Utility of serum ischemia modified albumin in the early diagnosis of acute coronary syndrome

Context: Coronary artery disease (CAD) is the leading cause of death. Early detection of myocardial ischemia can decrease morbidity and mortality due to CAD. Ischemia modified albumin (IMA) is a newer biomarker for early detection of myocardial ischemia in patients of acute coronary syndrome (ACS) as compared to cardiac troponin T (cTnT). Aims: The aim of this study was to test the utility of “IMA,” a new biomarker of myocardial ischemia, in the early diagnosis of ACS. Settings and Design: The cross-sectional study group consisted of 101 patients between the age group of 27 and 85 years having ACS and 100 control from a healthy population. Materials and Methods: Blood was collected from all the enrolled patients immediately after admission, and samples were analyzed for cTnT on the fully automated analyzer and IMA on a spectrophotometer. Statistical Analysis Used: Statistical analysis was carried out by calculation of the area under a curve using receiver operating characteristics (ROC) curve analysis for the IMA test in the 101 patients included in the study population. Results: ROC curve area for IMA was 0.89 (95% confidence interval 0.81-0.94). At the cut-off 95 U/ml, sensitivity, and specificity were 85% and 82% and positive predictive value and negative predictive value (NPV) were 88% and 78%, respectively. Conclusions: IMA is a useful ischemic marker for diagnosis of myocardial ischemia due to high sensitivity, and it also facilitates to rule out myocardial ischemia due to high NPV.

Key words: Cardiac troponin T, coronary artery disease, myocardial ischemia

INTRODUCTION

Acute coronary syndromes (ACSs) is a spectrum of disease including unstable angina, associated with reversible myocardial cell injury, Q-wave and non-Q-wave acute myocardial infarction (AMI), non-ST-segment elevation MI and ST-segment elevation MI with large areas of necrosis.[1] Myocardial ischemia occurs either due to a lack of blood flow resulting in a shortage of oxygen and accumulation of waste products in the ischemic tissue. This results in a number of biochemical changes in the cells and if continued, leads to irreversible damage and necrosis.[2] ACS is becoming more common in the developing world such that in India, cardiovascular disease is the leading cause of death in India.[3]

Every patient with acute chest pain presenting to Emergency Department should be identified and confirmed for ACS for accurate medical management.[4] Electrocardiography (ECG), troponin and creatine kinase-MB are conventional markers for diagnosis of ACS. These blood markers are extremely sensitive and specific for the identification of patients with myocardial necrosis, but their ability to identify patients with myocardial ischemia remains limited. ECG is rarely abnormal following transient myocardial ischemia, and the ECG at presentation is normal in approximately half of the patients with ACSs.[5] These highly specific markers might give negative results on admission but give positive results hours later because myocardial necrosis is time dependent. Thus, markers able to identify patients with myocardial ischemia without infarction might serve an important role in the clinical setting.[6] Only about 22% patients develop AMI out of total admission to the cardiac care center.[7]

Ischemia modified albumin (IMA) is a newer marker for the detection of myocardial ischemia, in which N-terminal amino acids of human serum albumin are modified by ischemia. The amino
terminal end (N-terminal) of the albumin molecule is a binding site for transitional metals such as cobalt, copper, and nickel. In the presence of ischemia, the binding capacity of N-terminal of human serum albumin decreases toward cobalt. The principle of this test is to measure unbound cobalt by colorimetric assay. An elevated concentration of IMA has, therefore, been anticipated as a marker of myocardial ischemic injury. This assay is reported to be positive within minutes of ischemia and remains so for up to several hours later, allowing detection before the development of myocardial necrosis.

**MATERIALS AND METHODS**

The study was undertaken from November 2009 to October 2010 after prior approval from hospital Ethical Committee. The cross-sectional study group consisted of 101 patients between the age group of 27 and 85 years, admitted in emergency department and in intensive cardiac care unit within 6 h of clinical signs and symptoms of ACS. The control group included 100 apparently healthy individuals (age range, 30-70 years) to determine the 95th percentile of a control reference population for the IMA test.

Patients had been diagnosed as ACS was depending upon signs and symptoms as well as ECG changes and status of cardiac troponin T (cTnT). The diagnosis of ACS was based on criteria defined by the Joint European Society of Cardiology/American College of Cardiology Committee.

A cross-sectional study was planned to select serum samples from two distinct patient populations: Group 1 (noncardiac chest pain): Group 1 included 39 out of 101 patients with acute chest pain but no subsequent evidence of myocardial ischemia, labeled as “nonischemic chest pain.” Group 2 (cardiac chest pain): Group 2 included 62 patients out of 101 patients with acute chest pain and evidence of early myocardial ischemia, including unstable angina and AMI. These were labeled as “ischemic chest pain.”

