Thyroid Echography-induced Thyroid Storm and Exacerbation of Acute Heart Failure

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Abstract

Hyperthyroidism and thyroid storm affect cardiac circulation in some conditions. Several factors including trauma can induce thyroid storms. We herein describe the case of a 57-year-old woman who experienced a thyroid storm and exacerbation of acute heart failure on thyroid echography. She initially demonstrated a good clinical course after medical rate control for atrial fibrillation; however, thyroid echography for evaluating hyperthyroidism led to a thyroid storm and she collapsed. A multidisciplinary approach stabilized her thyroid hormone levels and hemodynamics. Thus, the medical staff should be prepared for a deterioration in the patient’s condition during thyroid echography in heart failure patients with hyperthyroidism.

Key words: hyperthyroidism, acute heart failure, thyroid echography, thyroid storm


Introduction

Hyperthyroidism and thyroid storm may impact cardiac circulation in some conditions. Moreover, hyperdynamic states can cause high-output heart failure, while tachycardia causes tachycardia-mediated cardiomyopathy. Therefore, if physicians treat a heart failure patient with hyperthyroidism, they must perform thyroid therapy preceding or simultaneously with heart failure therapy.

A thyroid storm is relatively rare; it is reported in 0.20 people per 100,000 in the Japanese population annually and occurs in 0.22% of thyrotoxic patients (1). It has a mortality rate of over 10% in Japan (1), and it is induced by an acute event such as trauma, thyroid or non-thyroid surgery, infection, acute iodine load, or drug provocation. We herein describe a case where mechanical stimulation by thyroid echography induced a thyroid storm, which thus resulted in an exacerbation of acute heart failure.

Case Report

A 57-year-old woman working in the cleaning department presented with worsening dyspnea and general fatigue. Her medical and family history was unremarkable. She had never experienced pathological weight loss, sweating, tremors, diarrhea, nausea, or vomiting. She was alert and her blood pressure was 120/86 mmHg, heart rate was 206/min, body temperature was 36.1°C, respiratory rate was 30/min, and saturation was 96% (room air). Coarse crackles in all lung fields, bowel sound acceleration, and pretibial edema were detected, however, no goiter or exophthalmic or finger tremors were observed. The electrocardiogram showed an atrial fibrillation rhythm with a rapid ventricular response (heart rate 223/min), ST depression in leads V4-6, high voltage in leads V5-6, and inverted T in leads II, III, and the augmented vector foot lead (Fig. 1, left panel). The chest radiograph indicated pulmonary edema, enlargement of the cardiac silhouette and a cardiothoracic ratio of 64% (Fig. 1, right panel). Transthoracic echocardiography showed an almost completely preserved ejection fraction, enlargement of all cardiac chambers, moderate mitral regurgitation, moderate tricuspid regurgitation, and an inferior vena cava measuring 30 mm in diameter. Laboratory data showed the following values: 1.3 mg/dL of total bilirubin, 48 U/L of aspartate aminotransferase, 29 U/L of alanine aminotransferase, 0.34 mg/dL of creatinine, 69 mg/dL of total cholesterol, and 262.4 pg/ml of brain natriuretic peptide in the plasma.
According to these findings, we diagnosed the patient with acute decompensated heart failure. The suspected etiologies were as follows: rapid atrial fibrillation, tachycardia-mediated cardiomyopathy, diastolic dysfunction, and high-output heart failure. We first administered medical rate control using intravenous diltiazem (100 mg/day), oral bisoprolol (2.5 mg/day), and anticoagulation with heparin. This strategy was very successful and decreased the heart rate from 200 to 80/min, relieved the symptoms, and resulted in a body weight loss of 1.5 kg within 20 hours.

We confirmed that the patient had hyperthyroidism (thyroid stimulating hormone <0.01 IU/ml, free thyroxine 4.38 ng/dL, and free triiodothyronine 11.68 pg/ml; thyroid hormone receptor antibody 21.0%; thyroglobulin 163.9 ng/mL; anti-thyroid peroxidase antibody 40 IU/mL; anti-thyroglobulin antibody <10 IU/mL) on day 2. We subsequently performed thyroid echography to investigate hyperthyroidism, which revealed a heterogeneous thyroid of normal size and normal vasculature (Fig. 2).

