patients. Parathyroid carcinoma diagnosis is extremely difficult and depends on the presence of metastasis or local aggressiveness (113). Bone lesions show dense relatively avascular fibroblast-rich stroma, irregular spicules of woven bone with focal rim of osteoblasts (113).

Renal cysts may be numerous and resemble polycystic kidney disease. Renal neoplasms were reported to be mixed epithelial stromal tumor, adult nephroblastoma (Wilms tumor) and papillary RCC (54). Renal hamartomas and uterine carcinomas were also reported in this syndrome (113). Ossifying fibromas are related to the mutation in tumor suppressor gene, but not to hyperparathyroidism (114). Although immunohistochemistry for parafibromin in parathyroid tissue has been reported in HPT-JTS (115), there are no studies regarding renal neoplasms.

A6. PTEN Hamartoma Tumor Syndromes (Cowden Syndrome)(CS)

PTEN hamartoma tumor syndromes (Cowden syndrome, Bannayan–Riley–Ruvalcaba syndrome, Proteus syndrome, and Proteus-like syndrome), is characterized by multiple hamartomas and breast, thyroid, and endometrial carcinomas, is an autosomal dominant hereditary cancer syndrome associated with a germline mutation in PTEN gene (116, 117). The diagnosis is made based on the combination of pathognomonic, major and minor criteria as previously described (118). No genotype-phenotype association was
Other hereditary Syndromes with Genitourinary and the treatment has shown clinical benefit (122). In another study that included patients with clinical findings highly suggestive of CS, there was 16.7% penetrance (120). Unlike other hereditary renal syndromes, there is no family history in CS (119). Histologic types of RCC detected in PTEN hamartoma tumor syndrome are 75% papillary and 25% chromofobe RCC. No specific morphologic feature is present for this syndrome. Loss of immunohistochemical expression of PTEN has been demonstrated (119). Another genitourinary malignancy reported to be associated with CS is prostate carcinoma in younger patients and this has an aggressive clinical course (121).

Loss of heterozygosity in PTEN is common and leads to activation of the mTOR pathway; targeting this pathway in patients with germline PTEN mutation occurred in 30% (129). The penetrance of CNC is reported as 70-80% at 40 years of age (130). Familial history is present in 70% of cases, whereas a new mutation occurred in 30% (129). The penetrance of CNC is reported as 70-80% at 40 years of age (130). The diagnosis of CNC requires the presence of two of the disease manifestations or if one disease manifestation is present, additional supplemental criteria (such as effected first-degree relative or inactivating mutation in PRKAR1A gene) are needed (124).

PPNAD is the most frequent endocrine manifestation seen in CNC. This entity is the bilateral hyperplasia of adrenal glands that causes Cushing’s syndrome in 30% of cases (124). Adrenal glands consist of deeply-located, well-demarcated but unencapsulated brown nodules less than 1 mm in diameter. Lipofuscin is the pigment within the tumor cell cytoplasm that gives the macroscopic brown color. Adrenal glands in PPNAD are normal in size and weight (125). Although normal adrenal cortical cells do not express synaptophysin, PPNAD shows synaptophysin immunoreactivity (131). Transformation to adrenocortical carcinoma was also reported in the background of PPNAD (132-134).

Thyroid involvement is also seen in CNC. The lesion spectrum changes from hyperplastic nodules with or without cystic change to carcinomas, either papillary or follicular type (124, 128). Most of thyroid tumors are adenomas. Carcinomas are very rare. Patients with PRKAR1A mutation are more likely to have thyroid tumors than those without mutation (128).

LCCSCT are present in one third of males with CNC at presentation. Clinically, these tumors may produce hormones that may cause sexual precocity in young males with low gonadotropin levels, as well as gynecomastia due to aromatase overactivity (135). Infertility is also a feature in CNC with unclear underlying etiology in which obstruction by the tumor may be a contributory factor (136). LCCSCTs are multifocal and bilateral in CNC and rarely are sporadic. Tumors show macrocalcifications that can be palpated on physical examination. Grossly, they are well-circumscribed yellowish masses usually less than 4 cm in diameter. Cystic changes may be present. Calcifications and necrosis can also be found. The tumors are composed of eosinophilic cells within a myxoid to collagenous stroma with neutrophilic infiltration (Figure 3A,B). Tumor cells are arranged in sheets, nests, cords, ribbons, and trabeculae. Usually, focal solid tubule formation is present. Calcifications are usually prominent with large laminated nodule formation. Malignant LCCST is extremely rare; only one case has been reported in CNC (137).

B2. Congenital Adrenal Hyperplasia (CAH)

CAH is an autosomal recessive disease related to a defect in cortisol synthesis (138). 21-hydroxylase deficiency is the cause in more than 90% of the cases (139). Cortisol deficiency results in increased ACTH levels. ACTH stimulation on adrenal cortex leads to overproduction of adrenal androgen precursors. There are two forms of CAH, the classic form with salt wasting and simple virilizing subtypes, and the nonclassic form, which is more frequent. The classic form is severe, whereas the nonclassic form is mild and late onset. Recent developments in treatment strategies of this
syndrome lead to a wide spectrum of patient characteristics from birth to adult life. The pathophysiology, clinical details, and treatment are complex and beyond the scope of this article. Here we will only review testicular adrenal rest tumor related to CAH.

