The aim of this study was to evaluate the diagnostic accuracy of ischemia modified albumin (IMA) alone, or in combination with cardiac troponin T (cTnT) and electrocardiogram (ECG) findings for diagnosis of acute coronary syndrome (ACS). The study included patients with acute chest pain suggestive of ACS, recruited within 6 hours from onset. Patients were classified in ACS group and non-ischemic chest pain group (NICP). Of 84 patients, 49 were diagnosed with ACS and 35 with NICP. IMA was significantly higher in ACS group (p<0.0001). The area under receiver operating curve for IMA in ACS diagnosis was 0.95 (p<0.0001). Sensitivity and specificity of IMA for ACS diagnosis were 89.8% and 91.4%, respectively. IMA significantly (p<0.05) improved the sensitivity of ECG and cTnT, alone, and in combination. Sensitivity and negative predictive value of combination of IMA, ECG and cTnT for diagnosis of ACS were 100%. IMA is useful for diagnosis of ACS, in combination with ECG and cTnT.

Keywords: ischemia modified albumin; acute coronary syndrome; sensitivity; negative predictive value.

INTRODUCTION

The diagnosis of acute coronary syndrome (ACS) is currently based on evaluation of symptoms, electrocardiographic (ECG) findings and determination of markers of myocardial necrosis. An ACS may occasionally occur in the absence of ECG changes or elevations of biochemical markers, when the diagnosis is supported by the presence of previously documented coronary artery disease or subsequent confirmatory investigations [1]. ACS encompasses a wide spectrum of pathologic conditions, ranging from unstable angina (UA) to transmural acute myocardial infarction (AMI). Although these are different clinical presentations of ACS they usually share a common pathophysiologic substrate. In most conditions, the basic pathophysiologic mechanisms of ACS include atherosclerotic plaque rupture or erosion, with differing degrees of superimposed thrombosis and distal embolization, resulting in myocardial hypoperfusion [2].

Markers of myocardial necrosis have particular kinetics, thus can be detected in the blood 4-6 hours after insult. However, they can poorly discriminate UA from AMI patients. ECG changes are present at admission in about 50% of patients with ACS, and although widely used neither ECG findings nor determination of markers of myocardial necrosis can serve as a standard for the diagnosis of AMI. There is a constant need to establish early and sensitive markers of cardiac ischemia that will be able either alone or in combination with existing diagnostic tools to identify patients with ACS. Such marker could serve for initial identification of patients with ACS, followed by subsequent confirmation of the diagnosis of AMI versus UA using a sensitive and specific test for the presence of myocardial necrosis.

Albumin, with N-terminal part damaged or occupied with copper is referred to as ischemia modified albumin (IMA). One of the prominent features of IMA is inability to bind transition metal ions, such as cobalt. Bar-Or first reported a reduced binding of divalent cobalt ion added in vitro to human serum albumin in patients with ACS. On that basis, the test was developed in a prototype form, originally known as "assay for cobalt-albumin binding". Upon addition of cobalt solution to the serum, its attachment to albumin is measured using a colorimetric indicator dithiothreitol (DTT). In healthy individuals, the test was associated with less cobalt left to react with DTT. In patients with myocardial ischemia cobalt is not adhering to the N-terminal part of albumin, and more free cobalt is available to react with DTT [3].

STUDY OBJECTIVE

The aim of this study was to evaluate the diagnostic accuracy of IMA, alone and in combination with cardiac troponin T (cTnT) and ECG in patients presenting within the first 6 hours from the onset of acute chest pain suggestive for ACS.
MATERIAL AND METHODS

SUBJECTS

Patients admitted to the Coronary Care Unit, Medical Center, Kosovska Mitrovica, within 6 hours from the onset of acute chest pain suggestive of ACS were prospectively included in the study. Exclusion criteria were acute renal failure, peripheral vascular disease, congenital heart disease, autoimmune diseases, acute mesenteric ischemia, primary cardiomyopathy, brain ischemia, acute infection and albumin concentration outside of reference interval. All patients provided informed consent before participating in the study. This research was conducted following the tenets of the Declaration of Helsinki and approved by the institutional review board of Medical Faculty Pristina (Kosovska Mitrovica).

