Acute coronary syndrome with severe atherosclerotic and hyperthyroidism: A case report

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Abstract

BACKGROUND
Acute coronary syndrome (ACS) encompasses a spectrum of cardiovascular emergencies arising from the obstruction of coronary artery blood flow and acute myocardial ischemia. Recent studies have revealed that thyroid function is closely related to ACS. However, only a few reports of thyrotoxicosis-induced ACS with severe atherosclerosis have been reported.

CASE SUMMARY
A 33-year-old man, who had a history of hyperthyroidism without taking any antithyroid drugs and no history of coronary heart disease, experienced neck pain with occasional heart palpitations starting 3 mo prior that were aggravated after an activity. As the symptoms worsened at 21 d prior, he went to a hospital for treatment. The electrocardiogram examination showed a multilead ST segment elevation and pathological Q waves. Based on these findings and his symptoms, the patient was diagnosed with a suspected myocardial infarction and transferred to our hospital on July 2, 2020. He was diagnosed with a rare case of ACS due to coronary artery atherosclerosis in the anterior descending artery complicated by hyperthyroidism. A paclitaxel-coated drug balloon was used for treatment to avoid the use of metal stents, thus reducing the time of antiplatelet therapy and facilitating the continued treatment of hyperthyroidism. The 9-mo follow-up showed favorable results.

CONCLUSION
This case highlights that atherosclerosis is a cause of ACS that cannot be ignored even in a patient with hyperthyroidism.
Acute coronary syndrome (ACS) encompasses a spectrum of cardiovascular emergencies arising from the obstruction of coronary artery blood flow and acute myocardial ischemia. Patients with ACS frequently have a poor prognosis, and ACS is a major health and economic burden[1,2]. The symptoms of ACS arise from the functional destruction of the circulatory system. However, the pathogenic factors of ACS often arise from systems outside the circulatory system, and diseases related to endocrine system dysfunction, such as diabetes and hyperthyroidism[3], have an indispensable role to play. Such factors increase the difficulty in clinical treatment and improvement of patient prognosis. Therefore, in recent years, coronary heart disease (CHD) secondary to hyperthyroidism has gradually received considerable attention[4]. We describe a rare case of ACS due to coronary artery atherosclerosis in the anterior descending artery complicated by hyperthyroidism in a 33-year-old man. We also present relevant literature and discuss the interaction mechanism between complications and CHD to achieve a suitable treatment plan.

**CASE PRESENTATION**

**Chief complaints**

ACS encompasses a spectrum of cardiovascular emergencies arising from the obstruction of coronary artery blood flow and acute myocardial ischemia. Patients with ACS frequently have a poor prognosis, and ACS is a major health and economic burden[1,2]. The symptoms of ACS arise from the functional destruction of the circulatory system. However, the pathogenic factors of ACS often arise from systems outside the circulatory system, and diseases related to endocrine system dysfunction, such as diabetes and hyperthyroidism[3], have an indispensable role to play. Such factors increase the difficulty in clinical treatment and improvement of patient prognosis. Therefore, in recent years, CHD secondary to hyperthyroidism has gradually received considerable attention[4]. We describe a rare case of ACS due to coronary artery atherosclerosis in the anterior descending artery complicated by hyperthyroidism in a 33-year-old man. We also present relevant literature and discuss the interaction mechanism between complications and CHD to achieve a suitable treatment plan.
History of present illness
Three months ago, the patient experienced neck pain with occasional heart palpi-
tations that were aggravated after an activity. As the symptoms worsened 21 d prior,
he went to a hospital for treatment. The electrocardiogram examination showed a
multilead ST segment elevation and pathological Q waves. Based on the findings and
his symptoms, the patient was diagnosed with a suspected myocardial infarction.

History of past illness
The patient had a history of hyperthyroidism for 5 mo, without taking any antithyroid
drugs. There was no history of CHD.

Personal and family history
The patient smoked approximately 12 cigarettes a day for 10 years. He denied a family
history of related disease.

Physical examination
After hospitalization, the results of the diagnosis-related examinations were as follows:
body temperature, 36.6 ºC; breathing, 20 breaths/min; blood pressure, 120/80 mmHg;
and heart rate, 110 beats/min. The patient was of sound mind and had a slightly
enlarged thyroid with ocular signs. Since the onset of the disease, he has lost 6 kg of
weight.

Laboratory examinations
The electrocardiogram results were as follows: sinus rhythm, V1-5 ST segment
elevation 0.1-0.4 mv and pathological Q waves (Figure 1).

