CARDIOVASCULAR EFFECTS OF COCAINE - Case report

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CARDIOVASCULAR EFFECTS OF COCAINE

INTRODUCTION

Cocaine is one of the most commonly used illegal drug worldwide. It is highly psychologically addictive, but it has no physical withdrawal effects. The main mechanism action of cocaine is the inhibition of noradrenaline and dopamine re-uptake from the synaptic cleft which causes overstimulation of alpha and beta adrenergic receptors. [1] Cocaine abuse may cause acute and chronic cardiovascular diseases - myocardial ischemia, myocardial infraction, arrhythmias, cardiomyopathy, myocarditis, hypertension, aortic dissection. (Figure 1) Cocaine users have 4-8 times higher mortality compared to general population. It is also the most common cause of drug-related deaths.

CASE REPORT

A 39 year-old male with pneumonia and heart failure (de novo), with no previous of cardiac problems, presented shortness of breath. He admitted that he was taking cocaine 2-3 times a week for several years by inhaling it cocaine through his nose. Three weeks ago he underwent a laryngological surgery due to chronic sinusitis.

During physical examination the patient had blood pressure of 105/70 mmHg and his heart rate was 110 beats/min and regular. In cardiac auscultation a high-pitched holosystolic murmur on the apex, radiating to the armpit could be heard. Electrocardiogram revealed sinus rhythm with left ventricular overload and hypertrophy, negative T in leads I, II, aVL, V4-V6 (Figure 2).

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**FIGURE 1** Short-Term and long-term effects of cocaine use.

- SHORT-TERM
  - ARRHYTHMIAS
  - MYOCARDIAL INFARCTION
  - STROKE
  - AORTIC DISSECTION

- LONG-TERM
  - HEART FAILURE
  - CARDIOMYOPATHY
  - ENDOCARDITIS
  - HYPERTENSION
Troponin I levels peaked at 77 ng/ml, creatine was 0.97 mg/dl, B-type natriuretic peptide was 2003 pg/ml. Chest X-ray revealed an enlarged cardiac silhouette (Figure 3). An echocardiogram demonstrated akinesis of apex, anterior wall and interventricular septum, hypokinesis of other left ventricular walls, enlargement of all heart cavities, left ventricular ejection fraction of 25%. Coronarography didn’t show any abnormalities in coronary vessels (Figure 4). Cardiac MRI showed dilation and severe retardation of the systolic function of both chambers with no underlying cause. There wasn’t contrast enhancement typical to inflammation or myocardial ischemia.

COCAINE-INDUCTED ACUTE MYOCARDIAL INFARCTION

Overstimulation of beta adrenergic receptors, caused by cocaine, increases contractility and heart rate, which leads to a higher oxygen request. [2] Moreover cocaine is causing overstimulation alfa-1 and alfa-2 receptors, which causes coronary artery spasm. Ischemic effect appears when heart rate and blood pressure is higher, requiring more oxygen supply. (Figure 5) Cocaine also activates platelet aggregation, which is leading to formation of blood clots in the coronary vessels. It also accelerates atherosclerosis. [3] However coronarography may not show any abnormalities in coronary vessels. [4]

In patients with cocaine-related pain, a heart attack occurs in 6% of cases. Cocaine inducted acute myocardial infarction is the most common among population of patients between the ages of 18-45. [5-6] It needs to be suspected in young patients with chest pain.

The risk of a heart attack increases 24 times directly after using cocaine. Most infarctions happen during the first 3 hours after cocaine use[7]. Complications rarely occur after
12 hours, so patients should be monitored by ECGs and cardiac troponins for at least a 12 hours. [8] Cocaine is affiliated with an increased risk of acute coronary syndrome even without presence of coronary heart disease, due to coronary artery spasms.

The treatment is similar to the one that is used in acute myocardial infraction. Until recently it was considered that it is not recommended to use B-blockers in treatment, because of vasoconstriction properties. However because of the fact, that a lot of patients didn’t admit that they used cocaine, received B-blockers and they didn't affect them negatively, but even they had a beneficial effect. [9] B-blockers are causing lower request for oxygen. Treatment was associated with lower rates of deaths. The guidelines of ACC/AHA from 2012 states that it is acceptable to use B-blockers at the patients with hypertension or tachycardia, if they received vasodilator drug. [7] Additionally it is recommended to use benzodiazepines for the patients who are hyper excitable with hypertension and tachycardia. [10] Benzodiazepines reduce the hemodynamic effects of cocaine. Sedation is also indicated in patients with psychomotor agitation.

The lack of coronary artery stenosis in coronaryography, especially in young patients with acute coronary syndrome, should lead to suspicion of taking cocaine as a cause. The guidelines of ACC/AHA from 2008-2012 recommended to using bare-metal stents in cocaine abusers. [7]

The psychological state and euphoria associated with cocaine use, reduce the number of patients, who were referred to a hospital. That is why the reports on the number of heart attacks caused by cocaine use are underestimated.

With regard to the case report the possibility of myocardial infarction was ruled out because of the lack of chest pain and slightly increased TnI without typical dynamic, characteristic for myocardial infarction.

Coronarography didn’t show any abnormalities in coronary vessels.

FIGURE 4  Coronarography of presented patient.
AORTIC DISSECTION

Cocaine induces the apoptosis of smooth muscle cells, which build the vessel wall, making them more susceptible to damage. [11] Sudden increase of blood pressure up to high values, after exposure to cocaine, may cause aortic dissection. It is estimated that 0.5% is connected with cocaine usage. Should be expected aortic dissection especially in young patients with chest pain. First line treatment is controlling blood pressure, heart rate and level of pain. Further therapy is surgery. Patients have a high risk of dangerous complications and even of death.

