Well differentiated thyroid tumor of unknown malignant potential (WDT UMP) associated with COWDEN syndrome

Enaam Junainah*, Mohammed AlShanbari, Nashat Gandoura, Nadia Enani, Khalid Alnouri, Sahar Fatta, Jamal Junainah, Abdulrehman Sber, Saeed Alamoudi, Manal Junainah and Fawzi Junainah

Abstract
Cowden syndrome is a rare cancer predisposition syndrome inherited in an autosomal-dominant fashion. The syndrome is characterized by hamartomatous polyps that affect multiple organs: skin, mucous membranes, thyroid, breast, gastrointestinal tract, endometrium and brain. It is also associated with an increased risk of developing malignancy in many tissues but especially breast, thyroid and endometrium. Thyroid tumor is ranging from benign nodule to frank malignancy. We report a case of WDT UMP in 66 years old lady who has CS.

Keywords: WDT UMP, COWDEN syndrome, thyroid tumor

Case presentation
A 66- Year-old Saudi female patient known case of chronic bone & joint pain on Osteopenia & Sulfasalazine treatment following with Rheumatology. Presently complaining of neck swelling for 6 months associated with Dysphagia and shortness of breath. No history of hoarseness, no weight loss and no fever. No history of radiation exposure. No family history of thyroid carcinoma. past medical history revealed, multiple hamartomatous colonic polyp treated with endoscopic polypectomy. as well as uterine fibroid treated by hysterectomy. The following year she presented with breast nodules on two separate occasions. Surgical excision was performed on each occasion and histology on each of these occasions revealed benign fibrocystic disease of the breast.

On examination
She was found to have multiple abnormal physical findings. She had muffled face, additional skin lesions with multiple facial keratotic papules, and papillomatosis, particularly around the eyes, mouth, and nostrils, buccal mucosal papules with a 'cobblestone-like' appearance, and cutaneous verrucous papules in the acral portions of the hands, feet, palms, and soles. A breast examination revealed multiple palpable nodules. Additional physical findings included strabismus and skeletal abnormalities with kyphoscoliosis and pectusexavatum. Ear, Nose & Throat --> Unremarkable. Fiberoptic examination shows bilateral mobile vocal cord. Neck shows mid line neck swelling 4x5 cm with retrosternal extension, no tender and no palpable cervical lymph nodes lab investigations all with in normal limits, including thyroid function test-Excised buccal papules sent for histologic examination revealed acanthosis papillomatosis with keratosis epiderma. A breast ultrasound examination showed bilateral nodules compatible with fibrocystic disease.

Ultrasound Neck shows right thyroid lobe is normal average size with 3 hypo echoic nodules the biggest 6x4 mm. increase in area visibility. Isthmus enlarged 5mm, no focal nodules. Left thyroid lobe showed lobe showed large heterogeneous, soft tissue nodules measuring 8.4x5.4cm, occupying mid lower pole no enlarged cervical lymph node.

Impression
Bilateral thyroid nodules
CT Scan Neck also done, showing significant left thyroid lobe enlargement with retrosternal extension. No enlargement of cervical lymph seen (Figure 1) node Nodule FNA shows few group of typical follicular cell exhibit nuclear groove and over lopping. Patient underwent total thyroidectomy.

Histopathological
Examination of the surgically removed specimen (Figure 2) showed that Right lobe measured 4.5x3.5x2.5 cm with intact capsule. Serial sectioning revealed variable sized, well circumscribed and ill-defined nodules ranging from 0.5 to 1.5 cm in diameter. Cut surface was soft and brown with areas of cystic degeneration. The left lobe specimen consisted of single light brown, well circumscribed and capsulated solid nodule measuring 4.5x3x2 cm and weighing 79gram. Serial sectioning showed areas of hemorrhage and cystic degeneration. Microscopic examination revealed thyroid parenchyma with a well circumscribed nodule composed of closely packed follicles (Figure 3) containing scant colloid. Few areas showed variable sized follicles with abundant colloid. The follicles were lined by follicular cells. Fewareas had enlarged nuclei showing nuclear
clearing and mild overlapping. Scattered nuclei had nuclear grooves (Figure 4). Nuclear inclusions were not seen. Focal area show thick capsul with questionable capsular invasion (Figure 5), but no vascular invasion noted. Cytokeratin 19 (CK 19) was positive (Figure 6), Galctin3 (Figure 7) & HBME1 (Figure 8) show focal membranous & cytoplasmic positivity in 30% of the cells. Other areas revealed features suggestive of multinodular goiter.