The patient suddenly complained of nausea, thirst, and acute dyspnea. Thereafter, she rapidly lost her orientation and collapsed an hour after undergoing thyroid echography. Her body temperature was 38.4°C. The chest radiograph showed a butterfly shadow and laboratory data showed hyperlactatemia (169.5 mg/dL), hyponatremia (130 mEq/L), hypoglycemia (40 mg/dL), and hyperbilirubinemia (5.3 mg/dL total bilirubin and 3.4 mg/dL direct bilirubin), despite catecholamine support. We suspected a thyroid storm. An emergency administration of 200 mg intravenous hydrocortisone and 15 mg thiamazole was initiated with catecholamine support. The first dose immediately relieved the patient’s condition, thus we administered this dosage daily and tapered it gradually thereafter. After confirming the iodine uptake using technetium-99m pertechnetate scintigraphy (Fig. 3) on day 3, we diagnosed the patient with Basedow’s disease and the administration of inorganic iodine was also initiated. These multidisciplinary approaches stabilized the patient’s thyroid hormone levels (Table) and hemodynamics drastically (Fig. 4). After this acute phase, she recovered with an uneventful course and was discharged on day 22.

The patient’s thyroid and heart rate are currently being controlled in the outpatient clinic with 5 mg of thiamazole and 5 mg of bisoprolol. She has not experienced heart failure for approximately 2 years after the first admission. We obtained the patient’s informed consent to publish this case report.

Discussion

This case is the first published report of thyroid echography-induced thyroid storm and presents two important findings. Firstly, mechanical stress to the thyroid may lead to the exacerbation of acute heart failure in patients with hyperthyroidism. Secondly, the differential diagnosis for the etiology of acute heart failure is important to select the appropriate treatment to rescue the patient.

Mechanical stress to the thyroid may lead to the exacerbation of acute heart failure in patients with hyperthyroidism. Thyroid storms can have many triggers including trauma, thyroid or non-thyroid surgery, infection, acute iodine load, or drug provocation. The trigger in our case may be mechanical compression stress during thyroid echography. Some studies reported that suicide attempts by hanging (2), choking by assailants (3), or blunt trauma (4) resulted in a thyroid storm. In our institution, we observed ventricular fibrillation during thyroid echography in a heart failure pa-
The differential diagnosis of acute heart failure is important for selecting the appropriate treatment to rescue a patient. We treated the present patient by mainly targeting tachycardia on admission, which was successful. However, rate control therapy and catecholamine were not effective after thyroid echography. We immediately changed the main target etiology of heart failure to a thyroid storm from tachycardia. Although we used a relatively low dose of the treatment in this case, the standard therapy for a thyroid storm can include 20 mg of methimazole every 4 to 6 hours, 200 mg propylthiouracil every 4 hours, 10 drops of Lugol’s solution every 6 hours, 100 mg of hydrocortisone every 8 hours, or 4 g cholestyramine four times daily (7). If we had continued the heart rate-oriented therapy, we might have lost our patient.

Because our institution does not perform laboratory tests...
for thyroid measurement within the hospital setting, we obtained a delayed result on day 2. However, we were aware that heart failure and atrial fibrillation with a very rapid ventricular rate are often attributable to hyperthyroidism. Atrial fibrillation, especially in the presence of pre-existent heart disease, can result in clinical heart failure. Such heart failure may be due to an associated rapid ventricular response, which when sustained can lead to tachycardia-mediated cardiomyopathy. The loss of atrial contractile function and decreased diastolic filling time due to tachycardia may cause increased filling pressures, further contributing to this cardiomyopathy (8).

In conclusion, clinicians should be prepared for a possible deterioration in the patient’s condition during thyroid echography for heart failure patients with hyperthyroidism.

The authors state that they have no Conflict of Interest (COI).

References


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Table. Thyroid Hormone Levels during the Clinical Course.

<table>
<thead>
<tr>
<th>Day Post</th>
<th>TSH (NR: 0.35-4.94 μIU/mL)</th>
<th>Free Thyroxine (NR: 0.7-1.48 ng/dL)</th>
<th>Free Triiodothyronine (NR: 1.71-3.71 ng/dL)</th>
</tr>
</thead>
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<tr>
<td>2</td>
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<td>2.66</td>
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<tr>
<td>12</td>
<td>&lt; 0.01</td>
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<td>2.93</td>
</tr>
</tbody>
</table>

TSH: thyroid stimulating hormone, NR: normal range

Figure 4. Clinical course. DOA: dopamine, HDC: hydrocortisone, HR: heart rate, KI: potassium iodide, MMI: methimazole, NAD: noradrenaline, SBP: systolic blood pressure