The diagnosis of ACS was based on the Joint European Society of Cardiology/American College of Cardiology Committee (ESC/ACC) guidelines [2, 4 - 6]. The ACS group consisted of patients with persistent ST-segment elevation (STEMI), non-ST segment myocardial infarction (NSTEMI) and UA. The NSTEMI was defined as significant ST-segment depression and T-wave abnormalities and elevated cTnT. The UA was defined as normal cTnT but with significant ST-segment and T-wave abnormalities. This subgroup also included patients with normal or undetermined ECG and normal cTnT, whose clinical presentation was suggestive for myocardial ischemia, supported by previously diagnosed coronary artery disease or subsequent confirmatory investigations. The control group was consisted of patients with non-ischemic chest pain (NICP) in whom myocardial ischemia was finally excluded as the cause of symptoms.

Biochemical methods

The blood was collected at admission into the test tube without anticoagulant, and spun at 3000 rpm for 15 min to obtain serum for routine biochemical analyzes, including cTnT.

Measurement of IMA concentration

For determination of IMA we employed the method of Bar-Or, as described [3]. In brief, 200 µL of serum was gently mixed with 50 µL of cobalt chloride (Sigma, 7.7 mmol/L CoCl₂ x 6H₂O). After 10 minutes incubation at room temperature 50 µL of DTT (Sigma, 9.7 mm/L) was added, and after exactly 2 minutes the reaction was quenched by addition of 1.0 mL physiological saline solution. The intensity of maroon color was measured on UV/VIS spectrophotometer (Safas 2, Monaco) at λ = 470 nm against the sample blank in which cobalt solution was replaced with water. The results were reported in absorbance units (ABSU). The linearity of IMA assay was tested using commercial human serum albumin in a concentration range of 20.7 - 62.0 g/L. Within-run CV was assessed by analyzing ten replicates of a pooled human serum with ABSU close to optimal cut-off determined by ROC curve analyses. The same sample was also used for the between-day CV estimation over five consecutive days.

Other biochemical measurements

Serum cTnT was determined by electrochemiluminescence immunoassay on Roche Elecsys 2010, using Troponin T STAT third generation test (Roche Diagnostics, Mannheim, Germany). Concentration of cTnT > 0.03µg/L was adopted as manufacturer’s recommended criterion for diagnosis of myocardial infarction (MI). Serum albumin, glucose and lipid status were assessed on automatic analyzer Hitachi 902 (Roche Diagnostics, Mannheim, Germany).

Statistical methods

All data were analyzed using MedCalc 12.3.0.0 (MedCalc Software, Belgium). Data distribution and homogeneity of variance were tested by the Kolmogorov-Smirnov test. Continuous variables with normal distribution were presented as mean and standard deviation, or median and 95% confidence interval (CI) for median. Differences between groups were tested by Student’s t-test or Mann-Whitney test, where appropriate.

The receiver operating characteristic (ROC) curve was generated to evaluate the ability of IMA in diagnosis of ACS. We determined optimal cut-off value for IMA, as combination of the smallest number of false positive and false negative results. The optimal cut-off was in the same time the criterion value for maximal Youden index. Also, bootstrapped 95% CI was calculated for both the Youden index and corresponding criterion value.

The area under ROC curve (AUC) was calculated with 95% CI. IMA, cTnT and ischemic ECG changes were analyzed to determine sensitivity, specificity, positive and negative predictive value and likelihood ratio for diagnosis of ACS. A McNemar’s test was used to compare the sensitivity and specificity of individual variables and their combinations for diagnosis of ACS. The combination was positive when one or more variables were positive. The combination was considered negative when all variables were negative. The following findings were considered positive: cTnT>0.03 ng/mL, IMA-cut-off value (calculated from ROC curve), ECG changes suggestive for myocardial ischemia- significant ST-segment elevation and depression, T-wave inversion in at least two contiguous leads. In all other cases variables were regarded as negative. Correlation analysis was accomplished by calculation of Spearman’s rank correlation coefficient. Linear regression was used to determine the linearity of IMA assay. Statistical significance was set at p < 0.05.

