Thyrotoxic adenoma followed by atypical hyperthyroidism due to struma ovarii: clinical and genetic studies

Antonio Ciccarelli 1,4, Hernan Valdes-Socin 1, Jasmine Purma 2, Sok Kean Khoo 3, Jacqueline Schoumans 3, Annamaria Cologna 4, Etienne Hamoir 5 and Albert Beckers 1

1Department of Endocrinology, Liege University, Liege, Belgium, 2Institute of Interdisciplinary Research, Campus Erasme, Bruxelles University, Brussels, Belgium, 3Van Andel Research Institute, Grand Rapids, MI, USA, 4Department of Molecular and Clinical Endocrinology and Oncology, Federico II University, Naples, Italy and 5Department of Endocrine Surgery, Liege University, Liege, Belgium

(Correspondence should be addressed to Albert Beckers, Service d’Endocrinologie, CHU de Liege, Domaine Universitaire du Sart-Tilman 4000 Liege, Belgium; Email: albert.beckers@chu.ulg.ac.be)

(Antonio Ciccarelli and Hernan Valdes-Socin contributed equally to this study)

Abstract

Objective: Atypical forms of hyperthyroidism represent a diagnostic challenge for clinicians. Struma ovarii is an ovarian teratoma and constitutes a rare cause of ectopic thyroidal hormonal production. We describe a case of struma ovarii that combined two different sources of hyperthyroidism in the same patient and report genetic studies in order to contribute a better understanding of the autonomy and tumorigenesis of the struma ovarii.

Case report: A 73-year-old nulliparous woman presented a thyroid toxic adenoma that was successfully treated with 10 mCi radiiodine. Unexpectedly, a new onset of hyperthyroidism prompted us to look for a second etiology. A whole-body scan with 123I detected a pelvic hyperfixation suggesting struma ovarii, and a thyroid differentiated left ovarian teratoma 3 cm in size was surgically removed. We screened for mutations of thyroid-stimulating hormone receptor and Gs-α protein genes, as these mutations are common in thyroid adenomas. We did not identify any mutations. Androgen receptor study demonstrated a monoclonal status. Comparative genomic hybridization did not reveal any chromosomal abnormality. However, loss of heterozygosity analysis showed several structural abnormalities, compared with the majority of benign ovarian teratomas, which show a normal karyotype.

Conclusions: This is the first well-documented report of thyrotoxic struma ovarii revealed after treatment of a single thyroid toxic adenoma. We have shown in this case that struma ovarii originates from a single germ cell, and, albeit benign, this tumor presents several chromosomal abnormalities. Struma ovarii-induced hyperthyroidism is likely to be mediated by mechanisms different from those of the classical thyroid toxic adenoma.

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Introduction

Thyrotoxicosis is commonly caused by excessive production of thyroid hormones by the thyroid gland, as in Graves' disease, toxic adenoma or toxic multinodular goiter. These and other causes of hyperthyroidism are well associated with a high 24-h radioiodine uptake (RAI) in the thyroid bed. In contrast, some other causes of thyrotoxicosis are associated with a low 24-h RAI and are classified as atypical forms of hyperthyroidism. These latter represent a diagnostic puzzle for clinicians and may delay a proper management (1). As summarized in Table 1, atypical hyperthyroidism includes thyroid inflammation, iodine-induced hyperthyroidism and/or ectopic secretion of thyroid hormones.

Struma ovarii is a rare, mature ovarian teratoma. Since its first description in 1889, approximately 500 cases have been reported; only 5–15% are complicated by hyperthyroidism (2). Besides its rarity, the genotypic features of mature ovarian teratoma are disputed (3, 4), and the mechanisms underlying hyperthyroidism caused by struma ovarii are still largely unknown.

We report here a woman with atypical hyperthyroidism due to struma ovarii, co-existing with another source of typical hyperthyroidism. A genetic study of this teratoma was carried out to clarify its autonomy and tumorigenesis.