All the patients with renal diseases, liver cirrhosis, acute stroke, and lung cancer were excluded from this study.

Five milliliter of blood was collected from all the enrolled patients immediately after admission before any treatment was started, in a plain tube for estimation of cTnT and IMA. cTnT were estimated immediately after the separation of sera and then these tests sera were frozen at −20°C for IMA estimation at a later stage. Frozen samples were mixed thoroughly after thawing and re-centrifuged before analysis. Repeat freeze-thaw cycles were avoided to reduce deterioration of samples. The IMA testing was done in batches. All the enrolled patients had an ECG within 1 h of admission. ECG criteria used to support a clinical diagnosis of myocardial ischemia were ST segment elevation or depression of 1 mm or more or T wave inversion 2 mm or more in 2 contiguous leads.

**Albumin cobalt binding (ischemia modified albumin) test**

The colorimetric assay for ischemia modified serum albumin developed by Bar-Or et al. measures free cobalt in serum, after cobalt albumin binding has, takes place. N-terminal of albumin is modified by ischemia and decrease affinity toward cobalt, thus amount of free cobalt is an increase in the reaction mixture. The amount of cobalt bound-albumin and the intensity of the color formation are in an inverse relationship. The method of albumin cobalt binding assay (IMA) involved the addition of 200 µl of patient serum to 50 µl of a solution of 1 g/L cobalt chloride, followed by gentle mixing, and incubated the solution for 10 min. 50 µl dithiothreitol (1.5 g/L solution) was then added, mixed and further incubated for 2 min. Then, 1.0 ml of phenobarbitone buffer (pH-8.6) was added. The absorbance of assay mixtures was read at 470 nm with a ultraviolet/visible T60 UV/VIS Spectrophotometer (PG Instruments Limited, London). The blank was prepared similarly by serum and cobalt chloride without of dithiothreitol and absorbance was read at 470 nm on the spectrophotometer.

Albumin cobalt binding assay was standardized in the Department of Biochemistry by using different concentration of CoCl₂, ranging from 6 to 60.0 µg CoCl₂/ml. A standard curve was prepared in the range 6-60.0 µg CoCl₂/ml. One IMA unit was defined as “µg of free Co²⁺ in the reaction mixture per ml of serum sample.”

Calibrations were done before running each batch of samples, using freshly prepared CoCl₂ and dithiothreitol reagents. Before running each batch control was done using known patient sample from a previous batch, which had been frozen after the test. Imprecision statistics was determined from 20 replicate assays of an unknown sample. The coefficient of variation (CV%) was calculated using mean values and standard deviation (SD) of 20 replicates. CV% was 7.8% in this study.

**Cardiac troponin T assay**

Cardiac troponin T was measured quantitatively using a electrochemiluminescence immune assay based on electrochemiluminescence technology (fourth generation cTnT, Elecsys 2010, Roche, Mannheim, Germany). The lower detection limit of this assay is 0.01 ng/ml with a recommended diagnostic threshold of 0.03 ng/ml. Troponin T concentration ≥0.03 ng/ml was considered a positive result.

Receiver operating characteristics (ROC) curve was analyzed and calculation of the area under the curve was done for the IMA test in the 101 patients, included in the study population according to the method of Hanley and McNeil by using MedCalc statistical software Version 12.2.1.0 (MedCalc Software Mariakerke, Belgium). The optimum cut-off for the IMA test was selected from the ROC curve analysis to maximize sensitivity and specificity in the study population. The upper 95th percentile ACB Test value for apparently healthy individuals was calculated using parametric statistics.
RESULTS

A total of 101 patients admitted to Emergency Department were selected for the study in the age group of 27-85 years. Of these patients, 71 (70%) were male and 30 (30%) were female. The IMA values for the control reference population (n = 100) were between 60.3 and 104.5 U/ml (mean, 76.2 U/ml; median, 75.8 U/ml). The upper 95th percentile was 92.4 U/ml. When serum IMA levels in Group 2 (ischemic chest pain) were compared with Group 1 (nonischemic chest pain), it was found that serum IMA of Group 2 was between 68.9 and 137.2 U/ml (95% confidence interval [CI] 76-85 U/ml) with mean ± SD as 106.4 ± 13.3 U/ml. Whereas in Group 1 serum IMA level were between 53.7 and 109.6 U/ml (95% CI 103-110 U/ml) with mean ± SD as 80.7 ± 13.9 U/ml. The IMA levels in patients of chest pain with ischemia were significantly higher than patients of chest pain without ischemia and control population (P < 0.0001). In the Group 2, 54 (87%) samples were found to have increased IMA levels as compared to 32 (82%) samples in the Group 1 have normal IMA level, shown as a scattered distribution plot between Group 1 and Group 2 [Figure 1]. In Group 2, 26 out of 62 patients were troponin T negative at presentation, but actually they were troponin T positive when the test was repeated after 6 h.

