CORONARY ARTERY VASOSPASM SECONDARY TO TYPE I VARIANT KOUNIS SYNDROME: A CASE SERIES OF MEN. IS THE GENDER DIFFERENCES IMPORTANT?

Cuneyt Kocas¹, Ahmet Cagri Aykan², Okay Abaci¹, Gökhân Cetinkılı ¹, Sukru Arslan¹, Mustafa Yildız¹

Kounis syndrome is a well-known cause of acute coronary syndrome and more than 100 cases are reported with allergic reactions to various drugs, animal and insect bites, even with drug eluting stents and endovascular devices. In this paper we report five-patients with Kounis syndrome related to different drugs. The main characteristic of patients is given in Table 1. All of them were male and their age was from 18 to 35. All patients presented with ST- elevation myocardial infarction. Coronary angiography was performed in all patients and revealed normal coronary arteries. From history of allergic exposure, electrocardiographic, laboratory (Total IgE and tryptase levels) and angiographic findings the diagnosis was Kounis syndrome type I for all patients and was treated with oral antihistamines and prednisolone. Despite Type I KS is not associated with atherosclerotic risk factors and CAD all patients in our report and many patients in literature are male. Gender differences in KS should be investigated in further studies.


Key words: Kounis syndrome, gender.

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ВАЗОСПАЗМ КОРОНАРНЫХ АРТЕРИЙ СРЕДНЕГО ТИПА И ВАРИАНТ КОУНИС СИНДРОМА: СЕРИЯ СЛУЧАЕВ У МУЖЧИН. ВАЖНЫ ЛИ ГЕНДЕРНЫЕ РАЗЛИЧИЯ?

Cuneyt Kocas¹, Ahmet Cagri Aykan², Okay Abaci¹, Gökhân Cetinkılı ¹, Sukru Arslan¹, Mustafa Yildız¹

Kounis-синдром является известной причиной острого коронарного синдрома и сообщается о более 100 случаях заболевания при аллергических реакциях на различные препараты, укусы животных и насекомых, даже на стенды с лекарственным покрытием и эндокаверкулярные устройства. В этой статье мы доказываем о пяти пациентах с синдромом Kounis, по отношению к различным лекарствам. Главная характеристика больных приведена в Таблице 1. Все они были мужчинами, их возраст от 18 до 35 лет. У всех пациентов был указан инфаркт миокарда с подъемом ST. Коронарная ангиография выявила у всех больных нормальные коронарные артерии. Из истории аллергического воздействия, эпикардиографических, лабораторных (Общий IgE и уровня тритазы) и ангиографических данных, был поставлен диагноз синдрома Кounis типа I для всех пациентов, которых лечили перорально антигистаминными препаратами и преднизолоном. Несмотря на то, что синдром Kounis типа I не связан с факторами риска атеросклероза и ИБС, все пациенты в отчетах, и многие пациенты, описанные в литературе, являются мужчинами. Гендерные различия синдрома Kounis должны быть изучены в ходе дальнейших исследований.


Key words: Kounis syndrome, gender.

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Introduction

The first paper in the literature, examining the acute myocardial infarction (MI) related to prolonged allergic reaction was published in 1950 [1]. But the definition of Kounis syndrome (allergic angina) was in 1991 as “the coincidental occurrence of chest pain and allergic reactions accompanied by inflammatory mediators released during the allergic insult” [2]. In 1996 “allergic myocardial infarction” term was defined in literature [3]. As today, Kounis syndrome is a well-known cause of acute coronary syndrome and more than 100 cases are reported with allergic reactions to various drugs (antibiotics, analgesics, antineoplastics, contrast media, intravenous anaesthetics, no steroid anti-inflammatory drugs, anticoagulants, proton pump inhibitors), animal and insect bites, even with drug eluting stents and endovascular devices [4, 5].

Case Descriptions

In this paper we report five-patients with Kounis syndrome related to different drugs. The main characteristic of patients is given in Table 1. All of them were male and their age was from 18 to 35. Main symptom was chest pain in all patients whereas dyspnoea was present in case 1 and pruritus was also accompanied in case 2. All patients presented with ST- elevation myocardial infarction, case 1.3 and 5 with anterior MI, case 2 inferior MI and case 4 with inferolateral MI. Troponin levels were increased in all patients, total immunoglobulin E (IgE) levels were also increased in all of them, tryptase level could be measured in two cases (Case 2 and 5) and elevated. Whole blood count, D-dimer, antithrombin III, serum cholesterol levels, C3 and C4 levels and antinuclear antibody, anti-DNA tests were within normal limits. The reasons of Kounis syndrome were metimazol sodium in case 1, gadopentetic acid in case 2, amoxicillin/clavulanic acid in case 3 and case 5, acetaminophen in case 4. The medical history for bronchial asthma, any allergic disease or coronary artery disease was negative. The serologic tests for viral aetiology were also negative. Coronary angiography was performed in all patients and revealed normal coronary arteries. From history of allergic exposure, electrocardiographic, laboratory (Total IgE and tryptase levels) and angiographic findings the diagnosis was Kounis syndrome type I for all patients and was treated with oral antihistamines and prednisolone. All symptoms electrocardiographic and echo-cardiographic findings were resolved by the time of discharge.
The characteristics of patients with Kounis syndrome