RESULTS

The recruitment of patients in the study is reported in Figure 1. Of 114 eligible patients, 30 were excluded due to objective problems. Of remaining 84 patients, 49 were with final diagnosis of ACS (29 AMI, 20 UA) and 35 with NICP. In the UA group, a clinical diagnosis was established in 4 cases while in 16 cases the diagnosis was based on objective testing (dobutamine stress echocardiogram or exercise stress test).

Basic demographical and clinical characteristics of studied patients are presented in Table 1. There were no significant differences between groups in sex, cigarette smoking, diabetes mellitus and lipid status. Patients in ACS group were older than those in NICP group. Previous MI was more common in ACS than in NICP group.
Figure 1. STARD flow diagram presenting the recruitment of participants in the study

Figure 2. Receiver operating characteristic (ROC) curve of IMA absorbance units (ABSU) was derived for acute coronary syndrome versus nonischemic chest pain patients. Optimal cut-off value (in ABSU) is indicated in comparison to NICP group, the median serum IMA was significantly higher in ACS group (p<0.0001). Also, in AMI subgroup serum IMA was 0.762 ABSU (95% CI 0.687-0.913) and 0.674 ABSU (95% CI 0.534-0.695) in the NAP subgroup, and the difference was significant (p=0.033).

The ROC curve analysis of IMA in ACS versus NICP (Figure 2) revealed an area of 0.95 (95% CI, 0.879-0.986), that was highly significant (p<0.0001). Optimal cut-off value determined from ROC curve was 0.4847 ABSU. At that point the Youden index was 0.8122 (95% CI, 0.673-0.902). At a cut-off point of 0.4847 ABSU sensitivity and specificity were 89.8% (95% CI, 77.8-96.6) and 91.43% (76.9-98.2), respectively (Table 2). The positive predictive value was 93.62%, and the negative predictive value was 86.49%.

IMA showed good linearity in the range of albumin concentrations 20.7-62.0 g/L. The regression equation was as follows: IMA (ABSU) = 0.9875 - 0.0122 × albumin (g/L). The within-run CV was 4.1% at a mean IMA of 0.4954. The between-day CV was 3.6% at a mean IMA of 0.4864. The mean IMA was close to optimal cut-off value.

The diagnostic accuracy of IMA, ECG and cTnT, used alone and in combination, for diagnosis of ACS is presented in Table 2. In comparison to ECG, cTnT, alone or in combination, the sensitivity of IMA for diagnosis of ACS was significantly higher (p<0.05). The addition of IMA significantly (p<0.05) improved the sensitivity of ECG and cTnT, alone and in combination. When IMA was combined both with ECG and cTnT the sensitivity and negative predictive value for the diagnosis of ACS reached maximal 100%.

In comparison to NICP group, the concentration of albumin was significantly lower (p<0.0001) in ACS group (45.2±4.9 g/L) compared to NICP (49.8±7.9 g/L). All measured concentrations of albumin were within normal range. Correlations between albumin and IMA (rho=-0.05, p=0.71), IMA and cTnT (rho=-0.2, p=0.17), and albumin and cTnT (rho=-0.07, p=0.63) were not significant.

DISCUSSION

Despite of improvements in cardiac biomarkers and introduction of different diagnostic tools, the diagnosis of ACS still remains a significant challenge [5, 6]. For many reasons previously established diagnostic tools failed to recognize all ACS patients, as was also confirmed in our study. For example, ECG was shown to have low sensitivity (Table 2), while cTnT is by definition elevated in AMI. Thus, there is a constant need for new and more reliable cardiac biomarkers, of which IMA seems to be the most promising one.