Diagnostic values of IMA, ECG and cTnT together and alone are given in Table 1. When ROC curve was used to evaluate the value of IMA, the area under curve was 0.89 (95% CI 0.81-0.94) [Figure 2]. At the cut-off 95 U/ml, founded by ROC curve showed the maximum sensitivity of 85% and specificity of 82%. Positive predictive value (PPV) and negative predictive value (NPV) were 88% and 78%, respectively. When cTnT and ECG used alone showed the sensitivity of only 58% and 69%, respectively. Sensitivity and NPV of IMA, cTnT and ECG, when used in combination were 91%. Hence, IMA was able to diagnose more ischemic patient than ECG or cTnT. When IMA, cTnT and ECG used together for diagnosis for ACS, ROC curve analysis showed area under curve as 0.92 (95% CI 0.83-0.97), sensitivity and NPV of 91%, while specificity and PPV of 82% respectively [Figure 3]. These three biomarkers demonstrated the significantly greater value of sensitivity and NPV than individual parameters (P < 0.0001).

DISCUSSION

This study illustrated that IMA value was significantly higher in patients with ischemia chest pain as compared to control group and nonischemic chest pain group. In this study, cut-off value of IMA 95 U/ml, derived by ROC curve was relatively higher than 85 U/ml, because IMA was performed by manual method on

![Figure 1: Distribution of ischemia modified albumin value amongst patients of chest pain at presentation](image)

![Figure 2: Receiver operating characteristics curve for serum ischemia modified albumin level amongst ischemic and nonischemic patients](image)

![Figure 3: Receiver operating characteristics curve for serum ischemia modified albumin, cardiac troponin T and electrocardiography amongst ischemic and nonischemic patients](image)
spectrophotometer, while it was performed on fully automated chemistry analyzer in previous studies done by Sinha et al[14] and Collinson et al[16].

ROC curve analysis of IMA showed higher sensitivity (86%) and NPV (78%) than cTnT and ECG at cut-off IMA value of 95.1 U/ml were similar to the result of Sinha et al[14] and Takhshid et al[16] studies. If we decrease IMA cut-off value to 87 U/ml, it gives higher sensitivity (92%) and NPV (84%), but it will decrease specificity to 69%. In this study cut-off for reference population is 92.4 U/ml (95th percentile), but use of a cut-off of 95 U/ml was to minimize the number of false positive and false negative results. This was the reason for taking cut off 95 U/ml for this study.

Receiver operating characteristics curve analysis for IMA, cTnT and ECG together showed sensitivity and NPV of 91%, which was significantly greater than that of shown by IMA, cTnT and ECG alone. These findings were nearly similar to result of Sinha et al[14] and Takhshid et al[16] studies. Our study proved that IMA had high sensitivity for early diagnosis of ACS and high negative predictive to rule out cardiac ischemia in nonischemic chest pain patients, but the specificity of IMA was lower than cTnT in patients at presentation, and therefore IMA, can be used as diagnostic marker along with cTnT and ECG in ACS. If value of IMA and cTnT are <95 U/ml and <0.03 respectively with nondiagnostic ECG, NPV was 91% for ACS in this study. IMA, cTnT and ECG comprise an excellent combination for noninvasive tests. When the physician is trying to make a decision whether to send the patient home or keep him in the hospital, the information from these three tests in combination is very useful.

Limitation of this study was IMA test returns to baseline within 8-12 h after an ischemic event and many of the patients were excluded from this study due to late presentation.

CONCLUSION

From this study, it can be concluded that albumin cobalt binding test for IMA is easy and simple end point method that can be measured by spectrophotometer and can be easily established in a small scale laboratory. This study suggests that patients presenting with acute chest pain and have high IMA values should be assessed and followed up carefully, even if cardiac troponin and ECG are negative. When IMA, cardiac troponin, and ECG used in combination, clinicians can easily make a diagnosis of myocardial ischemia in patients presenting to Emergency Department. These three biomarkers could allow early discharge of nonischemic chest pain patients more safely and cost effectively, due to higher NPV of these biomarkers.

REFERENCES