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**Pheochromocytoma in a 49-year-old woman presenting with acute myocardial infarction: A case report**

Wu HY *et al*. Pheochromocytoma presenting with acute myocardial infarction

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**Abstract**

BACKGROUND

Pheochromocytoma is a rare endocrine tumor arising from chromaffin cells and having extensive and profound effects on the cardiovascular system by continuously or intermittently releasing catecholamines. The clinical manifestations of pheochromocytoma are diverse, and the typical triad, including episodic headache, palpitations, and sweating, only occurs in 24% of pheochromocytoma patients, which often misleads clinicians into making an incorrect diagnosis. We herein report the case of a patient with intermittent chest pain and elevated myocardial enzymes for 2 years who was diagnosed with pheochromocytoma.

CASE SUMMARY

A 49-year-old woman presented with intermittent chest pain for 2 years. Two years ago, the patient experienced chest pain and was diagnosed with acute myocardial infarction, with 25% stenosis in the left circumflex. The patient still had intermittent chest pain after discharge. Two hours before admission to our hospital, the patient experienced chest pain with nausea and vomiting, lasting for 20 min. Troponin I and urinary norepinephrine and catecholamine levels were elevated. An electrocardiogram indicated QT prolongation and ST-segment depression in leads II, III, aVF, and V3-V6. A coronary computed tomography angiogram revealed no evidence of coronary artery disease. Echocardiography showed left ventricular enlargement and a decreased posterior inferior wall motion amplitude. Contrast-enhanced computed tomography demonstrated an inhomogeneous right adrenal mass. The patient successfully underwent laparoscopic right adrenalectomy, and histopathology confirmed adrenal pheochromocytoma. During the first-year follow-up visits, the patient was asymptomatic. The abnormal changes on echocardiography and electro-cardiogram disappeared.

CONCLUSION

Clinicians should be aware of pheochromocytoma. A timely and accurate diagnosis of pheochromocytoma is essential for alleviating serious cardiac complications.

**Key Words:** Pheochromocytoma; Catecholamine; Cardiac complications; Acute myocardial infarction; Chest pain; Case report

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**Core Tip:** Pheochromocytoma releases excessive amounts of catecholamines, which can have mild to catastrophic effects on the cardiovascular system, such as hypertension, myocardial infarction, cardiomyopathy, arrhythmias, and heart failure. However, most of the cardiovascular complications are reversible after pheochromocytoma resection. Clinicians should be aware of pheochromocytoma. A timely and accurate diagnosis of pheochromocytoma is essential for alleviating serious cardiac complications.

**INTRODUCTION**

Pheochromocytoma is a rare endocrine tumor arising from chromaffin cells of the adrenal medulla or extra-adrenal paraganglion and having extensive and profound effects on the cardiovascular system by continuously or intermittently releasing catecholamines, mainly epinephrine and norepinephrine[1,2]. Pheochromocytoma has two main types, high norepinephrine/epinephrine secretion and high epinephrine/ norepinephrine secretion. The norepinephrine type affects mainly the cardiovascular system, manifesting mainly as hypertension, while the epinephrine type affects mainly the metabolism, manifesting mainly as hyperglycemia. A timely and accurate diagnosis of pheochromocytoma is essential for alleviating serious cardiac complications caused by an overdose of catecholamines. We report the case of a patient with intermittent chest pain and elevated myocardial enzymes for 2 years who was diagnosed with pheochromocytoma, and the abnormal changes in electrocardiogram (ECG) and echocardiography caused by pheochromocytoma disappeared after tumor resection.

**CASE PRESENTATION**

***Chief complaints***

A 49-year-old woman presented with intermittent chest pain for 2 years.

***History of present illness***

Two years ago, the patient experienced chest pain lasting for 20 min. The patient was diagnosed with acute myocardial infarction with a significant increase in troponin in a local hospital, and coronary angiography showed 25% stenosis in the left circumflex. The patient was treated with aspirin, clopidogrel, and statins. However, she still had intermittent chest pain, lasting for 5-10 min each time, after discharge. Two hours before admission to our hospital, the patient experienced chest pain with nausea and vomiting lasting for 20 min.

***History of past illness***

A history of hypertension and diabetes was denied.

***Personal and family history***

The patient had a free personal history and denied a family history of premature coronary artery disease.

***Physical examination***

Vital signs on arrival showed body temperature of 36.8 ℃, blood pressure of 140/80 mmHg, a regular pulse of 84 beats per minute, and a respiratory rate of 18 breaths per minute. Heart and lung examinations showed no abnormalities. Jugular vein engorgement or peripheral edema was not found.

***Laboratory examinations***

The troponin I level was 1.14 ng/mL (normal range < 0.04). The urinary norepine-phrine level was 296.2 nmol/24 h (normal range 80.3-164.0), and the urinary catecholamine level was 327.8 nmol/24 h (normal range 94.5-238.3). The fasting blood glucose level was 5.6 mmol/L (normal range 3.9-6.1). B-type natriuretic peptide, 24-h urinary epinephrine, hemoglobin, leukocytes, amylase, electrolytes, liver function, renal function, and D-dimer were not significantly abnormal.

