Diagnostic Performance of One-hour Delta High Sensitive Troponin in Acute Coronary Syndrome

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ABSTRACT

Objective: To determine the impact of an increase in the level of high sensitivite (HS) troponin-I between the time of Emergency Department (ED) arrival and one-hour after arrival (0h-1h), and between the time of ED arrival and three hours after arrival (0h-3h), upon the diagnostic performance of HS troponin-I to correctly diagnose AMI.

Study Design: Descriptive, analytical study.

Place and Duration of Study: Fethi Sekin City Hospital, Elazig, Turkey from January to June 2019.

Methodology: Patients presenting with chest pain were included in the study. The delta HS Troponin I (0h-1h) and delta HS Troponin I (0h-3h) values of the patients were calculated as percentage changes. These values were statistically compared between those who were diagnosed with acute myocardial infarction (AMI).

Results: The evaluation was performed on 114 patients. Delta HS Troponin I levels were examined, a change of >20% was observed in 48 patients (42.1%) for the 0-1 hr interval. Of the 20 patients ultimately diagnosed with an AMI, 16 (80%) had such a change at this time (p < 0.001). For 0-3 hr interval, a change of HS Troponin-I of >20% was observed in 52 patients (45.6%); and 17 of 20 AMI patients with an AMI were detected by such a change (p < 0.001).

Conclusion: Delta HS troponin I value in the first hour was nearly as sensitive for detection of an AMI as was delta HS Troponin I third-hour value.

Key Words: Troponin, Myocardial infarction, Acute coronary syndrome.

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INTRODUCTION

Acute myocardial infarction (AMI) takes place on the top among deaths in the world.¹ The diseases that cause the oxygenation of myocardial cells to deteriorate are acute coronary syndrome (ACS), which can be defined as a whole; sudden cardiac death consists of the subtitles of unstable angina pectoris, ST segment elevated myocardial infarction (STEMI) and non-ST segment elevation myocardial infarction (NSTEMI).

Today, the diagnosis of AMI is made by symptoms, electrocardiography (ECG) and cardiac biomarkers.^{2,3} Cardiac biomarkers are used in patients without ST segment elevation and with ischemic symptoms on ECG.² Troponin T and Troponin I, known as cardiac troponins, are among the most widely used biomarkers.^{4,5} An increase in Troponin I from extra-cardiac tissues has not been reported.²

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Received: June 25, 2020; Revised: September 17, 2020; Accepted: September 30, 2020 DOI: https://doi.org/10.29271/jcpsp.2020.09.906 In recent years, highly sensitive (HS) types of troponins are preferred because they respond earlier. When the troponin value exceeds 99 percantil upper referance limit (99th URL), it is an indication that myocard injury has occurred.² If there is an increase or decrease in the troponin value, this is an indication that the damage is acute and may be referred to as AMI.² It is stated in the literature that a 20% change in serial troponin measurements can be used as an AMI finding.^{2,6,7} As for the time of serial troponin measurements, the recommended arrival and 3rd hour troponin value should be seen.^{7,8} This change in troponin, called delta (Δ), may be undetectable due to the nature of the troponin ascent curve, if sampling the pattern is done too early or too late.²

The objective of this study was to compare the changes in troponin values at the 1^{st} hour and 3^{rd} hour, according to the arrival troponin value in terms of diagnostic performance on AMI (delta HStroponin 0-1h/delta HStroponin 0-3h).

METHODOLOGY

Patients who applied to the Emergency Department with ischemic symptoms suggestive of chest pain or ACS were included in the study. Pregnant women, those under the age of 18 years, those who did not agree to participate in the study, those who applied to the Emergency Room within the last 24

hours, those having a history of coronary intervention in the last month (cardiac bypass surgery, percutaneous coronary angioplasty), those who were in the haemodialysis programme, those with first troponin value <18ng/I (<99thURL), ST segment elevation in their ECG, as well as patients who were evaluated by the cardiologist for AMI with the first troponin result were excluded from the study.

The patients in this study were evaluated by emergency physicians blinded to the study design. Blood was drawn 3 times for troponin value from patients whose ECGs were taken at the time of application (0h), 1st hour and 3rd hour. Blood from gel tube (BD vacutainer SST II) studied through chemiluminescent immunoassay two step immunoenzymatic (sandwich) method with ACCESS hsTnI (RefB52699) kit in the Unicel DXI 800 accuTnI (Beckman Coulter, Inc., Fulerton, CA) device in the hospital centre laboratory. The results were obtained through the hospital data processing system. All patients were evaluated by a cardiologist with ECG, blood results, and echocardiography. Patients diagnosed with cardiac pathology were hospitalised. According to the HS troponin I results of the patients, delta HS troponin I (0h-1h), delta HS troponin I (0h-3h) and delta HS troponin I (1h-3h) values were calculated as percentage change. These values were compared statistically between those who were diagnosed with AMI and those who were not.