Mechanisms involved in ischemia-induced changes of albumin molecule could be associated with exposure to endothelial and extracellular hypoxia, free radical damage, membrane energy-dependent sodium and calcium pump disruption, and free iron and copper ion exposure [7-9]. Conditions implicated in alteration of metal binding site of human serum albumin are known to occur in vivo within a few minutes after the onset of myocardial ischemia [9, 10], as was previously confirmed in patients undergoing percutaneous coronary interventions [11, 12]. The study of Bar-Or et al. has also shown that acetylation or deletion of one or more amino acids on the N-terminal tripeptide region results in the loss of albumin cobalt binding capacity [11].

In the current study we tested the ability of IMA to distinguish patients with ACS. In agreement with other studies [3, 11, 13] we have found a significant elevation of IMA values in ACS compared to NICP group. Furthermore, IMA was significantly higher in the subgroup of AMI than in UA, suggesting that IMA actually reflects the extent of myocardial ischemia. However, there was no significant correlation between IMA and cTnT, probably due to different marker kinetics during the development of ACS [11, 12]. On the other side, Sinha et al. reported higher IMA in UA than in AMI group [14].

We determined the optimal cut-off value of IMA for diagnosis of ACS from the ROC curve. Although this ap-
Despite that, IMA alone failed to identify all patients. Variations indicate analytical reliability of IMA test. Moreover, the finding of low within-run and between-day imprecision of IMA in the current study serum IMA expressed high sensitivity of 91.4%. Biomarker sensitivity may be of crucial importance. In personalized patient evaluation, each biomarker has specific kinetics, it is hard to make comparisons. Another limitation is the lack of subfollow-up samples for IMA measurement, which would allow assessment of diagnostic accuracy of IMA in different time points from the onset of event. Furthermore, IMA is barely specific marker of myocardial ischemia, and altered cobalt binding to albumin can occur following strenuous exercise, as well as cerebral and intestinal ischemia [21, 22]. Although patients with these conditions were excluded from the study, the results of IMA should be carefully interpreted, even in combination with other diagnostic tools.

**CONCLUSION**

To conclude, our results indicate that the diagnostic performance of IMA to distinguish ACS from NICP can be enhanced by combination with ECG and cTnT findings.

### Table 1. Basic demographic and clinical characteristics of patients

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>ACS (N=49)</th>
<th>NICP (N=35)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>65±10</td>
<td>48±11</td>
<td>-0.0001</td>
</tr>
<tr>
<td>Male (%)</td>
<td>53%</td>
<td>54%</td>
<td>0.643</td>
</tr>
<tr>
<td>Smoking (%)</td>
<td>32.6%</td>
<td>20%</td>
<td>0.025</td>
</tr>
<tr>
<td>Hypertension (%)</td>
<td>87.6%</td>
<td>68.6%</td>
<td>0.026</td>
</tr>
<tr>
<td>Diabetes mellitus (%)</td>
<td>32.6%</td>
<td>17.1%</td>
<td>0.179</td>
</tr>
<tr>
<td>Hyperlipidaemia (%)</td>
<td>34.7%</td>
<td>31.4%</td>
<td>0.935</td>
</tr>
<tr>
<td>Previous MI (%)</td>
<td>26.3%</td>
<td>5.7%</td>
<td>0.03</td>
</tr>
<tr>
<td>IMA (ABSU)</td>
<td>0.696 (0.676-0.787)</td>
<td>0.395 (0.349-0.434)</td>
<td>-0.0001</td>
</tr>
</tbody>
</table>

Data in acute coronary syndrome (ACS) and non-ischemic chest pain (NICP) groups are presented as mean value ± SD, median and 95% confidence interval (in parenthesis), or frequency (%). Differences between groups were calculated using t-test, Mann-Whitney U-test, or chi-square test, where appropriate.

