A Patient With Graves’ Disease And Thyroid-Associated Orbitopathy Undergoing Radioactive Iodine in Dr. Soetomo General Academic Hospital, Surabaya

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Abstract

Graves’ Disease (GD) is the most common etiology of thyrotoxicosis, followed by toxic multinodular goiter and toxic adenoma. GD can be managed with anti-thyroid drugs (ATDs), surgery, or radioactive iodine (RAI). Thyroid-associated orbitopathy (TAO) or Graves’ Ophthalmopathy (GO) affects 25%-50% patients with GD, and its presence usually dissuades clinicians to use RAI in treating hyperthyroidism. The presence of GO is a relative contraindication use of RAI in patients with GD, as RAI can worsen existing GO. Corticosteroid prophylaxis can be given to such patients to reduce likelihood of worsening of GO. However, patient with moderate to severe active GO is currently advised against undergoing RAI. Established guidelines recommend the use of corticosteroid prophylaxis in these patients. We reported a patients with GD and orbitopathy who was treated with RAI and was given steroid prophylaxis to prevent worsening of GO.

Introduction

Graves’ disease (GD) is the most common etiology of hyperthyroidism, which generally occurs in 40-60 years’ old patients. The underlying pathology of GD is the presence of TRab, an antibody against thyroid stimulating hormone (TSH), although the exact mechanism is still unknown. The antibody, an IgG, binds to the thyrotropine receptors and activates it, thus leads to hyperthyroidism. This activation also stimulates follicular hypertrophy and hyperplasia, which causes gland enlargement and also increase of thyroid hormone synthesis. A study in the UK found the incidence of GD was 24.8 per 100000 people/year. The prevalence of GD was reported around 0.5% of the general population, with female predominance (5-10 : 1). In Indonesia, based on the Basic Health Research 2013, the prevalence of hyperthyroidism was 0.4 case per 10000 people.3

Graves’ ophthalmopathy (GO) is an orbital inflammation associated with GD, with incidence rate 25%-50% of all patients with GD. Symptoms of GO are corneal irritation, periorcular edema, lid retraction, conjunctival erythema/chemosis, and extraocular muscles dysfunction. Some patients progress to more severe manifestation such as significant inflammation/congestion, excessive proptosis, and corneal ulceration or optic neuropathy, which can lead to permanent blindness.4,5

According to guideline published by the American Thyroid Association (ATA) on management of GD, either antithyroid drugs (ATDs), thyrodeomy, or thyroid ablation using radioactive iodine (RAI) can...
be used as one of therapy modalities. The choice is based on patient’s preference and clinical circumstance, since long-term quality of life between these therapies is not different. RAI is usually advised for patients who are unwilling to take lifelong medication or for those who cannot achieve euthyroidism with ATD. However, RAI is known to induce GO in GD patients with certain risk factors; therefore, GD patients with GO are usually given other choices of therapy. If RAI is given to GD patients with GO or those with risk factors, corticosteroid is indicated to prevent progression of GO after RAI.

We present a case of a female patient with history of GD and GO, which was treated with RAI following her preference over ATD, which she had been consuming for years. She was then given corticosteroids to prevent progression of her GO, and she responded well to medication.

**Case**

A 27-year-old female presented to the endocrinology outpatient with orbital pain since 1 month before presentation. She had chief a complaint of intermittent orbital pain since 5 years ago, but it was intensified since she treated with RAI 1 month ago. She also complained of painless lump on her right neck since 5 years ago. She denied having blurred vision nor double vision, but she felt protrusion of the right eye. She denied any fatigue, palpitations, diarrhea, excessive sweating, nor fever. She had been diagnosed with GD since 5 years ago, and had been taking thyrozol and propranolol. She also intermittently takes methylprednisolone since 5 years ago whenever she felt any orbital pain. Her symptoms of thyrotoxicosis were well controlled with ATD which she had been taking for the last 4 years, but she sought another choice of therapy. Initially offered for thyroid surgery, however, she opted for RAI which was done 1 month prior to presentation. Her medication at presentation was thyrozol 5 milligrams once daily. Physical examination showed an alert condition, blood pressure 120/70 mmHg, pulse 86 beats per minute, respiratory rate 16 times per minute, peripheral oxygen saturation 98% on room air, axillary temperature 36.3°C, and body weight of 55 kg. Head and neck examination revealed proptosis and lid retraction. Hertel exophthalmometer: right eye 16 mm, left eye 20 mm. Her CAS score was 1. Her ATD had been stopped for 1 month. Her other medications were continued.