<table>
<thead>
<tr>
<th></th>
<th>Patient 1</th>
<th>Patient 2</th>
<th>Patient 3</th>
<th>Patient 4</th>
<th>Patient 5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, years</td>
<td>33</td>
<td>35</td>
<td>31</td>
<td>24</td>
<td>18</td>
</tr>
<tr>
<td>Sex</td>
<td>M</td>
<td>M</td>
<td>M</td>
<td>M</td>
<td>M</td>
</tr>
<tr>
<td>Atopy</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>CVRF</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Symptoms</td>
<td>Dyspnoea, chest pain</td>
<td>Pruritis, chest pain</td>
<td>Chest pain</td>
<td>Chest pain</td>
<td>Chest pain</td>
</tr>
<tr>
<td>Allergic cause</td>
<td>Metamizol 500 mg</td>
<td>Gadopentetic acid 20 ml</td>
<td>Amoxicillin/Clavulanic acid 1000 mg</td>
<td>Acetaminophen 500 mg</td>
<td>Amoxicillin/Clavulanic acid 1000 mg</td>
</tr>
<tr>
<td>ST segment elevation</td>
<td>Anterior</td>
<td>Inferior</td>
<td>Anterior</td>
<td>Inferolateral</td>
<td>Anterior</td>
</tr>
<tr>
<td>Trop I (ng/ml)</td>
<td>40</td>
<td>13</td>
<td>15</td>
<td>46</td>
<td>50</td>
</tr>
<tr>
<td>Total IgE (0–100 IU/ml)</td>
<td>190</td>
<td>117</td>
<td>120</td>
<td>202</td>
<td>350</td>
</tr>
<tr>
<td>Tryptase (5.6–13.5 µg/L)</td>
<td>-</td>
<td>20</td>
<td>-</td>
<td>-</td>
<td>45</td>
</tr>
<tr>
<td>CAG</td>
<td>Normal</td>
<td>Normal</td>
<td>Normal</td>
<td>Normal</td>
<td>Normal</td>
</tr>
</tbody>
</table>

Abbreviations: CVRF — cardiovascular risk factors, Trop I — troponine I, CAG — coronary angiography.

Discussion

Kounis syndrome was defined as an acute coronary syndrome that manifests as unstable vasospastic or nonvasospastic angina, and even as acute myocardial infarction triggered by inflammatory mediators following exposure to an allergic insult. There are currently 3 variants of Kounis syndrome [6]. The first variant is observed in patients with no cardiovascular risk factors and normal coronary arteries in whom acute release of inflammatory mediators such as histamine and leukotriens can trigger coronary spasm [7]. In Type-II variant of KS include patients with pre-existing atheromatous disease in whom acute release of these mediators can induce coronary artery spasm or rupture of atheromatous plaque. Recently, Type-III variant of Kounis syndrome has been defined in subjects with drug eluting coronary stent thrombosis [4]. In this variant stent components (nickel strut, polymer or impregnated drug) may play a role as allergic insult [8]. All of our patients were Type 1 Kounis syndrome. Several pathophysiologic mechanisms have been described to explain the association between an allergic reaction and acute coronary syndrome [6]. Mast cell degranulation follows after antibody antigen complex or sometimes allergen itself may start degranulation. After mast cell degranulation vasoconstricting and collagen degrading mediators such as, histamine, neutral proteases (tryptase, chymase), platelet activating factor and newly synthesized mediators are released locally and in the peripheral circulation [9]. These mediators appear to affect the myocardium directly. Histamine, through its H1 receptors, mediates coronary artery vasoconstriction, and increases vascular permeability, whereas activation of H2 receptors causes inotropic, chronotropic effects [10, 11]. Histamine can also activate platelets and potentiates aggregatory response. Tryptase level was elevated in two patients but not applicable in other patients. Total IgE levels were increased in all patients and this result may be the reflector of an allergic reaction. All patients in our report are male. CAD frequency is higher in male compared to female but Type 1 KS is not associated with atherosclerosis and defined as KS with normal coronary arteries. Although previous reports have shown mortality or malignant course of Kounis syndrome [4], our patients had an uneventful in-hospital course.

Conclusion

Although the exact pathophysiologic mechanism of Kounis syndrome is not clear, the increasing number of reports published in the last few year’s points that this syndrome should be kept in mind for consideration in the differential diagnosis of ischemic heart disease especially in young patients without cardiac risk factors and concomitant allergic symptoms. Gender differences in KS should be investigated in further studies.

References