***Imaging examinations***

A 12-lead ECG indicated QT prolongation (QTc 533 ms) and ST-segment depression in leads II, III, aVF, and V3-V6 (Figure 1). A coronary computed tomography angiogram revealed no evidence of coronary artery disease (Figure 2). Echocardiography showed left ventricular enlargement (systolic and diastolic diameters of 45 mm and 57 mm, respectively) and a decreased posterior inferior wall motion amplitude (left ventricular ejection fraction of 51%). Chest computed tomography showed no obvious abnormality, but abdominal computed tomography showed an adrenal mass. Contrast-enhanced computed tomography demonstrated an inhomogeneous right adrenal mass (6.1 cm × 3.9 cm, Figure 3).

**FINAL DIAGNOSIS**

The patient was diagnosed with pheochromocytoma.

**TREATMENT**

The patient was transferred to the urology department and underwent successful laparoscopic right adrenalectomy. Histopathology confirmed adrenal pheochro-mocytoma.

**OUTCOME AND FOLLOW-UP**

The patient was free of complications during hospitalization. During the first-year follow-up visits, the patient was asymptomatic and had normal blood pressure. Echocardiography showed a normal left ventricular size (systolic and diastolic diameters of 32 mm and 48 mm, respectively), no abnormal wall motion, and normal left ventricular function (left ventricular ejection fraction of 63%). ECG showed that QT prolongation and ST-segment depression in leads II, III, aVF, and V3-V6 disappeared (Figure 4). The patient showed no signs of recurrence and had normal urine norepinephrine and catecholamine levels.

**DISCUSSION**

Pheochromocytoma, which is derived from chromaffin cells, produces excessive amounts of catecholamines, especially epinephrine and norepinephrine, whose continuous or intermittent release can cause serious cardiovascular complications, such as hypertension, myocardial infarction, cardiomyopathy, arrhythmias, and heart failure[3,4]. It is estimated that the prevalence of pheochromocytoma during a lifetime is approximately 0.015%-0.04%, while its prevalence at autopsy is even higher (approximately 0.05%), which indicates that many pheochromocytomas are not diagnosed during the lifetime and may be the cause of death[5]. According to the laboratory results, the patient in our case belongs to high norepinephrine secreting pheochromocytoma.

Hypertension is one of the most common cardiovascular manifestations of pheochromocytoma, which can manifest as persistent or paroxysmal hypertension, mainly depending on the pattern of catecholamine secretion[6]. Approximately 90% of patients with pheochromocytoma have persistent or paroxysmal hypertension. Persistent hypertension is more common in patients with pheochromocytomas that continuously release high levels of norepinephrine. Paroxysmal hypertension is more common in patients with pheochromocytomas that intermittently secrete large amounts of epinephrine[7]. It is worth noting that a small proportion of patients with pheochromocytoma have normal blood pressure[8].

Patients with pheochromocytoma can develop catecholamine cardiomyopathy (including hypertrophic cardiomyopathy, dilated cardiomyopathy, and Takotsubo-like cardiomyopathy), myocardial ischemia, or even myocardial infarction. An increase in catecholamine levels can lead to increased production of reactive oxygen species, increased oxygen consumption, increased cardiac afterload, vasoconstriction, cell hypertrophy, cardiac remodeling, and even myocarditis[1,9]. The patient may present with chest pain, chest tightness, and sweating. ECG may show ST segment elevation or depression and T-wave inversion. Myocardial enzymes can also be elevated.

Approximately 50%-70% of pheochromocytoma patients complain of palpitations. Patients with pheochromocytoma may have a variety of arrhythmias, including atrial flutter, atrial fibrillation, supraventricular tachycardia, torsade de pointe ventricular tachycardia, ventricular fibrillation, and asystolic arrest[10]. Catecholamines can also cause the QT interval to be prolonged, which may induce torsade de pointes ventricular tachycardia and be life-threatening[11,12]. The reasons for arrhythmias caused by pheochromocytoma are considered to be multifactorial. Excessive catecholamine stimulation of beta-adrenergic receptors is one of the main reasons[13-16].

Although pheochromocytoma can cause many cardiovascular complications, most of them are reversible after pheochromocytoma resection[17-20]. In our case, the patient did not have the typical triad of pheochromocytoma but presented with acute myocardial infarction that caused abnormal changes in ECG and cardiac structure and function. However, the abnormal changes in ECG and echocardiography caused by pheochromocytoma disappeared after tumor resection.

**CONCLUSION**

The clinical manifestations of pheochromocytoma vary, and many patients do not have the typical triad, which easily leads to missing the diagnosis of pheochromocytoma. Pheochromocytoma causes an excess of catecholamines, which has a variety of potentially damaging effects on the cardiovascular system. However, most of the cardiovascular complications are reversible after pheochromocytoma resection. Clinicians should be familiar with clinical manifestations of pheochro-mocytoma, which helps raise clinical suspicion and facilitate the early diagnosis and treatment of pheochromocytoma.