The study was carried out in accordance with Helsinki Declaration after Ethics Committee's approval.

Variables in the study were analysed statistically using SPSS version 25.0 (IBM Corporation, Armonk, New York, United States) and PAST 3 (Hammer, Ø., Harper, D.A.T., Ryan, P.D. 2001. Paleontological statistics). Kolmogorov-Smirnov test and Shapiro-Wilk test were used to assess the conformity of the univariate data to normal distribution. In order to assess the multivariate normal distribution, Mardia (Dornik and Hansen omnibus) test was used. Mann-Whitney U-test was used in the comparisons of quantitative data of two independent groups. Friedman's Two-Way test was used in the comparison of two repetitive measurements of the dependent quantitative variables, and Dunn's Test was used for Post Hoc analysis. In the comparison of categorical variables with each other, Pearson Chi-square test was used. The odds ratio was used with 95% confidence intervals to indicate how many times higher the risk factor is compared with those not possessing a risk factor. In order to show the relationship between the categorisation created by the cut-off value which was calculated based on the variables - and the actual categorisation. The sensitivity, specificity and diagnostic accuracy likelihood odds were analysed and indicated by receiver operating characteristic (ROC) curve analysis. Neural Network (Multilayer Perceptron) was implemented both to determine the most important variable and for estimation as well. Gradient descent was used for optimisation algorithm, hyperbolic tangent was used as hidden layer activation function, and identity was used as output layer activation function. Furthermore, mini-batch method was used for training data selection, and it was set as 100% training set and 0% testing set. The quantitative variables were shown as median (25th percentiles (Q1)/75th percentiles (Q3)), while the categorical variables were shown as n (%) in the Tables. The variables were analysed at 95% confidence level, and the p-value was accepted as significant when it was lower than 0.05.

RESULTS

Between January and July 2019, 442 patients applied to the adult Emergency Department with a complaint suitable for the study. Sixty-four of these patients did not agree to participate in the study. Some patients were excluded from the study due to the various reasons as follows: Seven patients were on haemodialysis programme due to chronic renal failure; 9 patients left the emergency service at their own request while the treatment was continuing; 22 patients were diagnosed with STEMI at presentation or during follow-up; and 29 patients were considered AMI by the cardiologist with the first troponin result. The first troponin result of 197 patients was <18ng/I (99th URL). The final evaluation was made on 114 patients.

As a result of follow-up and consultations for 114 patients in the Emergency Department, 80 patients (70.2%) were discharged after their treatment was regulated. Twenty-six patients (22.8%) were hospitalised in coronary intensive care unit due to cardiac pathology, and 8 (7%) patients in other clinics. Twenty of the patients hospitalised in coronary intensive care had (17.5%) AMI, three patients had tachycardia, two patients had congestive heartfailure and one patient had pericarditis.

Of the 80 patients included in the study and discharged, 78 were reached by phone 7-11 days after their discharge. After discharge, the ACS and mortality histories of the patients were questioned. No ACS or mortality was detected in the patients reached. Two patients could not be reached due to missing contact information. It was checked through the national death reporting system, and no mortality was detected. There was no statistically significant difference by age and gender between those who were diagnosed with AMI and those who did not (Table I).

When delta HS troponin I levels were examined in 0-1-hour interval, a change of >20% was found in 48 (42.1%) of 114 patients; and 16 of these patients had AMI [p < 0.001, odds ratio: 7.8, 2.4-25.1)]. While 52 (45.6%) patients with delta HS troponin I >20% were detected between the range of 0-3 hour. In the AMI group, it was found to be 17 patients in this range, [p < 0.001, odds ratio: 9.6 (2.6-34.9, Table I]. Between 1-3 hours, while delta HS troponin I was >20% in 45 (39.5%) patients, this rate was found to be 17 patients in the AMI group, [p < 0.001 odds ratio: 13.4 (3.6-49.2, Table I).