### Table 2. Diagnostic accuracy of several biomarkers in diagnosis of acute coronary syndrome

<table>
<thead>
<tr>
<th></th>
<th>IMA (ABSU)</th>
<th>ECG</th>
<th>cTnT</th>
<th>ECG + cTnT</th>
<th>IMA + cTnT</th>
<th>IMA + ECG</th>
<th>IMA + cTnT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sensitivity</td>
<td>(77.8-96.6)*</td>
<td>(38.3-67.5)</td>
<td>(19.9-47.5)</td>
<td>(52.5-80.1)</td>
<td>(92.7-100)*</td>
<td>(83.1-98.7)*</td>
<td>(92.7-100)*</td>
</tr>
<tr>
<td>Specificity</td>
<td>91.4</td>
<td>91.43</td>
<td>100</td>
<td>91.43</td>
<td>80</td>
<td>88.57</td>
<td>80.0</td>
</tr>
<tr>
<td>Positive PV</td>
<td>93.6</td>
<td>89.7</td>
<td>100</td>
<td>91.7</td>
<td>87.3</td>
<td>92</td>
<td>87.5</td>
</tr>
<tr>
<td>Negative PV</td>
<td>86.4</td>
<td>58.2</td>
<td>83.5</td>
<td>66.7</td>
<td>96.6</td>
<td>91.2</td>
<td>100*</td>
</tr>
<tr>
<td>Positive LR</td>
<td>10.4</td>
<td>6.19</td>
<td>7.86</td>
<td>4.9</td>
<td>8.21</td>
<td>5</td>
<td>8</td>
</tr>
<tr>
<td>Negative LR</td>
<td>0.11</td>
<td>0.51</td>
<td>0.67</td>
<td>0.36</td>
<td>0.026</td>
<td>0.069</td>
<td>0</td>
</tr>
</tbody>
</table>

Specificity, sensitivity, predicted value (PV), and likelihood ratio (LR) were calculated in 84 patients for diagnostic accuracy of ischemia modified albumin (IMA), electrocardiographical finding (ECG) and cardiac troponin T (cTnT) alone, or in combination, in diagnosing acute coronary syndrome. Data are presented as percentage and 95% confidence interval (in parenthesis). *p<0.05 in comparison to ECG, cTnT, alone and in combination.
REFERENCES


ZNAČAJ ISHEMIJOM MODIFIKOVANOG ALBUMINA (IMA) ZA DIJAGNOZU AKUTNOG KORONARNOG SINDROMA (AKS) KOD PACIJENATA SA AKUTNIM BOLOM U GRUDIMA

Dragojević I.1, Kisić B.1, Mirić M.2, Puhalo Sladoje D.3.
1 Institut za Biohemiju, Medicinski fakultet Priština, Kosovska Mitrovica, Srbija
2 Institut za Fiziologiju, Medicinski fakultet Priština, Kosovska Mitrovica, Srbija
3 Medicinski fakultet, Univerzitet Istočno Sarajevo, Republika Srpska, Bosna i Hercegovina

SAŽETAK

Cilj istraživanja je bio procena vrednosti ishemijom modifikovanog albumina (IMA) zasebno, i u kombinaciji sa srčanim troponinom T (cTnT) i elektrokardiografskim nalazom (EKG), u dijagnozi akutnog koronarnog sindroma (AKS). Istraživanje je obuhvatilo pacijente sa akutnim bolom u grudima, suspektnim na AKS, od čije pojave do pregleda je proteklo manje od 6 sati. Pacijenti su klasifikovani u dve grupe, sa AKS i grupu sa bolom u grudima neishemijskog porekla. Istraživanje je obuhvatilo 84 pacijenta, 49 sa AKS i 35 sa bolom u grudima neishemijskog porekla. Vrednosti IMA su bile značajno veće u grupi sa AKS (p<0.0001). Polje ispod ROC (receiver operating characteristic) krive je bilo 0.95 (p<0.0001). Senzitivnost i specifičnost IMA u dijagnozi AKS je bila 89.8% i 91.4%. Kombinacija IMA sa EKG nalazom i koncentracijom cTnT, pojedinačno ili sa oba, je značajno povećala njihovu senzitivnost u dijagnozi AKS. Senzitivnosti i negativna prediktivna vrednost kombinacije IMA, EKG i cTnT u dijagnozi AKS je bila 100%. Primena IMA u kombinaciji sa EKG nalazom i vrednošću cTnT se pokazala korisnom za dijagnozu AKS.

Ključne reči: ishemijom modifikovani albumin, akutni koronarni sindrom, senzitivnost, negativna prediktivna vrednost.