One of the challenging genitourinary and endocrine manifestations of CAH is testicular adrenal rest tumor (TART), which is the most important cause of infertility. TART cause infertility due to both mechanic and paracrine effects (140). They are small tumors that vary from mm to cm, are mostly bilateral and located close to or within the rete testis. They are palpable if they reach >2 cm. The gross examination shows yellow-tan multilobular masses. Microscopy resembles that of adrenal cortex. Tumors are well-demarcated but nonencapsulated. Tumor cells have abundant eosinophilic cytoplasm arranged in sheets and cords (Figure 4A,B). Tumor cells are separated by thin fibrovascular septa. Typical zonation of adrenal cortex is absent. Lipofuscin pigment can be seen within the tumor cells (Figure 4C).

TARTs are composed of adrenal-like cells with ACTH and angiotensin II receptors, and also adrenal-specific enzymes and steroids (141). The pathogenesis of TART was investigated through embryology. The adrenal gland and gonad are very close to each other in the embryo and during emigration, some adrenocortical cells may settle in the rete testis (Figure 4B). Increased ACTH concentrations induce these cells to proliferate. Both the concentrations and duration of ACTH exposure determine the tumor development (142, 143). The other proposed theory is that hilar pluripotent cells react to increased ACTH production and proliferate (144).

TART and Leydig cell tumors are difficult to differentiate morphologically, the latter have Reinke crystals 25-40%.

Figure 3: Large cell calcifying Sertoli cell tumor with A) Calcification and prominent neutrophils in the stroma B) Tumor cells arranged in cords, trabecula, and tubules (H&E).
Bilaterality is more frequent in TART, whereas malignant transformation may be seen in Leydig cell tumor but not TART. Lack of cytologic atypia, low mitotic activity, dense fibrous septa, lymphoid aggregates, adipose metaplasia, and lipofuscin pigment favor a diagnosis of TART. However, TART may show endocrine atypia with greater degree of nuclear pleomorphism (145)(Figure 4D). TARTs are positive for CD56, synaptophysin, and negative for androgen receptor. On the other hand, Leydig cell tumors express androgen receptor but show no or weak expression for CD56 and synaptophysin (145, 146).

**B3. Beckwith-Wiedemann Syndrome (BWS)**

BWS is a pediatric overgrowth syndrome that predisposes to tumor development (147). The hallmarks of BWS are exomphalos, macroglossia, and gigantism. However, not all cases exhibit these features. No sex predilection is present for the frequency of BWS (148). The pathogenesis of BWS is highly complex and thus we should first remember normal status and understand genomic imprinting. Autosomal genes are mostly expressed from both maternal and parental alleles. However, in less than 100 genes the allele from only the parental or maternal allele is expressed. The other allele that is not expressed becomes silenced. This mechanism is called genomic imprinting and is controlled by epigenetic mechanisms such as DNA methylation, histone modification, and noncoding RNAs. This process leads to differences in DNA methylation. These imprinted genes present in clusters called imprinted domains and these are regulated by imprinting centers, which are on the same chromosome. Epigenomic or genomic deregulation of imprinted gene clusters on 11p15.5 region results in BWS (147). This critical region includes various genes: insulin-like growth factor 2 (IGF2), H19, cyclin-dependent kinase

---

**Figure 4:** Testicular adrenal rest in a child with congenital adrenal hyperplasia. **A)** The tumor separated by fibrohyaline bands, **B)** Localized next to the rete testis, **C)** Show granular cytoplasm with lipofuscin pigment, **D)** Focal endocrine atypia (H&E).
inhibitor 1C (CDKN1C), potassium channel voltage-gated KQT-like subfamily member 1 (KCNQ1), and KCNQ1-overlapping transcript 1 (KCNQ1OT1, or long QT intronic transcript 1). The genetic alterations that occur in BWS are associated with specific phenotype–epigenotype/genotype correlations, which lead to clinical heterogeneity. BWS are usually sporadic (80%) but autosomal dominant inheritance may be present with a frequency of 15% (147).

The clinical features of BWS vary and include macroglossia, prenatal and postnatal overgrowth, abdominal wall defects, pancreatic islet hyperplasia (hypoglycemia) (149). The components of the syndrome vary depending on the underlying genetic/epigenetic alterations. Patient types regarding genotype and phenotype are beyond the scope of this article. We only include adrenal and genitourinary tract involvement in this paper. Adrenal cytomegaly is defined as the presence of markedly enlarged cortical cells with eosinophilic cytoplasm. The nuclei are pleomorphic and hyperchromatic. There are intranuclear cytoplasmic pseudoinclusions in the cortical cells. This entity is present in neonates but more common in premature infants with Rh incompatibility. BWS is another condition in which adrenal cytomegaly is seen. Both adrenal glands are involved and hyperplastic in BWS (149). Adrenal glands are enlarged with a gross cerebriform appearance. Cytogemalic cells are greater in number than isolated adrenal cytomegaly and they mimic CMV infection but lack typical eosiophilic intranuclear inclusions (149). Hemorrhagic adrenal macrocysts may also occur (150). Adrenal adenomas and carcinomas also occur in BWS with increased frequency (151, 152). Various pediatric malignant tumors also develop in patients with BWS such as nephroblastoma (Wilms’ tumor), neuroblastoma, and hepatoblastoma. Renal abnormalities like nephromegaly, nephrocalcinosis, and later development of medullary sponge kidney are some other genitourinary tract abnormalities of BWS (149).

CONFLICT OF INTEREST
The authors have declared no conflict of interest.

REFERENCES