Case report

A 73-year-old, nulliparous, Caucasian woman had suffered a new onset of palpitations, tachycardia and a weight loss of 14 kg during the year before admission. Physical examination revealed a tender nodule 2 cm in size in the right thyroid lobe. Laboratory findings were
as follows: thyroid-stimulating hormone (TSH) < 0.01 mU/ml (0.1 – 4.0), free thyroxine (FT4) = 2.7 ng/ml (0.7 – 2) and free triiodothyronine (FT3) = 5 pg/ml (1.5 – 5.5), with thyroglobulin antibodies (AbTg), thyro-peroxidase antibodies (AbTPO), TSH-receptor antibodies (TRAb), thyroglobulin and calcitonin in the normal range. A 13 × 8 × 15 mm cervical nodule at ultrasonography showed intense 123I uptake. Hypothyroidism was achieved 3 months later after 10 mCi 131I treatment, and levothyroxine replacement was started progressively at a dose of 75 mg daily. However, the patient developed tachycardia and biochemically detected hyperthyroidism, prompting us to re-evaluate the thyroid profile without levothyroxine (TSH, 0.01 mU/ml, FT4 = 2.5 ng/ml, FT3 = 3 pg/ml, with AbTg, AbTPO and TRAb in the normal range) on the hypothesis of a residue of toxic adenoma, but a new RAI showed <1% uptake in the cervical tract. Since the patient had high consumption of marine salt, a low-salt diet was initiated. One month later, the urinary iodine level was normal, but hyperthyroidism persisted. Therefore, a whole-body scan with 123I was performed to detect atypical hyperthyroidism. Intense tracer activity was detected in the pelvis. An abdominal radiograph in the planar view demonstrated a pelvic soft tissue density containing calcification; ultrasonography of this region revealed a 12 × 8 cm mass, containing both solid and cystic components, located in the left side. No hypervascularization was found by Doppler ultrasound. Magnetic resonance imaging (MRI) of this lesion showed a bilocular mass in T1-weighted images; the solid part of the tumor was heterogeneous and hyperintense in T2-weighted images.

The mass, originating in the left ovary and firmly affixed to the rectosigmoid, was surgically removed. Total anexohysterectomy and partial resection of the sigmoid were performed (Fig. 1). Two weeks postoperatively, TSH had risen to 8 μU/ml and FT4 was 1.2 ng/ml. A control whole-body iodine scan showed a cervical residual uptake. Five years later, the patient was free of symptoms and was receiving replacement therapy with 87.5 μg of levothyroxine.

Pathologic studies
Histology revealed brown thyroid tissue associated with respiratory epithelium and bone tissue. Thyroid tissue measured 2.5 × 2.5 × 3 cm. Thyroid follicles varied in size with small and medium follicles consisted of cubic epithelium surrounding colloid material that stained positively with thyroglobulin but negatively with calcitonin. Fibroinflammatory cells and moderate cytoplastic vacuolization suggested secondary effects to radiiodine treatment.

Figure 1 A: Anexohysterectomy and partial resection of sigmoid. B: Magnification of struma ovary arising from left ovary. C: Histologic examination of struma ovarii (staining type haematoxylin-eosin; magnification 40 ×). The thyroid follicles (■) varied in size, with middle and small follicles consisted of cubic epithelium surrounding colloid material staining for thyroglobulin, but not for calcitonin. Fibroinflammatory cells (□) and moderate cytoplasmic vacuolization suggested secondary effects to radiiodine treatment.

Table 1 Causes of thyrotoxicosis with low radioiodine uptake in thyroid bed.

| Subacute thyroiditis |
| Struma ovarii |
| Factitious ingestion of thyroid hormone |
| Metastatic functional thyroid cancer |
| Iodine-induced hyperthyroidism |

Genetic investigations

1-TSH receptor and Gs-α subunit mutational studies
Genomic DNA and RNA were isolated from the patient’s lymphocytes and tumor. DNA and RNA were analyzed by PCR and RT-PCR respectively and submitted to direct sequencing by standard protocols, to detect either a known germline or somatic mutation of the receptor of TSH (TSHR) and Gs-α subunit genes. PCR products were purified and sequenced using the Big Dye cycle sequencing kit (Applied Biosystem, Foster City, CA, USA), and products of sequencing were analyzed on a 377 or 3100 Genetic Analyser (Applied Biosystem). Mutations were investigated using Factura and Sequence Navigator software.

TSH receptor gene The human TSHR gene is located on chromosome 14q31 (distal position). Most of its known activating mutations are located in exons 9 and 10 (from residue 430 to codon stop). Exon 9 was amplified using intronic primers for genomic DNA, as previously described (5), and exonic primers for cDNA
(primer direct \textit{5′}TGT AAA ACG ACG GCC AGT GCA TGC TGG ACG TGT CT3′, the underlined sequence corresponding to universal primer M13D; primer reverse \textit{5′}CAG TTT GTA GTG ACT AGT GAG GAG AAT AAG3′). Region 430-stop of exon 10 was amplified using three fragments, as previously described (6). No mutations were found, but a polymorphism, D727E, was identified in the heterozygote state in both the struma ovarii and the lymphocytes.

