Thyroid storm in the second stage of labour: a case report

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SUMMARY
A thyroid storm (or thyroid crisis) is an emergency in endocrinology. It is a form of complication of hyperthyroidism which can be life-threatening. Inadequate control of hyperthyroidism in pregnancy can develop into thyroid storm, especially in the peripartum period. We present a woman came in the second stage of labour, with thyroid storm, superimposed pre-eclampsia, acute lung oedema and impending respiratory failure. Treatment for thyroid storm, pre-eclampsia protocol and corticosteroid was delivered. The baby was born uneventfully, while the mother was discharged after 5 days of hospitalisation. Delivery is an important precipitant in the development of thyroid storm in uncontrolled hyperthyroidism in pregnancy. Although very rare, it can cause severe consequences. Diagnosis and treatment guidelines for thyroid storm were available and should be done aggressively and immediately. Uncontrolled hyperthyroidism should be prevented by adequate control in thyroid hormone levels, especially before the peripartum period.

BACKGROUND
Hyperthyroidism in pregnancy is rare, occurring in only 1–4 in 1000 pregnancies. The incidence of hyperthyroidism in pregnancy is about 0.2% and most cases are subclinical.1 Thyroid storm is a rare complication of improper treatment of hyperthyroidism in pregnancy.2 A thyroid storm is precipitated when hyperthyroidism effects surpass the patient’s ability to compensate cardiovascular, thermoregulatory and metabolic systems.3 The risk of developing thyroid storm in pregnant hyperthyroid patients can be 10 times higher than in the general population.4 Precipitant of thyroid storm includes thyroid surgery, withdrawal of antithyroid drugs, infection, emotional stress and trauma.1 Several obstetric conditions that can precipitate thyroid storm include pre-eclampsia, induction of labour, surgery and pregnancy itself.4

CASE PRESENTATION
A 20-year-old woman was admitted to the emergency room due to the second stage of labour on gravida two parity one 29 weeks of gestational age, singleton lives intrauterine, mother with superimposed pre-eclampsia with severe features and history of uncontrolled thyroid disease. Patient was diagnosed as hyperthyroidism since 5 years before admission, but never took adequate treatment. She had bad antenatal care and did only once visit to the midwife without any written data, but she was told that her blood pressure was high and given nifedipine 10 mg once per day without her taking the medicine. During this pregnancy, she had no history of screening for hyperthyroidism, neither does she took any medicine to lower her thyroid levels. The patient came to emergency room by herself due to regular contraction 6 hours before admission. The patient was in agitation state of consciousness, she had Glasgow Coma Score of 12 consisted of 3 for eye opening response score, 4 for verbal response score and 5 for motoric response score. Her blood pressure in the emergency room was 220/120 mm Hg, and her heart rate 156 times/min. She had fever with temperature of 38.4°C and an increased respiratory rate of 40 times/min with oxygen saturation 95% in room air.

INVESTIGATIONS
In the general examination, the patient had bilateral protruding eyes suggesting exophthalmos. Diffuse struma in the neck was palpable, with no bruit. Auscultation in both lungs grade 3/6 loudest on the apex with rales in the bilateral bottom side of the lungs. Extremities were warm, there was bilateral pitting oedema. In the obstetric examination, fundal height was 30 cm, with regular contractions of four times in 10 min with a duration of 45 s. Fetal heart rate was 140 beats/min. Vaginal touch examination revealed a fully dilated cervix, no amniotic membrane was present and fetal feet were palpable on station +2. Patient was underwent laboratory examination, such as blood gas analysis, urinalysis, ureum creatinine level, aspartate aminotransferase (AST)/alanine aminotransferase (ALT) level and antigen swab of SARS-CoV-2 as the COVID-19 screening protocol in our hospital. Her blood gas analysis showed a little bit increasing of pH (7.455) while the HCO3 decreased to 17.8 mmol/L and her pCO2 decreased to 25.1 mmol/L, in urinalysis examination there was proteinuria +2 and haematuria +1. In addition, her electro gram showed sinus tachycardia. Her thyroid stimulating hormone (TSH) and FT4 level come afterwards, and showed low TSH (<0.02 µIU/mL) and high free T4 (3.75 ng/dL), her lactate dehydrogenase (LDH) was 547 U/L as well.