The next month, she presented with much improved symptoms of thyrotoxicosis and orbital pain. Physical examination revealed right eye proptosis and lid retraction. Hertel exophthalmometer: right eye 16 mm, left eye 20 mm. Her CAS score was 1. Her thyroid function tests were as follows: FT4 1.05 ng/mL, TSH 0.194 ng/mL (Table 1). She was given propranolol 10 milligrams once daily, and her other medications were continued.

The following month, she came to the outpatient clinic with episodic palpitations, fatigue, and excessive swating. Her thyroid function tests were as follows: FT4 1.22 ng/mL, TSH 0.743 ng/mL (1 year prior to presentation) and 6.01 IU/L (4 years prior to presentation) (Table 1). Ultrasound of the neck showed diffuse enlargement of thyroid gland, with a hypechoic nodule measuring 2.3 x 1.6 cm in the right lobe without calcification and vascularization. Fine needle aspiration was done on the nodule, which showed no malignant cells. Her thyroid scan showed a nodular goitre with non-toxic laboratory result (Figure 2). The head MSCT showed 0.5 cm proptosis of the right eye and hypertrophy of extraocular muscle, conforms with a thyroid-associated orbitopathy (Figure 3). She was diagnosed with GD (post RAI), mildly active GO, and benign thyroid nodule. She was given thyrozol 2.5 milligrams once daily and methylprednisolone 8 milligrams thrice daily, and she was told to follow-up the next month.

A month after her first visit, she came to the outpatient clinic with chest discomfort and fatigue without orbital pain which accordance with a clinical manifestation of hypothyroidism. Her vital signs were within normal limit. Physical examination revealed proptosis and lid retraction of the right eye. Hertel exophthalmometer: right eye 20 mm, left eye 16 mm, and her CAS score was 1. Her ATD had been stopped for 1 month. Her thyroid function tests were as follows: FT4 1.05 ng/mL, TSH 4.815 ng/mL (Table 1). Propranolol had been stopped then changed with thyrozol 2.5 milligrams every two days alongside with tapered down of methylprednisolon dose.

The following month, she came to the outpatient clinic with chest discomfort and fatigue without orbital pain which accordance with a clinical manifestation of hypothyroidism. Her vital signs were within normal limit. Physical examination revealed proptosis and lid retraction of the right eye. Hertel exophthalmometer: right eye 20 mm, left eye 16 mm, and her CAS score was 1. Her ATD had been stopped for 1 month. Her thyroid function tests were as follows: FT4 1.03 ng/mL and TSH 11.197 ng/mL (Table 1). She was diagnosed with GD post RAI and subclinical hypothyroidism, then was given levothyroxine 50 microgram once daily. Her subsequent visits were symptom-free and her thyroid functions were within normal range under the daily levothyroxine, and she was scheduled for eyelid surgery to correct the lid retraction. Table 1 summarizes her thyroid function throughout monthly visits.

**Table 1. Patient’s thyroid function throughout her monthly visits**

<table>
<thead>
<tr>
<th>Date</th>
<th>FT4 (ng/dL)</th>
<th>TSH (ng/dL)</th>
<th>TRAb (U/l)</th>
</tr>
</thead>
<tbody>
<tr>
<td>29/11/14</td>
<td>3.05</td>
<td>&lt;0.01</td>
<td>6.01</td>
</tr>
<tr>
<td>12/9/17</td>
<td>1.12</td>
<td>1.22</td>
<td>2.82</td>
</tr>
<tr>
<td>28/4/18</td>
<td>1.05</td>
<td>0.194</td>
<td></td>
</tr>
<tr>
<td>30/6/18</td>
<td>1.03</td>
<td>4.815</td>
<td></td>
</tr>
<tr>
<td>27/7/18</td>
<td>0.82</td>
<td>11.197</td>
<td>3.3</td>
</tr>
</tbody>
</table>

**Note:** The data in the table is based on patient’s thyroid function throughout her monthly visits. The values are given in micrograms per milliliter (µg/mL) for TSH, and in International Units per liter (IU/L) for TRAb. The values are rounded to two decimal places for simplicity.
Figure 1. Clinical appearance of the patient during her first visit to endocrinology outpatient. Note the ptosis of left upper eyelid.