\textbf{Gs-α gene} The Gs-α gene is localized on chromosome 20q13 (distal position). Exons 8 and 9 were amplified by PCR using intronic primers for genomic DNA, as previously described (7), and exonic primers for cDNA (primer direct \textit{5′} ACT ATG TGC CGA GCG ATC AG 3′; primer reverse \textit{5′} CCC GGA TGA CCA TGT TGT AG 3′). No mutations could be detected.

\section*{2-Monoclonality studies}

The androgen receptor (AR) gene is mapped to chromosome Xq11.2 – q12 (centromeric position). The human AR gene assay (HUMARA) for the analysis of clonality in tissue from female patients examines the inactivation pattern of this gene on the X chromosome. This method relies on the length polymorphism of a human AR gene exon, which has restriction sites for methylase-sensitive enzymes (HpaII). Only methylated inactive alleles are preserved after HpaII digestion, and they are amplified by PCR. If the subject is heterozygote at the HUMARA after HpaII digestion, displaying two different alleles would indicate polyclonal origin, whereas the displaying of only one allele would indicate monoclonality. The allele pattern of AR was compared in the patient’s tumoral and lymphocytes DNA, as previously described (8). In our case, digestion by HpaII has not been necessary because, by HUMARA PCR amplification of undigested samples, we found a loss of heterozygosity for the AR gene in the tumor tissue, a clear sign of monoclonality (Fig. 2).

\section*{3-Comparative genomic hybridization (CGH)}

High molecular weight DNA was extracted from the fresh frozen tumor using standard methods. CGH was performed as previously described (9). Briefly, tumor DNA was labeled with FITC-12-dUTP (DuPont, Boston, MA, USA) by nick translation, and normal reference DNA was labeled with Spectrum Red (Vysis Inc., Downers Grove, IL, USA). Tumor and reference DNAs were mixed with unlabeled Cot-1 DNA (Gibco Invitrogen, Inc., Carlsbad, CA, USA), denatured and applied onto denatured metaphase slides of normal lymphocytes (Vysis Inc.). After hybridization at 37°C for 48 h, the slides were washed in 0.4 $\times$ SSC/0.3% NP-40 at 74°C for 2 min and in 2 $\times$ SSC/0.1% NP-40 at room temperature for 1 min. After air-drying, the slides were counterstained with 0.1 mg/ml 4,6-diamidino-2-phenylindole (DAPI) (Sigma) in an antifade solution (Vectashield, Vector Inc, Burlingame, CA, USA). A control hybridization was also performed using normal female DNA against normal male DNA.

Ten tri-color digital images (DAPI, Applied Spectral Imaging, Carlsbad, CA, USA; FITC and Spectrum Red fluorescence Vysis Inc., Downers Grove, IL, USA) were collected from each hybridization using a Zeiss Axioplan 2 (Carl Zeiss Jena GmbH, Jena, Germany) epifluorescence microscope and a Sensys (Photometrics, Tucson, AZ) camera.
AZ, USA) charge-coupled-device camera interfaced to an IPLab Spectrum 10 workstation (Signal Analytics Corporation, Vienna, VA, USA). Relative DNA sequence copy number changes were detected by analyzing the fluorescence intensities of tumor and normal DNAs along the length of all chromosomes in each metaphase spread. The absolute fluorescence intensities were normalized so that the average green-to-red ratio of all chromosomes in each metaphase was 1.0. The final results were plotted as a series of green-to-red ratio profiles and corresponding standard deviations (S.D.) for each human chromosome from p-telomere to q-telomere. At least 12 ratio profiles were averaged for each chromosome to reduce noise. Green-to-red ratios over 1.20 were considered as gains of genetic material, and ratios under 0.80 as losses. Heterochromatic regions, the short arm of the acrocentric chromosomes and chromosome Y were not included in the evaluation.

We did not find any chromosomal abnormalities in the tumor.

4-Loss of heterozygosity (LOH) analysis

We further analyzed the tumor and normal tissues for loss of heterozygosity (LOH). One, two or three microsatellite markers, preferably one from the short (p) arm and another from the long (q) arm, were chosen from ABI Prism Linkage Mapping Set v. 2 (Applied Biosystems, Foster City, CA, USA) for each chromosome: D1S2667, D1S498, D1S249, D2S168, D2S112, D3S1289, D3S1271, D4S391, D4S413, D5S630, D5S436, D6S470, D6S262, D7S513, D7S640, D8S550, D8S284, D9S286, D9S1776, D11S1338, D11S925, D12S99, D13S175, D13S170, D14S283, D14S68, D15S165, D15S127, D16S404, D16S515, D17S831, D17S798, D17S949, D18S452.

