Shock Thyroid in a Patient with Septic Shock: 
A Case Report and Literature Review

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Shock thyroid is a rare manifestation of the CT hypotension complex and can be diagnosed when thyroid and perithyroidal edemas are observed on CT during the onset of shock. Shock thyroid can be a useful CT sign for decompensated shock. This condition is reversible and recovers rapidly with adequate treatment. We present the case of an 84-year-old female with septic shock, exhibiting CT features consistent with a shock thyroid. We also reviewed the clinical and radiological findings reported in the literature. The present case emphasizes that shock thyroid can be an early indicator of impending hemodynamic instability and has important prognostic and therapeutic implications.

Index terms Thyroid; Septic Shock; Hypotension; Multidetector Computed Tomography

INTRODUCTION

Shock thyroid is an uncommon part of the CT hypotension complex (1, 2). Shock thyroid manifests as thyroid and perithyroidal edema, which is incidentally noted in CT of patients with shock conditions (1, 2). To our knowledge, only seven cases have been reported in the literature (1, 2), and because this disease is largely unknown to radiologists and clinicians, it is frequently underdiagnosed (2). The relationship between shock thyroids and poor prognosis remains unclear. However, shock itself is a life-threatening condition that can lead to multiple organ failure. Prompt diagnosis of shock is very imp-
important for proper treatment of the underlying condition. Thus, shock thyroid is a helpful imaging indicator to identify inadequate perfusion in patients with shock (1, 2).

Here, we present the case of a female with septic shock secondary to *Staphylococcus aureus* bacteremia of unknown source, and conduct a comprehensive review of the literature. Our study emphasizes that shock thyroid is not related to thyroid injury or primary disease, but is related to perfusion alterations. Shock thyroid must be distinguished from other pathologies of the thyroid gland to avoid unnecessary tests and to establish an appropriate treatment plan.

**CASE REPORT**

An 84-year-old incapacitated female with underlying Alzheimer’s dementia presented to our emergency department with a chief complaint of fever. The patient had been admitted to our hospital 39 days previously, due to pulmonary thromboembolism and deep vein thrombosis. Her initial vital signs were as follows: blood pressure, 87/44 mm Hg; heart rate, 105 bpm; and body temperature, 38.2°. Laboratory analyses showed leukocytosis, elevated levels of C-reactive protein, and serum lactate level of 3.57 mmol/L. Her clinical presentation was characterized by septic shock. Initial blood culture revealed methicillin-resistant *Staphylococcus aureus* bacteremia.

Approximately 4 hours after arriving at the emergency department, the patient underwent contrast-enhanced chest and abdominal CT to identify the infection focus. Chest CT revealed multiple peripheral nodules in both lungs, indicating septic embolism. Additionally, CT demonstrated mild swelling and heterogeneous enhancement of the thyroid gland with perithyroidal fluid collection (Fig. 1A, B). CT scan obtained at the previous admission due to pulmonary thromboembolism did not show abnormalities in the thyroid gland, except a probable benign hypo-attenuating nodule in the right thyroid gland. The radiologic findings in the thyroid gland were interpreted as shock thyroid. However, there were no accompanying signs of the CT hypotension complex, such as shock bowel, collapse of the inferior vena cava.

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**Fig. 1.** An 84-year-old female with septic shock and shock thyroid.

A, B. Axial and coronal reconstruction images of the contrast-enhanced chest CT obtained at the emergency department demonstrate diffuse parenchymal swelling and heterogeneous enhancement of the thyroid gland. Perithyroidal fluid collection is also observed.

C. Follow-up contrast-enhanced CT image obtained 22 hours after (A) and (B) showing remarkable improvement. The thyroid gland demonstrates a normal size and homogenous enhancement with near-complete resolution of the perithyroidal fluid collection.
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The patient was admitted to the intensive care unit. The source of methicillin-resistant Staphylococcus aureus bacteremia was uncertain. Glycopeptide antibiotic and fluid resuscitation, with diluted norepinephrine, were administered through a central venous catheter. The patient became hemodynamically stable after resuscitation therapy. The attending physician performed follow-up neck CT and thyroid function tests to differentiate the primary thyroid gland disease. Neck CT taken 22 hours after the initial chest CT demonstrated normalization of size and enhancement pattern of the thyroid gland (Fig. 1C). Perithyroidal fluid collection was mostly resolved. The thyroid function test on the same day showed normal concentrations of serum free thyroxine (T4, 1.23 ng/dL) and serum thyroid-stimulating hormone (0.53 mIU/L), and low concentration of serum triiodothyronine (T3, 58.5 ng/dL). After 4 days in the intensive care unit, the patient was moved to the general ward.

This study was approved by the Institutional Review Board of our institution and the requirement for informed consent was waived (IRB No. 2020-07-033).

DISCUSSION

The CT hypotension complex refers to a spectrum of CT findings with severe hypotension (3, 4). It is mainly present in abdominal organs, but multiple organs can exhibit various atypical features (3, 4). In 2006, Brochert and Rafoth (1) reported thoracic CT findings of three traumatized patients with hypovolemic shock, showing heterogeneous thyroid contrast enhancement and fluid surrounding the thyroid. They described these CT findings as “shock thyroid” and considered as part of the CT hypotension complex rather than direct traumatic injury (1).

The exact incidence of shock thyroid remains unknown, and to our knowledge, only seven cases have been reported in the literature (Table 1) (1, 2). It is thus certain that shock thyroid is a relatively rare manifestation of the CT hypotension complex. In a previous study involving traumatized or non-traumatized 41 patients with shock bowel on abdominal CT, no shock thyroid cases were found (4). However, because of the low awareness about this disorder, it might be frequently misdiagnosed, as the findings might be attributed to other conditions, such as thyroiditis or traumatic injury (2).