The isthmus show evidence of has himothyroidaites. Based on findings, it was diagnosed as well differentiated tumor of uncertain malignant potential.

Colonoscopy diagnosed diffuse colorectal polyposis with multiple lesions in the sigmoid and rectum Biopsies of these
polyps. Our patient was diagnosed as having CS based on the syndrome testing criteria adapted by the US National Comprehensive Cancer Network (NCCN) [3].

Discussion and literature review
Cowden syndrome is an autosomal dominant syndrome predisposing to cancer, characterized by the occurrence of hyperplastic hamartomatous and tumoral lesions affecting various organs [23]. The disease mainly affects Caucasian women [24,25]. CS is most often diagnosed during the third decade of life (age range, 13 to 65 years) [24,25].

This syndrome is characterized by a combination of ectodermal, mesodermal, and endodermal alterations that may involve various organs: the skin, mucous membranes, breast, digestive tract, thyroid, and central nervous system. The characteristic skin signs such as facial trichilemmomas, acral keratosis and mucocutaneous papillomas, occur in 99% to 100% of patients and are preferentially localized in the peri-oral and facial regions [24]. These lesions are significant for diagnosis and have little malignant potential [26]. Breast lesions with fibrocystic disease, as observed in our patient, occur in approximately 75% of women. Breast carcinoma has been described in 30% to 50% of patients, and a recent review reported an 81% lifetime risk of breast cancer in patients with CS [27]. No breast cancer was detected in our patient [28,29]. Thyroid disease occurs in two-thirds of patients including goiter, thyroiditis, and thyroid cancer. Gastrointestinal involvement may be found throughout the gastrointestinal tract, frequently in the colon, but rarely in the small bowel [30]. "Gastrointestinal involvement is predominantly in the form of hamartomatous colorectal polyps". Other polyps such as lipomatous, fibromatous, hyperplastic inflammatory and adenomatous lesions have also been described [31]. Esophageal glycogenic acanthosis is present in 40% to 60% of patients with CS and should be pathognomonic [31]. Until recently, it was reported that gastrointestinal involvement was not neoplastic. Heald et al., in a prospective study of 127 PTEN mutation carriers reported that 13% of patients undergoing colonoscopy were diagnosed as having colorectal cancer [33]. This study confirms that patients with CS are at increased risk for cancer.

Additional findings include abnormalities of the female reproductive tract presenting as ovarian cysts (24%), leiomyoma (44%), and endometrial carcinoma (10%) [1]. Central nervous system tumors, ganglioneuromas, neurofibromas, intracranial hypertension, granular cell myoblastoma and meningioma have all been reported in patients with CS [23]. Skeletal abnormalities are described in 37% of patients with CS including adenoid facies, kyphoscoliosis, syndactyly, and brachyphalangia. Other abnormalities include eye dysfunction, pulmonary lipoma, lung cysts, and cardiovascular problems [32]. Our patient had many of these documented physical findings of CS including skin and oral mucosal lesions, thyroid tumor, gastrointestinal polyps, skeletal abnormalities and a history

![Figure 6](image1)

Figure 6. High power examination, original magnification x50
Show strong +3 positive membranous stain for CK19.

![Figure 7](image2)

Figure 7. High power examination, original power x40 show focal +2 membranous and cytoplasmic positivity with Glucin 3.