Figure 3 Electropherograms showing LOH in struma ovarii. LOH values were calculated from the height of the major peak of each allele (numbers in boxes). Microsatellite markers D7S640, D10S196, D10S1686, DXS990 and DXS1227 are located at 7q33, 10q11.23, 10q23.1, Xq21.32 and Xq27.2 respectively. N is the matched normal tissue and T is the tumor tissue.
D18S61, D19S209, D19S884, D19S571, D20S889, D20S196, D21S914 and D22S283. For chromosomes 10 and X, we used D10S547, D10S1653, D10S548, D10S197, D10S208, D10S196, D10S1652, D10S537, D10S1686, D10S185 and D10S192; and DXS1060, DXS8051, DXS987, DXS1226, DXS1214, DXS1068, DXS993, DXS991, DXS986, DXS990, DXS1106, DXS8055, DXS1001, DXS1047, DXS1227, DXS8043, DXS8091 and DXS1073, respectively. PCR was performed in a 7.5 μl reaction volume containing 0.17 μM each of HEX-labeled forward and unlabeled reverse primer (Invitrogen, Carlsbad, CA, USA), 4 mM MgCl2, 0.3 units AmpliTaq Gold polymerase and 1 X buffer II (Applied Biosystems), 250 μM dNTPs (Invitrogen), and 15 ng of genomic DNA. Amplification was performed in a DNA Engine Tetrad (MJ Research, Incline Village, NV, USA) with an initial denaturation of 95 °C for 10 min, followed by 10 cycles of 94 °C for 15 s, 55 °C for 15 s and 72 °C for 30 s; 20 cycles of 89 °C for 15 s, 55 °C for 15 s and 72 °C for 30 s; and a final extension at 72 °C for 10 min. A volume of 1 μl of each amplicon was added to 10 μl of Hi-Di formamide (Applied Biosystems) and 0.2 μl of ROX 400HD size standard (Applied Biosystems), denatured at 95 °C for 5 min and loaded into an ABI Prism 3700 Genetic Analyser (Applied Biosystems). LOH was assessed using Genescan v. 3.7 and Genotyper v. 3.7 software (Applied Biosystems). LOH was defined according to the following formula: LOH index = (T2/T1)/(N2/N1), where T was the tumor sample, N was the matched normal sample, and 1 and 2 were the intensities of smaller and larger alleles respectively. If the ratio was under 0.67 or over 1.3, the result was determined to be LOH. We found LOH at 4p15.2 (D4S391), 7p21.3–7q33 (D7S513, D7S640), 10q11.23 (D10S196), 10q23.1 (D10S186), 13q12.11 (D13S175), 14q11.2 (D14S283), 16q23.1 (D16S515), Xp22.1–Xq21.32 (DXS1226, DXS991, DXS986, DXS990) and Xq27.2 (DXS1227) (Fig. 3).

Discussion

Struma ovarii is a rare ovarian teratoma that consists mainly of thyroid tissue, biologically and microscopically identical to normal thyroid (10).

Benign and malignant, struma ovarii represent 0.3–1% of all ovarian neoplasms and 2–4% of all ovarian teratomas (1, 2). Most patients are diagnosed because of an asictic syndrome or a pelvic tumoral mass, whereas hyperthyroidism, with low radiodine uptake in the thyroid bed, is present in only about 5–15% of cases (2, 11, 12).

The case we report has the peculiarity of involving two different mechanisms of hyperthyroidism. The first mechanism is the occurrence of a toxic thyroid adenoma, according to ultrasonographic and scintigraphic evidence of a hot nodule, confirmed by transient hypothyroidism and a blunted cervical scintigraphy after radiodine treatment. Second, unexpectedly, we found thyrotoxicosis of new onset due to atypical hyperthyroidism, as demonstrated by a whole-body scan with 123I, clearly depicting a pelvic uptake suggestive of an ectopic source of thyroid hormone production. Pelvic ultrasonography and MRI detected a cystic mass 12 cm in diameter, and specific pathologic examination identified a restraint mass, 3 cm in size, containing thyroid tissue.

The synchronous presentation of thyroid toxic adenoma and struma ovarii is an extremely rare condition. Two previous cases of struma ovarii described in a series of 25 patients were diagnosed after thyroidectomy for Plummer’s disease (11). Nevertheless, these cases were diagnosed before 1970, and the contribution of teratomas to the thyrotoxic state seems difficult to assess retrospectively (13). The coexistence of struma ovarii and Graves’ disease has rarely been described (14).