Although the CT hypotension complex was reported for the first time in traumatic patients with hemodynamic instability (5), it may also occur in multiple other hypovolemic conditions, including neurogenic shock from head or spinal injury, blood loss, sepsis, and cardiac arrest (4). Similarly, shock thyroid can be present in patients with hypotension regardless of the underlying etiology (2). One of the seven previously reported patients was a case of shock thyroid secondary to septic shock, as in our patient (2).

The mechanism of the shock thyroid is yet clearly established. The CT hypotension complex is thought to reflect alterations in perfusion secondary to hypotension or hypovolemia affecting sympathetic splanchnic stimulations. Hypoperfusion may affect the thyroid gland, which is a highly vascular organ, resulting in cellular edema or death, as well as intracellular fluid exudation (1, 2). Alternatively, severe hypovolemia may trigger shock thyroid since it stimulates transient thyrotoxicosis, which maintains cardiac output by increasing the heart...
**Table 1. Summary of Cases of Shock Thyroid in the Literature**

| Reference                           | Underlying Disease               | Cause                          | Age | Sex | TFT | Other Accompanying CT Hypotension Complex | Follow Up CT                        | Outcome       |
|-------------------------------------|----------------------------------|--------------------------------|-----|-----|-----|-------------------------------------------|-------------------------------------|---------------|
| Brochert and Rafoth, 2006 (1)       | Trauma (gunshot)                 | 29 F                           | NA  | Shock bowel | Post-trauma days 2 and 7; normalization Discharge | Discharge |
| Brochert and Rafoth, 2006 (1)       | Trauma (vehicle collision)       | 31 F                           | NA  | Shock bowel | Post-trauma, 11 hours later and day 28 | Discharge |
| Brochert and Rafoth, 2006 (1)       | Trauma (vehicle collision)       | 12 M                           | NA  | Shock bowel | Post-trauma day 12 | Discharge |
| Han et al., 2017 (2)                 | Parkinson’s disease              | Septic shock secondary to pneumonia | 79  | M   | NA  | NA | NA | Discharge |
| Han et al., 2017 (2)                 | Bipolar I disorder               | Cardiac arrest of unknown origin Post-ROSC | 25  | F   | NA  | NA | HD 14 | Expire on HD 20 |
| Han et al., 2017 (2)                 | Trauma Post-ROSC                 | 22 F                           | NA  | Shock bowel | Peripancreatic edema | NA | Expired on HD 5 |
| Han et al., 2017 (2)                 | Trauma (bicycle accident) Post-ROSC | 17 M                           | NA  | Shock bowel, periportal edema Small-caliber IVC | NA | Expired on HD 2 |
| Our case, 2020                       | Alzheimer’s dementia             | Septic shock secondary to UTI  | 84  | F   | Low T3 level | Normal fT4 and TSH level | 22 hours after initial CT | Discharge |

F = female, fT4 = free thyroxine, HD = hospital day, IVC = inferior vena cava, M = male, ROSC = return of spontaneous circulation, TFT = thyroid function test, TSH = thyroid-stimulating hormone, T3 = triiodothyronine
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rate (1, 2). However, in our patient, these morphological changes were not associated with alterations in thyroid function. Although, the results of the thyroid function test showed slightly decreased T3 levels, this finding may be attributed to the normal aging process (6). In the abovementioned previous study, thyroid function tests were performed only in one of the seven patients, and as in our patient, the levels were in the normal range (1).

Typical CT findings of shock thyroid include inhomogeneous enhancement or heterogeneous parenchymal attenuation of the thyroid gland with perithyroidal fluid collection, representing thyroid and perithyroidal edema (1, 2). These changes are reversible; however, the recovery time for the shock thyroid remains unknown (1, 2). In our patient, follow-up CT taken 22 hours later showed a normal enhancement pattern of the thyroid gland. Previous studies reported that the thyroid gland was almost normalized after only 11 hours in a patient with trauma (1). These findings suggest that regardless of the underlying cause, proper management of hypovolemia and hypotension results in rapid normalization of the thyroid gland and adjacent tissue. Shock thyroid may be accompanied by a constellation of findings, including diffusely thickened bowel loops with enhancing walls (shock bowel), collapse of the inferior vena cava and aorta, variable enhancement of the pancreas with peripancreatic fluid, decreased enhancement of the spleen and liver, and increased enhancement of the adrenal glands (1-3). None of these findings were observed in our patient.

The clinical implications of the shock thyroid are unclear. Usually, the CT hypotension complex indicates poor prognosis (3-5). However, the relationship between poor prognosis and shock thyroid is unclear due to the lack of adequate reports and large studies. Although three out of eight reported patients expired shortly after the diagnosis, it might be dangerous to jump to hasty conclusions due to the limited number of reported cases (1, 2). However, early recognition of shock thyroid is essential because it is a critical condition leading to permanent organ damage and death. Clinicians and radiologists must recognize that shock thyroid is an indicator of hypoperfusion and initiate proper management immediately. We believe that awareness of this disorder will help distinguish shock thyroid from other pathologies.

In conclusion, shock thyroid is a useful early indicator of impending hemodynamic instability and is a reversible phenomenon. It is important for radiologists and clinicians to recognize and report shock thyroid as a manifestation of the CT hypotension complex, rather than a condition resulting from direct injury or other primary thyroid disorders.

Author Contributions
Conceptualization, all authors; data curation, K.W.H., K.M.S.; investigation, K.W.H., K.M.S.; project administration, K.M.S., K.J.H., L.K.H.; resources, L.J.H.; supervision, K.J.H., L.K.H., L.J.H.; visualization, K.W.H., K.M.S.; writing—original draft, all authors; and writing—review & editing, K.W.H., K.M.S.

Conflicts of Interest
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