Figure 2. Patient’s Head CT (Orbital-Focused). There is marked hypertrophy of medial and lateral right rectus muscles (red arrows), which pushes ocular bulb anteriorly, which causes proptosis +/- 0.5 cm. Radiologist’s remark: Thyroid-Associated Orbitopathy

Discussion
Graves’ Disease is an autoimmune disease mediated by autoantibody, with thyrotoxicosis as its cardinal manifestation. GD is diagnosed by the presence of diffuse enlargement of the thyroid gland, moderate to severe hyperthyroidism, and orbitopathy. When these signs and symptoms are present, the diagnosis of GD is usually straightforward. In the absence of these features, elevated TRAb and characteristic thyroid scan (RAIU) uptake can aid in the diagnosis. Although GD is commonly known to involve thyroid gland, the other organ systems can be affected as well, either as a part of the autoimmune process or as complications of thyrotoxicosis. Systemic autoimmune involvement of GD includes eye (GO) and skin (Graves’ dermatopathy), as tissues in these organs possess receptor to TRAb.

Graves’ ophthalmopathy (GO) is define as an inflammation of orbita associated with GD, however, not all GD patients will develop GO. GD patients with established GO or suspicion of GO must be evaluated by ophthalmologist. In terms of severity, GO can be classified into mild, moderate, and severe, according to NOSPECS (No physical signs or symptoms, Only signs, no symptoms, Soft tissue involvement, Proptosis, Extraocular muscle involvement, Corneal involvement, Sight loss) criteria or EUGOGO (EUropean Groups On Graves’ Orbitopathy) criteria. In terms of disease activity, GO can be divided into active or inactive based on CAS (Clinical Activity Score) scoring system. This classification will guide clinicians to determine the best course of action in treating the ophthalmopathy, with overall goal is to avoid visual-threatening disease and to achieve minimal disease activity.

In GD patients with GO, therapy of GD with RAI should be done with consideration, because RAI can worsen the existing ophthalmopathy, especially when the patient develops symptomatic hypothyroidism after RAI. Guideline from ATA recommends systemic corticosteroids be given to GD patients with GO under RAI treatment within the setting of mild and active ophthalmopathy, with/without risk factors of worsening of GO. RAI is not recommended for GD patients with moderate to severe GO.

A RCT recommends giving 0.4-0.5 mg/kgBW/day of prednisone as prophylaxis for GD patients with ophthalmopathy treated with RAI, which can be started as early as 1-3 days after RAI. The dose should be given for 1 month, followed by 2 months of gradual tapering. In the other hand, a retrospective cohort study reported that lower dose and shorter duration of steroid (prednisol 0.2 mg/kgBW/day for 6 weeks) has a similar efficacy in terms of preventing new onset GO and worsening of ophthalmopathy in patients with preexisting GO. Currently, most guidelines recommend administering 30 mg of prednisone daily followed by tapering down, with total duration of 6-8 weeks. After achieving a low disease activity, any residual anatomical disruption of the orbita can be corrected by surgery.

Conclusion
In summary, RAI can be chosen as therapy for patients with GD presenting with mild and active GO, as per 2016 guideline from American Thyroid Association. Prophylactic steroid is given after RAI to prevent worsening of GO. Post-RAI care should be given to prevent hypothyroidism after RAI, which can also worsen GO. This patient was treated with RAI over surgery after taking ATD for years despite her ophthalmopathy, and was given prophylactic steroid. She achieved hypothyroidism several months after RAI, and did not show any deterioration of ophthalmopathy.

Conflict of Interest
The author stated there is no conflict of interest

References