As previously stated, struma ovarii has been found to be biochemically and histologically identical to the thyroid tissue. This case also suggests that hyperfunctioning struma ovarii can have a clinical presentation similar to that of thyroid toxic adenoma. The underlying patho-physiological mechanisms of autonomous thyroid nodules have been described only recently. Thyroid toxic adenomas were found to be the consequence of monoclonal proliferation (15). As thyroid cell proliferation is mainly regulated by TSH and the way of cAMP, mutations of TSHR and Gs-α protein were investigated in toxic adenomas (16). In a Belgian series of 33 thyroid toxic adenomas, TSHR and Gs-α mutations were found in 82% and 6% of hyperfunctioning tumors respectively (6), as confirmed by other studies (17, 18). Therefore, we have initially tested these hypotheses in this struma ovarii, but we were unable to identify any mutation of TSHR or Gs-α coding genes. This evidence suggests that (an)other mutation(s) as yet unknown or (an)other pathophysiological mechanism(s) could be implicated in the development of the hyperthyroidism caused by struma ovarii.

The struma ovarii is biochemically and histologically identical to thyroid tissue. However, it is a benign neoplasia that develops during embryogenesis. Is it monoclonal in analogy with other teratomas?

Theories on the origin of ovarian teratomas are controversial. Earlier studies showed that teratoma cells are homozygous for enzyme markers that are heterozygous in host cells (19), suggesting that teratomas arise from a single germ cell during the meiotic process. Later studies showed that distal loci of chromosome arms from benign ovarian teratomas are heterozygous for genetic recombination during the long prophase of meiosis I, whereas centromeric markers are homozygous, suggesting that teratomas might originate from a germ cell that had completed the first meiotic division but failed the second (20, 21). However, other studies
could not detect any homozygous genetic composition of teratomas while identifying heterozygous centromeric markers in a subset of the tumors, suggesting the possibility of either a postmeiotic or a premeiotic origin of these tumors (4, 22). Currently, the origin of teratomas remains debatable, but, generally, they appear to originate from a single germ cell after a defective meiosis, with a low incidence of numerical and/or structural chromosomal aberrations (3, 4).

As already mentioned, the struma ovarii is a rare ovarian teratoma; thus, genetic studies are very scant. In this case, we have had the opportunity to employ the most recent genetic techniques, probably for the first time.

We evaluated clonality using the AR. The struma ovarii cells displayed a LOH for the AR, as also confirmed by the LOH analysis of chromosome X (LOH at Xp22.1–Xq21.32), suggesting the monoclonality of this tumor and that it derives from a single germ cell.

Furthermore, we analyzed the cytogenetic features of the struma ovarii, using CGH and ‘conventional’ LOH analysis to detect possible chromosomal anomalies. CGH could not detect any chromosomal anomalies, but several chromosomal alterations were identified by using LOH, confirming the sensitivity of the LOH approach. Our results showed multiple loss of regions in chromosomes 4, 7, 10, 13, 14, 16 and X. Other cytogenetic analysis of mature ovarian teratomas showed a very low incidence of numerical and/or structural aberrations (5–7%) with no consistency or recurrence (4, 23, 24). Furthermore, some reports described chromosomal anomalies in mature ovarian teratomas that undergo malignant transformation (23, 24).

Some of the chromosomal anomalies described here have been previously reported in other cases of mature teratomas. However, some of them were also described in immature teratomas and in benign mature teratomas that undergo malignant transformation. Noumoff et al. observed a del(X)(q11) in the struma ovarii cells. As already mentioned, the struma ovarii is a rare ovarian teratoma, thus, genetic studies are very scant. In this case, we have had the opportunity to employ the most recent genetic techniques, probably for the first time.

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In conclusion, to our knowledge, this is the first report of thyrotoxic struma ovarii revealed after treatment of a single thyroid toxic adenoma, illustrating to clinicians how a tree may hide a forest. This peculiar case allowed us to gain further insights into the pathophysiology of this teratoma and of struma ovarii-induced hyperthyroidism. We have shown in this case that 1) struma ovarii, part of a unique group of tumors, originate from a single germ cell; 2) albeit benign, they present several chromosomal abnormalities (mostly deletions); and 3) struma ovarii-induced hyperthyroidism is likely to be mediated by mechanisms different from those of the classical thyroid toxic adenoma.

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