After quick examination, we concluded this patient had second stage of labour on gravida two parity one 29 weeks of gestational age, singleton lives intrauterine, mother with thyroid storm, superimposed pre-eclampsia with severe features, acute lung oedema and impending respiratory failure.
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DIFFERENTIAL DIAGNOSIS

Based on the hypertension condition with mean arterial pressure more than 120 mm Hg and we found proteinuria followed by increasing LDH level, we first suggested this patient got pre-eclampsia condition compounded by HELLP (hemolysis, elevated liver enzyme and low platelet) syndrome. Nevertheless, there was no increasing of AST/ALT level in this patient and her platelet count still in normal limit, the diagnosis can be eliminated. We also found her blood glucose increased to 143 mg/dL but we thought it as reactive hyperglycemia condition because the blood glucose level was decreasing after we did re-evaluation.

Moreover, the patient came to the hospital with dyspnoea, with her respiratory rate 40 times/min and her oxygen saturation 95% in room air. In this pandemic condition, initially we suspect SARS-CoV-2 infection. However, without any previous coughing, fever symptom, her antigen swab of SARS-CoV-2 IgG/IgM showed non-reactive result and her chest X-ray represented cardiomegaly with lung oedema appearance compounded by suspected bilateral pleural effusion, we temporarily set aside this differential diagnosis until the PCR swab result come out.

TREATMENT

The patient was given a non-rebreathing mask with oxygen 15 L/min, hereinafter her oxygen saturation level was going up. Pre-eclampsia protocol was carried out immediately. We gave bolus magnesium sulphate (MgSO₄) 40% 4 g continued by 1 g per hour as anticonvulswnt agent and neuroprotector for the baby. We added nifedipine 10 mg every 20 min four times to lower her blood pressure. For her thyroid condition we delivered propranolol 40 mg, propylthiouracil (PTU) 600 mg, and 5 drops of Lugol iodine 2%. Furthermore, as this patient came with bad antenatal care and we were not sure about her gestational age, also to prevent respiratory distress and interventricular haemorrhage we gave dexamethasone 12 mg single shot. Fetal foot extraction along with episiotomy was performed afterwards, and delivery was successful 5 min after this patient came to the emergency room. A baby girl was born, with a birth weight of 1950 g, length of 38 cm and a 5 min apgar score of 7/8. Amniotic fluid was scanty and the placenta was delivered completely after 15 min. Episiotomy wound was closed by perineorrhaphy. In the third phase of labour her blood pressure slowly decreased to 180/100 mm Hg, her pulse was still in 140 times/min, saturation oxygen was 96% using non-rebreathing mask oxygen 15 L/min, her respiration rate came to 30 times/min but her temperature were not going down.

In the fourth phase of labour, the patient’s blood pressure slowly decreases to 134/88 mm Hg and her pulse slowed down to 120 times/min, her temperature normalised to 37°C and she was getting alert with consciousness level became comatose.

The baby was born with APGAR to score 7/8. Then, she got CPAP (continuous positive airway pressure) with FiO₂ 21% and PEEP (positive end expiratory pressure) 7 to help her breathing.

OUTCOME AND FOLLOW-UP

Collaborative management was done by endocrinologists from internal medicine and obstetricians, and the patient was discharged after 5 days of treatment. The paediatric checked for the baby’s free T4 and the result was within normal limit (1.87 ng/dL). The baby’s Ballard score similar with 35 weeks of gestation age which showed a discrepancy between prior gestation age that can be caused by her irregular menstruation cycle and no antenatal care data available. The mother’s TSH and free T4 level come afterwards, and showed low TSH (<0.02 µIU/mL) and high free T4 (3.75 ng/dL) which similar with hyperthyroid condition. Patient was discharged and the follow-up was continued in outpatient clinic.