![Figure 8](image3)

Figure 8. High power examination, original power x40 show focal +2 positive membranous and nuclear stain with HBME 1.
of fibrocystic breast disease. Most patients with CS have a germ-line mutation in the tumor suppressor gene PTEN. The role of PTEN in tumorigenesis has been demonstrated and the loss of PTEN function contributes to cellular transformation, increasing the risk of cancer development in patients at an earlier age. *The mutation is identified in only 80% of patients who meet the clinical criteria. The interesting finding in our patient that she developed thyroid follicular tumor which consider of low malignant potential, as this category of thyroid tumor been recently introduced by WHO, in which The conventional grouping of thyroid carcinoma into the main categories of papillary, follicular, medullary and anaplastic (undifferentiated) carcinoma based on morphological features according to WHO classification [1] has made the diagnosis of these entities very simple and straightforward. However, the argument arise in cases of encapsulated follicular variant of papillary thyroid carcinoma (FVPTC) also called as Lindsay's tumor. This was described originally by Lindsay in 1960 and then later reemphasized by Chen and Rosai [3-5]. These tumors are defined as an encapsulated neoplasm of follicular cells with follicular architecture and having archetypal nuclear characteristics of papillary thyroid carcinoma (PTC). The presence of capsular and/or blood vessel invasion is not a prerequisite for making the diagnosis. On the other hand, nuclear features of papillary thyroid carcinoma in the form of clearing, overlapping, grooves and pseudo inclusions should be wide spread. Treatment of the lesions is not different from management of conventional papillary and follicular thyroid carcinoma since Chang et al., pointed out the blood born metastatic pattern of some cases of FVPTC [10]. To circumvent the low threshold of the endocrine pathologist in the diagnosis of these conditions two categories were proposed in 2000 regarding the terminology of thyroid tumors by the Chernobylpathologists group. These novel definitions were based on the fact that there are several diagnostic uncertainties in terms of recognition of certain varieties of encapsulated thyroid tumors with a follicular architecture [6-11]. Further more, there is propensity to over diagnose the follicular variant of papillary thyroid carcinoma. To overcome this diagnostic dilemma, an editorial was published proposing the category of well differentiated tumor of uncertain malignant potential (WD TUM P). Due to the vagueness of the clinical behavior of these tumors, clinicians and surgeons are often puzzled regarding the treatment of these tumors. More over, there are no guidelines or protocols for the management of these tumors as well. In the management of our case, we also encountered the similar problem and treatment uncertainty but ultimately decided to treat the patient with RAI post thyrodectomy which later on proved to be a sensible clinical decision due to the finding of pulmonary metastasis found on post RAI whole body scan. On the basis of the reported findings, it is tempting to speculate that these new entities should be treated with total thyrodectomy followed by RAI until more long term data is vailable documenting the clinical out come of these lesions.

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

Authors’ contributions

| Authors’ contributions | EJ | MA | NG | NE | KA | SF | JJ | AS | SA | MJ | FJ |
|------------------------|----|----|----|----|----|----|----|----|----|----|----|
| Research concept and design | ✓ | -- | -- | -- | -- | -- | -- | -- | -- | -- | -- |
| Collection and/or assembly of data | ✓ | ✓ | -- | -- | -- | -- | -- | -- | -- | -- | -- |
| Data analysis and interpretation | ✓ | -- | -- | -- | -- | -- | -- | -- | -- | -- | -- |
| Writing the article | ✓ | -- | -- | -- | -- | -- | -- | -- | -- | -- | -- |
| Critical revision of the article | ✓ | -- | -- | -- | -- | -- | -- | -- | -- | -- | -- |
| Final approval of article | ✓ | ✓ | -- | -- | -- | -- | -- | -- | -- | -- | -- |
| Statistical analysis | ✓ | -- | -- | -- | -- | -- | -- | -- | -- | -- | -- |

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