Clinical Reports

Riedel’s thyroiditis associated with hypothyroidism and hypoparathyroidism

F. Marín, R. Araujo, C. Páramo, T. Lucas and L. Salto

Servicio de Endocrinologia, Clínica Puerta de Hierro, Universidad Autónoma, 28035 Madrid, Spain.

Summary: Hypoparathyroidism secondary to Riedel’s thyroiditis is rare, only 2 previous cases having been reported. We present the case of a 36 year old woman with Riedel’s thyroiditis which developed into hypothyroidism and hypoparathyroidism.

Introduction

Riedel’s thyroiditis is a rare disorder in which the normal thyroid parenchyma is replaced by a dense fibrous tissue which extends beyond the limits of the thyroid gland and invades adjacent structures in the neck. The incidence has been calculated to be 0.06% of a total of 56,700 thyroidectomies. We present the case of a female patient with Riedel’s thyroiditis which developed into hypoparathyroidism and hypothyroidism.

Case report

The patient was a 36 year old woman without familial background of thyroid pathology and who did not originate from a geographical area of endemic goitre. She had not undergone cranio-cervical radiation. Nine months before being sent to our hospital, she began to suffer asthenia, somnolence, constipation, weight gain and dryness of skin. Two months later, she noted a painless lump on the front of her neck which led her to seek medical attention. It was discovered that she had intense hypothyroidism with a thyroxine (T₄) level of 15.87 nmol/l and thyroid stimulating hormone (TSH) of 110 mU/l. Echography defined her disorder as diffuse goitre which, in the scintigraphic study, showed no radioisotopic uptake. Her erythrocyte sedimentation rate was 56 mm in the first hour. Antithyroglobulin antibodies were negative and antimicrosomal antibodies were positive at a titre of 1:25,000. The remainder of the analytical study, including blood calcium, was normal. With the diagnosis of hypothyroidism due to Hashimoto’s thyroiditis, she was treated with thyroxine, resulting in the diminution of her goitre and clinical improvement.

One month before coming to our hospital, she began to experience dysphagia with solids and liquids, dysphonia, nocturnal stridor, anorexia and weight loss of 5 kg. She felt a sharp pain in the thyroid area which radiated to both ears.

Examination revealed an asthenic individual with hoarseness of voice. Her thyroid gland was 3 times its normal size, of strong hard consistency, and firmly adherent to the adjacent tissue planes. It was not painful and palpation indicated the absence of adenopathy. The remainder of the physical examination was normal.

Under treatment with 100 µg/day of thyroxine, she presented a serum T₄ of 79.8 nmol/l (normal range: 51.4–154.4 nmol/l), triiodothyronine (T₃) of 1.1 nmol/l (normal range: 1.54–3.08 nmol/l), TSH of 18.38 mU/l (normal range: 0.2–5 mU/l) and a T₃-resin uptake of 24.50% (normal range: 25–35%). Antithyroglobulin antibodies were negative and antimicrosomal antibodies were positive at a titre of 1:6,400. There was no uptake of the radioisotope by the thyroid and echography revealed a diffuse hyperplasia with multiple, poorly defined, solid nodules distributed throughout the gland. Radiography of the thorax showed tracheal stenosis.

Blood calcium, calculated as the mean of three determinations, was 1.85 mmol/l (normal range: 2.17–2.67 mmol/l) and blood phosphorus was 1.56 mmol/l (normal range: 0.95–1.71 mmol/l). Serum proteins, haemogram, creatinine, electrolytes, lipids, liver function and coagulation were all normal. Her parathyroid hormone (C-terminal PTH) level, as determined by commercial kit (Immune Nuclear Corp., Minnesota), was 0.58 µg/l (normal: 0.55–1.4 µg/l).
0.66 ± 0.33 μg/l). There was no clinical alteration derived from hypocalcaemia, nor was there sign of intestinal malabsorption. Fine-needle puncture-aspiration was performed, affording abundant follicular elements and Hürthle cells, which resulted in a cytological diagnosis of follicular neoplasia.

It was decided to operate, and during surgery it was observed that the thyroid gland had been replaced by a fibrous mass of stony hard consistency which infiltrated the neighbouring muscles. Simple isthmectomy was performed to remove 7 g of tissue, but without manipulation of the remainder. Histologically, the specimen was intensely collagenized fibrous tissue with focal areas of lymphocytic infiltration (Figure 1). No thyroid or parathyroid tissue was observed. After surgery, the patient’s local compressive symptoms improved. Serum calcium (mean of three determinations) was 1.77 mmol/l; phosphorus, 1.56 mmol/l; and PTH, 0.40 μg/l. She was treated with thyroxine and vitamin D₃ to maintain normal blood calcium and thyroid hormone levels within satisfactory limits. Eighteen months post-surgery, she remains asymptomatic except for a progressive increase in her hoarseness. Laryngological examination has revealed bilateral recurrent paralysis.

**Figure 1** Riedel’s thyroiditis. The normal thyroid tissue is largely replaced by a densely collagenized connective with some chronic inflammatory infiltrate. Observe some residual thyroid follicles, × 225.

**Discussion**

The appearance of hypoparathyroidism in Riedel’s thyroiditis is a rare occurrence. We have found only one well-documented case and reference to another possible instance in the literature.³ In our patient, as in the two aforementioned cases, there were no clinical manifestations secondary to hypocalcaemia, probably because the involvement of the parathyroid glands was produced by slow but progressive infiltration by fibrous tissue. The clinical and complementary data permit us to reasonably rule out other pathological entities which would justify the presence of hypocalcaemia together with normal PTH levels.

Hypothyroidism is not a usual manifestation of Riedel’s thyroiditis either. Woolner et al.⁴ found 5 myxoedematous patients in a series of 20 cases of Riedel’s thyroiditis. In all of them there was an almost total destruction of the thyroid gland, which is the mechanism invoked to explain the thyroid hypofunction. In spite of the fact that antimicrosomal antibodies were detected in our patient, the levels did not reach those characteristic of Hashimoto’s thyroiditis. Moreover, the marked infiltration of the adjacent tissue confirmed during surgery and the histological study rules out lymphomatous thyroiditis. In this regard, it is of interest to point out that in the initial descriptions of Riedel’s thyroiditis, it was suggested that it might be considered a final stage of Hashimoto’s thyroiditis.³ Presently, however, the two are generally considered to be nosologically different disorders.⁶

To distinguish between Riedel’s thyroiditis and undifferentiated thyroid carcinomas is an important aspect of the diagnostic assessment of this disease. Clinically, it is impossible to make an absolutely certain diagnosis. In our case, fine-needle puncture-aspiration was of no use either since only follicular cells were aspirated, rather than the fibrous material characteristic of this type of thyroiditis. It has even been observed that, on occasion, biopsy itself may be insufficient as there are cases of undifferentiated carcinomas with a significant amount of fibrous component.⁷ The clinical evolution of our patient after more than two years of survival allows us to rule out the existence of undifferentiated carcinoma.

With respect to treatment, isthmectomy is the technique of choice and its application improved the compressive picture of our patient. The control of the thyroxine and PTH deficiencies by routine means was satisfactory. Nevertheless, the patient presents progressive paralysis of the vocal cords, presumably secondary to fibrosis of the recurrent nerves, which demonstrates the occasionally aggressive character of this disorder.
References