DISCUSSION

Our patient was admitted to the emergency room due to second stage of labour on gravida two parity one 29 weeks of gestation age, singleton lives intrauterine; mother with thyroid storm, superimposed pre-eclampsia with severe features, acute lung oedema and impending respiratory failure. Previously, this patient had inadequate hyperthyroid treatment. She has been diagnosed with hyperthyroidism since she was 15 years old, but the patient only admitted to buying over-the-counter drugs without a prescription and took them only in the presence of symptoms. Our patient also had bad antenatal care. She came to the midwife once and lost her antenatal care data, but she was told that her blood pressure is high and she was given nifedipine 10 mg once per day without her taking the medicine. During this pregnancy, she had no history of screening for her hyperthyroidism, neither does she took any medicine to lower her thyroid levels.

Thyroid storm is most diagnosed clinically. Laboratory findings are of little importance since they cannot distinguish uncomplicated thyrotoxicosis with impending thyroid storm, and could not provide a diagnosis. Thyroid storm is diagnosed in patients with a Burch-Wartofsky Point Scale (BWPS) of ≥45 or Japanese Thyroid Association categories of thyroid storm 1 or 2.³ The BWPS took into accord the patient’s theremoregulatory dysfunction, central nervous system effects, gastrointestinal-hepatic dysfunction, cardiovascular dysfunction and precipitant history.³ Patients meeting these criteria with a systemic decompensation need to be managed aggressively. In patients with impending thyroid storm (BWPS of 25–44), aggressive therapy could be done based on clinical judgement.³ In this case, the patient’s Burch and Wartofsky criteria score is 50, highly suggestive of thyroid storm, needing immediate aggressive treatment. We use Modified Early Warning Score (MEOWS) for assessing the severity of this case. MEOWS is early detection of clinical signs of deterioration in women who were developing critical illnesses. Her MEOWS was 29 and categorised as red criteria.

We gave emergency treatment for our patients consecutively based on the airway, breathing and circulation. We ensured her airway is clear before giving further treatment. As this patient had hypertension with blood pressure up to 220/120 mm Hg, we assumed she got pre-eclampsia. We decided to give magnesium sulphate (MgSO₄) 40% bolus 4 g, continued by 1 g/hour and nifedipine 10 mL every 20 min four times to treat the condition. MgSO₄ is the main treatment for pre-eclampsia as anticonvulsant which works in the neuromuscular junction while the nifedipine works as calcium channel blocker to lower the blood pressure. Because this patient did not give adequate information about her previous antenatal care and we need to do delivery in short period, so we gave single shot dexamethasone 12 mg to prevent respiratory distress and interventricular haemorrhage for the baby. We also gave propranolol 40 mL which works as a beta-blocker to slower the heart rate, this drug is in Food and Drug Administration (FDA) category C for pregnant women. Furthermore, PTU was given in 600 mL dosage to inhibit the production of new thyroid hormone in the thyroid gland. It acts by inhibiting the enzyme thyroid peroxidase, which usually functions to convert iodide to iodine molecule and incorporate the iodine molecule into amino acid tyrosine. This drug is in old

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FDA category D for pregnancy but depend on the trimester of pregnancy as the result may vary. Other drug which can be used for hyperthyroid is methimazole. For the first trimester, PTU is the drug of choice because it has less effect on fetal birth defect. Since, PTU has more rapid onset of action and the additional benefit of inhibition of peripheral deiodinase enzyme-mediated conversion of T4 into T3, this drug is preferred in thyroid storm case. Lugol’s solution or as saturated solution of potassium iodide decreases its transport into the thyroid, inhibits iodide organification (the Wolff-Chaikoff effect), and rapidly blocks the release of T4 and T3 from the gland. Potassium iodide is in FDA category D for pregnancy.