In this particular case report aortic dissection is not expected because there was no chest pain. Also there was no evidence of aortic dissection in imaging studies.

CARDIOMIOPATHY

Cardiomyopathy in cocaine users is caused by direct toxic effect on myocytes, oxidative stress, calcium channel dysfunction and also excessive adrenergic stimulation. Toxic influence of cocaine is causes inflammation of the heart muscle, fibrosis and as a result heart failure. Oxidative stress has toxic effect on myocardium. It is caused by high level of peroxides and free radicals, which are damaging all components of cardiomyocytes. [12] Raised calcium inflow into smooth muscle cells results in increased contraction force, but when it is chronic, it weakens the myocardium, causes overgrowth and decreases the left ventricle ejection fraction. [13] Excessive adrenergic stimulation caused by the inhibition
of noradrenaline and dopamine reuptake causes increase of heart rate and elevated blood pressure. [14] This affects the remodeling of the myocardium. Improvement of the left ventricle ejection fraction is possible after reaching total abstinence from cocaine.

Cocaine-associated cardiomyopathy should be suspected in young patients with heart failure. Symptoms reported by patients are similar to those with dilated cardiomyopathy. Mostly it is dyspnea and fatigue. [15] However it was revealed that symptoms such as leg oedema are much less intense. Cardiomyopathy usually has a more severe course. Usually patients also present symptoms of increased adrenergic stimulation, such as tachycardia, hypertension and those associated with the psychotropic action of cocaine. In physical examination it’s possible to notice symptoms of cocaine usage – depending on the route of administration of the drug such as scars after intravenous injections or damage of the nasal septum.

It is reasonable to use B-blocker treatment in patients with cocaine-related cardiomyopathy who had an abstinence for more than 6 months, as standard therapy of the left ventricular dysfunction.

The clinical picture of this patient led to the diagnosis - cardiomyopathy. Patient presented dyspnea and impaired of effort tolerance. It was confirmed by the results of additional tests that includes among others echocardiography and MRI.

ARRHYTHMIA

Cocaine increases the risk of arrhythmia, due to excessive adrenergic stimulation and it also affects the blockage of sodium and potassium channels. In small doses the effect of stimulation of the adrenergic system dominates, in large doses it blocks ion channels. [16] Usage of cocaine induces ventricular and supraventricular arrhythmias. It can cause sinus tachycardia or atrial fibrillation, when arrhythmia is the result of excessive adrenergic stimulation. Blockade of potassium channels causes QT interval prolongation, which can lead to ventricular tachycardia, including torsade de pointes. Cocaine-induced myocardial ischemia can cause ventricular tachycardia or ventricular fibrillation. [17]

The first-line treatment is B-blockers in combination with alpha-blockers. Sodium bicarbonate and lidocaine are also used. Ca-blockers have been shown to reduce arrhythmia. It is important to correct electrolyte disturbances. Arrhythmias resistant to pharmacological treatment can be corrected by cardioversion or defibrillation. It has been proven that ablation is feasible, safe and effective in patients with drug-resistant cocaine-induced tachycardia. [18] The most common cause of sudden death in people addicted to cocaine is ventricular fibrillation. [19] The main therapeutic goal of a patient experiencing cocaine-related arrhythmia is the abstinence of using cocaine.

During the hospitalization of the patient in the ECG monitoring, no arrhythmias were recorded.

ENDOCARDITIS

Cocaine is one of the most commonly used drugs for intravenous use. This route of administration is associated with high risk of endocarditis.

Cocaine also has a damaging effect on the vascular endothelium. Endocarditis in drug users is mostly on the tricuspid valve. [20] Treatment consist of multi-week antibiotic therapy but the most important goal is to stop using cocaine.

The results of laboratory tests and the result of MRI allowed to rule out endocarditis at the patient.
STROKE

Strokes occur most often in older patients. However recently there was an increase in number of incidences among younger people, which is related to the use of illegal drugs. Cocaine increases the risk of ischemic and hemorrhagic stroke. [21] The most common causes of ischemic strokes in patients using cocaine are atherosclerotic stenosis of the large artery and occlusion of the small cerebral vessels.

Vasoconstriction and prothrombotic properties increase the risk of ischemic stroke. The use of cocaine accelerates the process of atherosclerosis. [22] Long-term use of cocaine increases the risk of stroke, including endothelial changes, atherosclerosis and vasculitis. Most ischemic cocaine-related strokes occur in people with a distant history of cocaine use.

Hemorrhagic strokes are associated with a sudden rise in blood pressure after cocaine ingestion. They are more common in active cocaine users. The most common cause is rupture of the aneurysm. [23] It has been proven that brain aneurysms are likely to happen to people who use cocaine. The use of cocaine is associated with a greater probability of hemorrhagic stroke than ischemic stroke. [24]

The patient did not show symptoms of neurological deficit. Diagnosis for stroke has not been extended.

CONCLUSION

As demonstrated, cocaine has many negative effects on the cardiovascular system. One of the main goals of treatment is total abstinence. An important aspect is to make patients aware of the consequences of further cocaine use. Cocaine use can cause both short-term and long-term effects.

In young patients presenting cardiovascular disease symptoms, but not having any other loads, the drug use should be considered in the diagnosis, because it is most often occurs in this age group.

In this case report, the patient was diagnosed for toxic cocaine cardiomyopathy. The direct cause of the disease was long-term and regular use of this drug.

The patient was qualified for ICD implantation as the primary prevention of sudden cardiac death.

REFERENCES