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**Footnotes**

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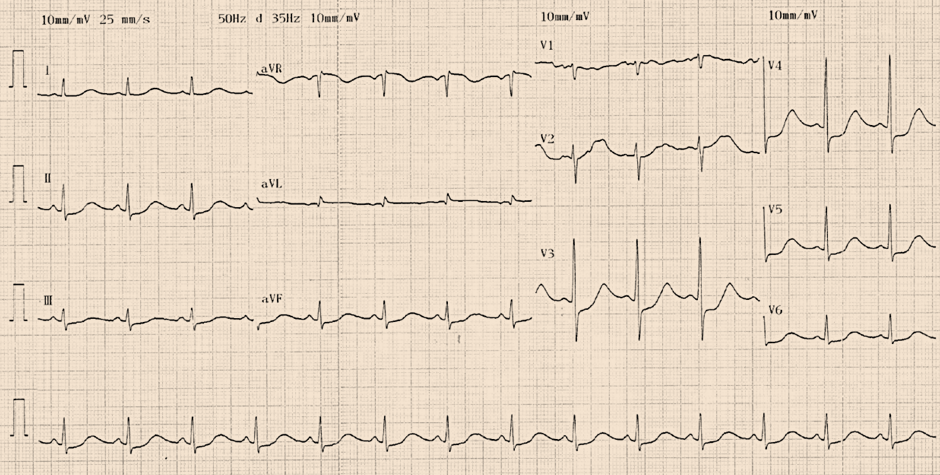
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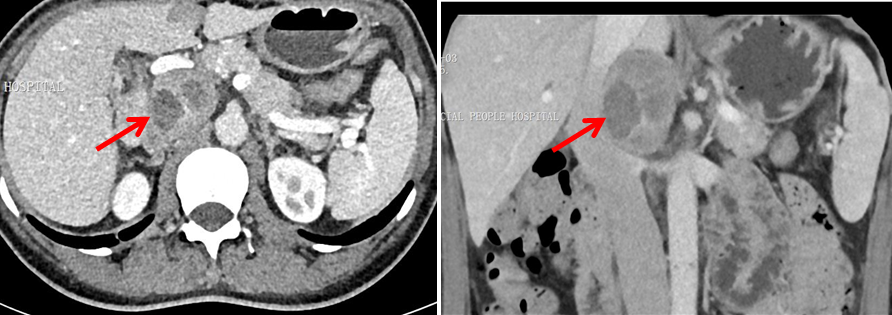
**Figure Legends**

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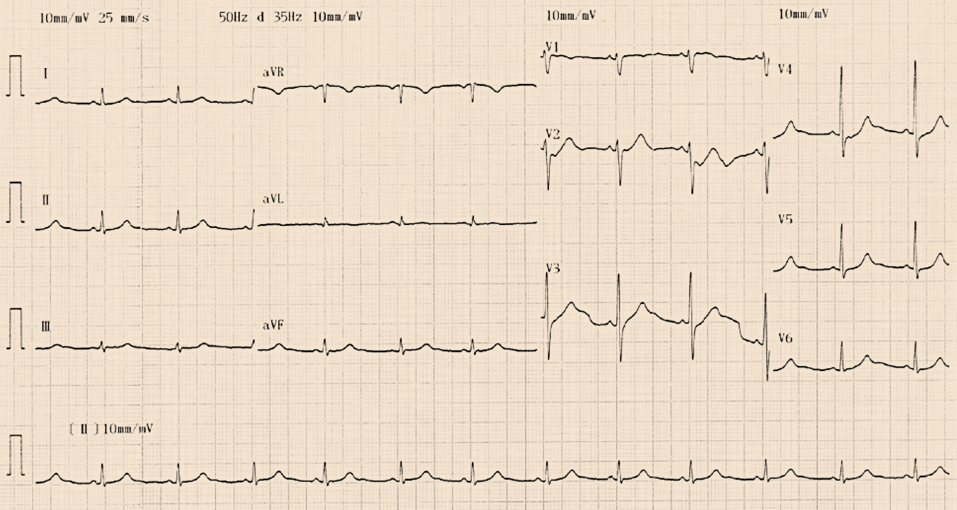
**Figure 1** **Twelve-lead electrocardiogram indicated QT prolongation (QTc 533 ms) and ST-segment depression in leads II, III, aVF, and V3-V6 at admission.**

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**Figure 2 Coronary computed tomography angiography revealed no evidence of coronary artery disease.** A and B: Left coronary artery; C: Right coronary artery.

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**Figure 3 Contrast-enhanced computed tomography demonstrated an inhomogeneous right adrenal mass (6.1 cm × 3.9 cm, orange arrows).**

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**Figure 4 Twelve-lead electrocardiogram indicated QT prolongation and ST-segment depression in leads II, III, aVF and V3-V6 disappeared at the 1-mo follow-up after pheochromocytoma resection.**



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