Kounis syndrome

Kounis syndrome is a group of symptoms manifested as unstable vasospastic or non-vasospastic angina, or even as acute myocardial infarction. Inflammatory mediators released into a bloodstream during some allergic reaction are most frequently mentioned to be a trigger for Kounis syndrome. We present a case of acute myocardial infarction associated with an allergic reaction after a honeybee sting, in an 79-year-old Serbian male.

Key words: insect bites and stings; hypersensitivity, immediate; myocardial infarction.
CASE REPORT

A 79-year-old male was admitted to the Emergency room in Clinical Centre Kragujevac two hours after he was stung on the thumb of his left leg by a honeybee while working in his field. After stinging he had a short period of dizziness and respiratory discomfort without loss of consciousness. After a while he felt sudden chest pain, itching, dyspnea and fatigue so he was transferred to our hospital by his relatives. He had a history of treated arterial hypertension and consumption of alcohol; he was a non-smoker, with no history of dyslipidaemia, diabetes mellitus, coronary artery disease or other significant illness, and denied previous allergy, rhinitis, bronchial asthma, dermatitis or eczema. On arrival he was conscious, well oriented, afebrile and slightly cyanotic. His blood pressure was 105/60 mmHg, and the pulse was 90/min regular. Auscultation of the lungs and heart revealed normal breathing sound, rhythmic heart action with clear sounds. He had a erythematous rash on the left foot with a mild edema. Examination of other organ systems showed no abnormality.

An electrocardiogram performed immediately revealed sinus rhythm, heart rate was 89/min, with 1-1.5 mm ST-segment elevation in leads II, III, aVF, 2 mm depression of ST-segment with biphasic T wave in leads V2 and V3 (Figure 1). With the diagnosis of ST-segment elevation myocardial infarction he was transferred to Coronary Unit.

Initial laboratory analysis showed rise in Troponin I levels (34.68), CK-MB (49.3), markers of inflammation (leukocytosis (WBC=17.9*10^9 /L) with predomination of eosinophils – 15%), CRP – 65 mg/L; fibrinogen – 5.879 g/L), mild hyperglycemia (10.2 mmol/l). All other laboratory findings were inside the reference range. Echocardiography revealed mild left atrial and ventricle dilatation (LA 43 mm, EDDLV 60 mm; ESDLV 44 mm), severe hypokinesis of posterior wall, mild hypokinesis of basal segment of inferior left ventricle wall and mild hypokinesis of basal segment of ventricle septum. Due to a suspicion on Kounis syndrome we consulted an allergologist who suggested additional laboratory analysis (histamine, immunology with the accent on IgE antibodies, complement component C).

Patient received dual antiplatelet therapy and sent to primary PCI. Coronarography revealed significant stenosis of distal right coronary artery (RCA) branch – posterolateral artery (PLA) (Figure 2). During the procedure aspiration of fresh thrombus was made and extracted thrombus was sent to PH analysis with hematoxilin-eosin, which revealed the presence of eosinophils.
Results of additional laboratory parameters confirmed the presence of allergic reaction (histamine – 2.7 ng/mL; with IgE and complement component C above normal range).

During the hospitalization we administrated dual anti-platelet therapy, anticoagulant therapy, beta blockers (after normalization of blood pressure), H1 inhibitors, corticosteroids (methylprednisolone). After stent was implanted there was a significant resolution of ECG changes, over 50%, followed by clinical improvement and diminishing the angina discomfort.

During the hospital follow-up he had a complete resolution in ECG changes (Figure 3) as well as decrease in cardio specific enzymes (troponin I – 0.260 μg/L; CK-MB – 23.1 U/L). Patient was discharged after 5 days with advice to attend 3 and 6-month follow-up controls.
DISCUSSION

Kounis syndrome was named after a Greek author, Kounis, who was first to describe the progress of allergic angina into acute myocardial infarction. In 1991 syndrome was described by Kounis and Zavras as „the coincidental occurrence of chest pain and allergic reactions accompanied by clinical and laboratory findings of classic angina pectoris caused by inflammatory mediators released during the allergic insult”. Later, in 1995, Kovanen investigated the specificity of coronary arteries in patients who died after acute myocardial infarction, and have found that they had more significant mastocyte degranulation in eroded or ruptured plaque area than in nearby cells. Constantinidis showed that even some common allergic reactions could provoke plaque rupture, which happened with our patient who experienced acute myocardial infarction after he was stung by honeybee.

In 1998, Braunwald, in his editorial, categorized allergic angina as a subgroup of dynamic coronary occlusive lesions induced by allergic reactions noting that vasospastic angina can be induced by „allergic reactions with mediators such as histamine or leukotriens acting on coronary vascular smooth muscle”. As previously described, blood histamine levels can be elevated in patients suffering from different types of ischaemic heart disease including Kounis syndrome.

Today, Kounis syndrome is described as the occurrence of acute coronary syndrome with mast cell activation induced by allergic or hypersensitivity and anaphylactoid reactions. It can manifest as unstable vasospastic or nonvasospastic angina, and even as acute myocardial infarction triggered by the release of inflammatory mediators. Our patient had a typical allergic reaction to honeybee sting followed by angina pain and development of myocardial ischaemia with histamine increase.

There are two types of Kounis syndrome described so far with a proposition to include the third variant in a current classification. Type I refers to patients without cardiovascular risk factors and with healthy coronary arteries in whom allergic insult triggers coronary vasospasm that causes chest pain, ischaemic changes on electrocardiogram and normal/elevated markers of myocardial damage. This can be manifestation of endothelial dysfunction or microvascular angina.

Type II occurs in patients with pre-existing coronary disease where mediators may induce coronary vasospasm with normal cardiac enzymes, or erosion/rupture of the atherosomatous plaque, resulting in acute myocardial infarction. We classified our patient into type II Kounis syndrome according to his findings. Inferior myocardial infarction was shown on ECG as ST-segment elevation in inferior leads and confirmed by hypokinetic inferior wall on electrocardiography. Severe hypokinesis of posterior wall was present because of the occlusion of posterolateral branch of right coronary artery.

Type III, that has been proposed in recent years, refers to patients with drug-eluting stent thrombosis with the presence of mast cells and eosinophils revealed with Giemsa ad hematoxylin-eosin staining.

We administrated therapy according to acute coronary syndrome protocol plus type I Kounis treatment (corticosteroids, H1 and H2 blockers, dual antiplatelet and anticoagulant therapy). Although beta blockers should be administrated carefully because they can exaggerate coronary spasm, we had to administrate them because of tachycardia and occasional PVCs in our patient.

Several causes have been reported as capable of inducing Kounis syndrome. These include some drugs, foods and insects stings. In the case of insect stings and bites, the venom or saliva contains proteins, peptides and vasoactive amines that can cause direct cardiotoxicity but also behave as allergens, with activation and degranulation of the mast cells, resulting in the release into the systemic bloodstream of a number of vasoactive mediators and proteases. The existence of mastocytes in heart tissue and their participation in the anaphylactic reaction that triggers tachycardia, coronary vasoconstriction, dysfunctional ventricular contractility and blockade of atrioventricular conduction was already demonstrated.
LITERATURA