Although the high-sensitivity troponin T (commonly referred to as TNT-Hs) test
result was 29.2 pg/mL on admission and 17.1 pg/mL on day 5 of admission (reference
range: 0-0.04 pg/mL), the myocardial enzyme test did not show abnormal results. We
also tested the patient’s thyroid function (Table 1 and 2), blood lipids (Table 3), and
cogulation function (Table 4). The thyroid function test showed high levels of free
triiodothyronine (commonly referred to as FT3), free thyroxine (commonly referred to
as FT4) and thyroid-stimulating hormone receptor antibody (commonly referred to as
TRAb), and low level of third-generation thyroid-stimulating hormone (commonly
referred to as TSH-3GEN).

Imaging examinations
Cardiac color Doppler ultrasound showed uncoordinated left ventricular wall motion
and weakened interventricular septal motion. Thyroid color Doppler ultrasound
showed that the bilateral thyroid glands were diffusely enlarged with rich color flow,
and bilateral cervical lymph nodes were visible.

FINAL DIAGNOSIS
The final diagnosis of the presented case was ACS due to coronary artery athero-
sclerosis in the anterior descending artery complicated by hyperthyroidism.

TREATMENT
Preoperative intravascular ultrasound (IVUS) examination showed plaque formation
in the middle of the anterior descending branch (Figure 2) and severe lesions in the
proximal segment. The lesions were rich in lipids and fibrous plaques. The minimum
lumen cross-sectional area (referred to as CSA) was 2.03 mm² (Figure 3), and
percutaneous coronary intervention was performed. After the anterior descending
branch guide wire was passed, a 2.0 × 20 Abbott Balloon (MINI TREK) 10 atm was
administered to predilate the proximal lesion, and a 3.5 × 10 cutting balloon
(FlxtoneTM Cutting Balloon™) was cut twice at 8 atm. Then, a 3.5 × 20 drug-coated
balloon (SeQuent; B. Braun Melsungen AG) was expanded at 6 atm for 60 s. Finally,
the original narrowest lumen CSA increased to 5.58 mm² by IVUS examination. There
was no local dissection or hematoma formation and coronary angiography showed
significant relief of the stenosis (Figure 4 and 5). The patient’s condition improved 4 d
after the interventional therapy, and he was discharged. The foregoing oral medication
### Table 1 Thyroid function test results from time of admission

| Item         | Result | Reference range | Unit   |
|--------------|--------|-----------------|--------|
| FT3          | > 30.00| 1.71-3.71       | pg/mL  |
| FT4          | 3.59   | 0.7-1.48        | ng/dL  |
| TSH-3GEN     | 0.0017 | 0.4700-4.6400   | IU/mL  |
| A-TPO        | 2.40   | 0-5.61          | IU/mL  |
| A-TG         | 0.84   | 0-4.11          | IU/mL  |
| TRAb         | 12.56  | < 1.75          | IU/L   |

A-TPO: Thyroid peroxidase; A-TG: Thyroglobulin; FT3: Free triiodothyronine; FT4: Free thyroxine; TRAb: Thyroid-stimulating hormone receptor antibody; TSH-3GEN: Third-generation thyroid-stimulating hormone.

### Table 2 Thyroid function test results from 1 wk after anti-hyperthyroidism treatment

| Item         | Result | Reference range | Unit   |
|--------------|--------|-----------------|--------|
| FT3          | 11.85  | 1.71-3.71       | pg/mL  |
| FT4          | 2.85   | 0.70-1.48       | ng/dL  |
| TSH-3GEN     | 0.0016 | 0.4700-4.6400   | IU/mL  |

FT3: Free triiodothyronine; FT4: Thyroxine; TSH-3GEN: Third-generation thyroid-stimulating hormone.

### Table 3 Blood lipid test results

| Item | Result | Reference range | Unit   |
|------|--------|-----------------|--------|
| CHOL | 3.56   | 2.80-5.17       | mmol/L |
| TG   | 1.16   | 0.56-1.70       | mmol/L |
| HDL  | 0.81   | 0.96-1.15       | mmol/L |
| LDL  | 2.63   | 0-3.10          | mmol/L |

CHOL: Cholesterol; HDL: High-density lipoprotein; LDL: Low-density lipoprotein; TG: Triglycerides.