According to American Thyroid Association guidelines, 5 treatments for thyroid storm are; (1) decreasing thyroid hormone secretion and synthesis, (2) blocking thyroid hormone effects in the tissue level, (3) reversal of systemic decompensation, (4) treating precipitating event/coexisting illness and (5) definitive therapy. Therapy includes PTU 500–1000 mL loading dose followed with 250 mL q4h (PTU is more recommended than methimazole), propranolol 60–80 mL q4h, iodine 5 drops (250 mL) q6h and hydrocortisone 300 mL loading dose followed by 100 mL q8h. 6 Our patient was treated with PTU, propranolol and Lugol solution, along with pre-eclampsia treatment of nifedipine and MgSO4.

Thyroid storms can cause life-threatening complications, with mortality rates 8%–25%. A case report by Kitazawa et al presented a 41-year-old G1P1 woman with respiratory failure following delivery. 6 The women had uncontrolled hyperthyroidism throughout pregnancy, had an unremarkable delivery, but respiratory failure occurred 9 min after delivery. In our patient, her respiratory rate was 40 times/min with oxygen saturation of 95%, and rales are heard in most of her lung surface, suggesting acute lung oedema with impending respiratory failure.

Both our patient and Kitazawa et al’s had uncontrolled hyperthyroidism, and a thyroid storm occurred within the peripartum period. It is known that heart failure occurs in 10% of cases of thyrotoxicosis in pregnancy. This is due to a decrease in peripheral vascular resistance and increased cardiac output during pregnancy. 6 It was assumed that in our case, chronic uncontrolled hyperthyroidism leads to cardiac insufficiency, and added by increased stress in the delivery process caused the failure of cardiac compensation. Fortunately, our patient was successfully managed and no further complications arise.

Hyperthyroidism reduces fertility, hence, there are very few cases of hyperthyroidism in pregnancies. Studies are rarely conducted due to their rarity, and preventive acts are seldom mentioned. 8 Despite the rarity, clinicians must always be aware of uncontrolled hyperthyroid in pregnant women, since it can cause devastating effects for the fetus, such as growth restriction, premature birth, low birth weight, miscarriages and also for the mother, such as pre-eclampsia, respiratory failure and death. 8 General prevention of hyperthyroidism during gestation could be done by ensuring at least 200 µg of iodine intake per day. Proper antenatal care should be done every month, including in it a clinical screening for hyperthyroidism or Grave’s disease. Pregnant women should always avoid radiation therapy. 10

Newborn who was born from hyperthyroid mother should undergo the examination to ensure there is no sign of congenital hyperthyroidism. The clinician can measure the plasma levels of free T4, T3 and TSH. Even if these are normal, they should be repeated 3–7 days later because of the possibility of the delayed appearance of hyperthyroidism. 11 Some suggestion included assessing maternal TSH receptor antibodies from the cord blood. If the results are negative, no specific neonatal follow-up is needed. But if the examination is unavailable or the result is positive, we can regard the newborn as ‘at risk’ newborn for the development of hyperthyroidism. 12

Follow-up during the first postpartum year for the mother is highly recommended in woman with hyperthyroid during pregnancy. Recurrence in the first 3 months post partum could be due to Graves’ hyperthyroidism or postpartum thyroiditis. The postpartum thyroiditis occurs in 7%–10% of postpartum women, although this varies depending on iodine intake and genetic factors. Thyroid function tests are indicated at three and 6 months post partum in these women and those with known autoimmune disease, previous postpartum thyroiditis or chronic viral hepatitis. The pregnancy should be avoided until the thyroid function become normal. 13

The patient was discharged after 5 days of treatment.

Thyroid storm is a rare complication of hyperthyroidism.

Delivery is an important precipitating factor, especially in pregnancies with uncontrolled hyperthyroidism.

Thyroid storm can cause life-threatening complications, including acute decompensated heart failure, acute lung oedema and respiratory failure.

Immediate and aggressive treatment needs to be done in patients with thyroid storms, especially in the peripartum period.

Our case of thyroid storm in the second stage of labour is unique, and patient outcome was satisfying, despite the appalling condition at the time of admission.

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