Case Report

Concurrent Renal Chronic Kidney Disease-Metabolic Bone Disorder (Renal Hyperparathyroidism) and Noninvasive Benign Thymoma

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Abstract

Common embryological origin of the inferior parathyroid glands and thymus can lead to rare concomitant lesions of these two glands. The authors reported a 59 year-old female with confirmed chronic kidney disease-metabolic bone disorder (CKD-MBD) on hemodialysis from 12 years whose extensive surgical exploration occasioned by total parathyrectomy enabled discovery and complete removal of a retromanubrial nonmyasthenic thymoma. The clinical and pathological features of parathyroid nodular hyperplasia and noninvasive thymoma are discussed. The case presented here can be added to the few published observations describing a coincidental association between CKD-MBD (i.e. renal hyperparathyroidism) and thymoma.
Keywords: CKD-MBD, renal hyperparathyroidism, thymoma, myasthenia

Introduction

Synchronous or metachronous coexistence of two or more distinct pathological entities often raised problems about their pathogeny, diagnosis and best therapy, but occasionally epitomized new syndromes. Owing to their anatomical connexions and common embryological origin from the third pharyngeal pouch, the rare but possible association between lesions causing hyperparathyroidism i.e. adenoma, hyperplasia and thymoma can be explained. Our aim is to present an unusual case of coincidental association between CKD-MBD and
nonmyasthenic thymoma. To our knowledge this is one of the first observations reported in literature.

**Case report:** Sixty-two patients who underwent surgical exploration of the neck for renal hyperparathyroidism between 1994-2012 were revised. Thirty-seven of these patients had a total parathyroidectomy (9 with autotransplantation). Excision of retromanubrial areolar fibrofatty tissue searching ectopic or supranumerary parathyroids was done in 19 cases revealing one nonmyasthenic thymoma, one thymic cyst and thymic cellular rests in 6 cases.

A 59-year old female with a medical history of CKD-BMD, on hemodialysis from 12 years for an ESRD consecutive to a chronic glomerulonephritis of streptococcal origin is referred to our clinic by a nefrodialysis unit. She accused intense bone pains,
persistent disturbing itching, generalized muscle weakness, depression and massive weight loss (15 kg in few months). Preoperative serum calcium was 2.2 mmol/l, and iPTH level was 1150 pg/ml. Ultrasound evidenced four parathyroid glands of 12-15 mm diameter and a normal thyroid. (Fig. 1)

The patient underwent a standard total parathyroidectomy without autotransplantation. The four unequally enlarged parathyroid glands weighting=300 mg showed nodular hyperplasia with a higher proportion of clear and vacuolised chief cells, no-table nuclear pleomorphism, rare oxyphyls and reduction of the normal fat. (Fig.2,3), However, at the exploration of the retromanubrial space searching ectopic or supra- numerary glands, a round, well defined greyish mass of 20 mm 0 was identified, progressively mobilized and removed
through cervical incision together with the surrounding connective tissues.

Fig. 1: Ultrasound Evidencing the Four Parathyroids and the Normal Thyroid.
Fig. 2: Operative Specimen: the Parathyroid and the Thymic Tumor.
Topography and gross appearance suggested the diagnostic of noninvasive thymoma confirmed by microscopic examination which evidenced a completely encapsulated tumor composed by epithelial oval-shaped cells lacking nuclear atypia and rare non-neoplastic lymphocytes, separated by fibrous bands (type AB stage I benign thymoma)(Fig.2,4) The postoperative course was uneventful and the patient continues the hemodialysis without any evidence of symptoms recurrence after 12 and 24 months followup.
Fig. 3: Nodular Parathyroid Hyperplasia Higher Magnification HE x 200
Fig. 4: Benign Epithelial Thymoma low Magnifications
HE x 100
Discussion

Close anatomical and embryological relations between the parathyroid glands and the thymus can lead to the appearance of rare but various associations between hyperparathyroidism producing lesions with benign or malignant thymoma. Therefore, outside primary hyperparathyroidism, ectopic secretion of PTH or PTH related protein and MEN type 1, our case is one of the few observations of glandular hyperplasia in CKD-MBD coexisting with thymoma described by Palmer and Sawyer (1978), Byrne et al (1989), Suzuki T et al (1997), Suzuki K et al (1998), De Toma et al (20010 and Palin et al (2009).

The routinely excision of the upper retrosternal tissue during surgery of RHp recommended by Rizzoli et al (1994), Welch et al (2012) and Maria et al (2013) can reveal outside of ectopic or
supernumerary glands the presence of cellular nests or even constituted thymic tumors. The clinical aspects of this concurrent pathology are mixed, the long standing evolution on hemodialysis and annoying combined features of ESRD and parathyroid hyperfunction prevailing the discrete presence of a benign noninvasive thymoma or even the symptomatic course more or less rapid of a malignant tumor. Therefore the progressive osteoarthicular pains, neuromuscular and even psychiatric complaints, ectopic calcifications and calciphylaxy or intense pruritus dominate and masks the asymptomatic evolution or nonspecific manifestations of an incipient or even malignant thymoma: fever, sweating, weight loss or belated appearing dyspnea, cough, compressive signs etc. Laboratory usual tests revealed normal calcemia but raised iPTH and of course uremia and hypercreatininemia of renal HP. Selected
investigations can be used only to look for detecting associated problems or tumor spread in thymoma: full blood count electrophoresis and anti-acetylcholine-receptors (indicative for malignancy).

Imaging resort to ultrasound and sestamibi scan in RHp and to chest X-ray, high un-hanced contrast CT scan and MR scan for thymoma as Mari et al (1999) and Ceriani et al (2008) and the lesion may be sampled with a CT guided needle biopsy. A third of these tumors are discovered serendipitous due to chest radiography or CT for imaging of other conditions, for screening purpose or, as in our case, intraoperatively at surgery for any cervical or thoracic lesions. However the final diagnosis in thymoma is the histopathological examination of the surgical specimen.

Despite various advances in medical therapy, parathyroidectomy
is a pragmatic symptomatic therapy in cases of RHp appearing in patients under maintenance dialysis but the optimal technique must be individualized in each case and still to be debated. For us, the best solution for aggressive, refractory forms of RHp is total para-thyroidectomy without auto-transplantation. Even if our patient had no clinical signs and symptoms for myasthenia gravis complete exeresis of the thymic tumor occurred as the most favorable curative alternative.

**Conclusion**

Coexistence of the main types of hyperparathyroidism and thymoma is probably coincidentally and in the lack of myasthenic features the diagnosis can be established only by chance, imaging methods or surgery on a neighboring organ. Although rare the
operating surgeon must be aware of possible presence of such pathology. Concomitant exeresis of all hyperplastic parathyroids together with the thymic tumor will achieve cure of both lesions.

The authors declare that they have no competing interest. All authors read and approved the final manuscript. Author’s contribution: DMR- conceived the study and performed surgery, GM- performed surgery, G – expertise in pathology, DS - drafted the paper, corresponding author

References