### Table 4 Coagulation test results

| Item       | Result | Reference range | Unit |
|------------|--------|-----------------|------|
| PT-sec     | 13.0   | 11-14           | s    |
| PT-INR     | 1.09   | 0.8-1.2         | N/A  |
| APTT       | 47.7   | 27-45           | s    |
| FIB        | 2.83   | 2-4             | g/L  |
| TT         | 18.6   | 0-20            | s    |
| D-Dimer    | 0.52   | 0-1             | mg/L |

APTT: Activated partial thromboplastin time; FIB: Fibrinogen; N/A: Not applicable; PT-sec: Prothrombin time in seconds; PT-INR: Prothrombin time international normalized ratio; TT: Thrombin time.

regimen was continued after discharge.
Figure 1 Electrocardiogram examination. A: On admission; B: Day after admission.

Figure 2 Presence of plaque in the middle of the anterior descending coronary artery. A: Plaque burden was 50%; B: Plaque thickness was 1 mm.

OUTCOME AND FOLLOW-UP
The 9 mo follow-up showed that the patient was in good condition. On March 11, 2021, the cardiac ultrasound showed that left ventricular wall movement was roughly coordinated, and systolic function was normal.
DISCUSSION

The thyroid hormone has a significant effect on the metabolism of sugar, fat, and protein[5] and acts on all tissue cells. Recent studies have revealed that thyroid hormone widely affects the physiological and pathological processes of the cardiovascular system[4] and is associated with cardiomyocyte injury[6], thrombus burden[7], coronary artery spasm[8], and coronary atherosclerosis[9,10]. The effects of the thyroid hormone on the cardiovascular system include increased resting heart rate, left ventricular contractility, blood pressure (volume) and decreased systemic vascular resistance.

Thyroid function is closely related to ACS. Coronary atherosclerosis, one of the most important pathogenic factors leading to ACS, is a long-lasting and continuously evolving disease[11]. Patients with thyrotoxicosis-induced ACS are rare, and almost all reported cases have been associated with Graves’ disease. Coronary angiography usually shows zero disease, and coronary artery spasm occupies a large proportion of data[8,12,13]. However, in this case ACS was accompanied by severe atherosclerosis.

In our patient, the main manifestation was neck pain, which is atypical. The patient was diagnosed with hyperthyroidism because he had a goiter and showed ocular protrusion symptoms. ACS was not diagnosed until the electrocardiogram and TNT-Hs test were completed. Patients with hyperthyroidism often exhibit a high metabolic state[14], and the incidence of diabetes and dyslipidemia in such patients is lower than that in ordinary patients, indicating that the conditions for atherosclerosis are lacking and patients are less likely to have CHD. It is important to understand the factors that caused atherosclerotic plaque in this young male patient with hyperthyroidism. There were several risk factors, including his former work as a courier, a fatty diet, smoking, poor sleep, irregular lifestyle, and stress. During the first half of 2020, the patient had...
to stay home for several months because of the COVID-19 epidemic, resulting in a lack of physical exercise.

It is possible that severe atherosclerotic plaques already existed in the coronary arteries before the onset of hyperthyroidism, and the environmental risk factors promoted the development of coronary atherosclerosis. It is also possible that newly developed hyperthyroidism induced coronary artery spasm or accelerated the progression of atherosclerosis and subsequent plaque disruption or erosion that led to ACS. Given that IVUS did not accurately reflect the composition of the plaque surface or image the microstructure, optical coherence tomography detection provided a better understanding of the mechanism of ACS onset.

We managed the disease using drug-coated balloon technology. During the treatment, we used the IVUS examination to evaluate the treatment effect of the cutting balloon + drug balloon, which can limit the use of iodine-containing contrast media and reduce the effect of iodine on patients with hyperthyroidism. Patients with hyperthyroidism need to take antithyroid drugs, such as thiourea, for an extended period, which may lead to complications, such as neutropenia. If metal stents were used, the patients may be unable to withstand long-term double antiplatelet therapy because of neutropenia or bleeding. Therefore, we chose the paclitaxel-coated drug balloon as an implant-free interventional therapy to avoid the use of stents, reduce the time of antiplatelet therapy, and facilitate the continued treatment of the patient with subsequent hyperthyroidism.

CONCLUSION

ACS with hyperthyroidism is easy to miss clinically. In addition to coronary spasms, the mechanism of coronary atherosclerosis is a cause of ACS that cannot be ignored. For young patients, the dangers of smoking, a fatty diet, and sedentary lifestyle should be emphasized. In terms of clinical treatment, intensive drug therapy and implant-free interventional therapy are better options for patients with ACS and hyperthyroidism.

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