The cut off values were determined as the most appropriate increase rate in troponin value for AMI diagnosis, as follows: delta HS troponin I >66.7% (sensitivity 75.0%, specificity 79.8%, odds ratio: 11.84) 0-3h delta HS troponin I >148.3% (sensitivity 80.0%, specificity 85.1% odds ratio: 22.86) in the range 1-3h delta HS troponin I >47.2% (sensitivity 80.0%, specificity 89, odds ratio: 33,59, Table II).

		Total (n=114) Median (Q1-Q3)	ACS Absent (n=94) Median (Q1-Q3)	ACS Present (n=20) Median (Q1-Q3)	Р
		n (%)	n (%)	n (%)	
Age		64 (53-74)	63 (52-72)	68 (56-79)	0.100 ^u
Sex Female Male		44 (38.6) 70 (61.4)	36 (38.3) 58 (61.7)	8 (40.0) 12 (60.0)	0.887 ^p
		Total (n=114) n (%)	ACS Absent (n=94) n (%)	ACS Present (n=20) n (%)	Р
delta hs Trop I (0-1)h $\pm 20\%$					
Non risc		66 (57.9)	62 (66.0) "	4 (20.0)	<0.001 ^p
Risc of ACS		48 (42.1)	32 (34.0)	16 (80.0)	7.8 (2.4-25.1) ^{or}
delta hs Trop I (0-3)h ±20%					
Non risc		62 (54.4)	59 (62.8) "	3 (15.0)	<0.001 ^p
Risc of ACS		52 (45.6)	35 (37.2)	17 (85.0)	9.6 (2.6-34.9) ^{or}
delta hs Trop I (1-3)h ±20%					
Non risc		69 (60.5)	66 (70.2) "	3 (15.0)	<0.001 ^p
Risc of ACS		45 (39.5)	28 (29.8)	17 (85.0)	13.4 (3.6-49.2) ^{or}
		Median (Q1Q3.)	Median (Q1Q3.)	Median (Q1Q3.)	
delta hs Trop I 1h-0h %		12.3 (1.7-91.4)	9.9 (1.4-43.0)	130.8 (53.5-335.2)	<0.001 "
delta hs Trop I 3h-0h %		18.2 (0.6-200.3)	14.8 (0.5-77.7)	820.5 (290.5-2056.5)	<0.001 "
delta hs Trop I 3h-1h %		8.4 (-4.3-46.1)	4.1 (-5.2-20)	133.5 (51.5-347.6)	<0.001 "
hs Trop I Oh ng/I (A)		24.2 (8.9-45.1)	22.5 (7.1-39.4)	46.3 (33.4-85.7)	<0.001 "
hs Trop I 1h ng/I (B)		33.1 (19.1-90.5)	30.5 (12.5-51.5)	153.6 (91.6-418.3)	<0.001 "
hs Trop I 3h ng/l (C)		39.1 (20.9-142.8)	30.6 (14.6-58.1)	387.9 (249.1-1059.2)	<0.001 "
P value for intra groups ^f		<0.001	<0.001	<0.001	
	A-B	<0.001	<0.001	0.027	
Pairwise comparation	A-C	<0.001	<0.001	<0.001	
	B-C	0.017	0.109	0.027	
" Mann Whitney u-test, ^P Pearson 25%, Q2.: Percentile 75%, (I, II) s	Chi-square shows mean	Test, ^{or} Odds Ratio (95% Confid ingfulness according to groups.	ence Intervall), ^f Friedman Te	st; Post Hoc Test : Dunn's Te	st , Q1.: Percentile

Table II: Cutoff variables determined according to the delta time at HS troponin I.

	Delta HS Trop I 1h-0h %	Delta HS Trop I 3h-0h %	Delta HS Trop I 3h-1h %				
	(A)	(B)	(C)				
Cut off	>66.7	>148.3	>47.2				
Sensitivity	75.0%	80.0%	80.0%				
Specificity	79.8%	85.1%	89.4%				
Accuracy rate	78.95%	84.21%	87.72%				
AUC±SE.	0.755 ± 0.072	0.809 ± 0.074	0.838 ± 0.066				
Р	<0.001	<0.001	<0.001				
Pairwise comparation for AUC	P (A-B)=0.291	P (A-C)=0.323	P (B-C)=0.613				
Odds Ratio (SE.)	11.84 (0.70)	22.86 (0.77)	33.59 (0.81)				
Roc (Receiver Operating Curve) Analysis (Honley&Mc Nell - Youden index I) AUC: Area under the ROC curve SE: Standard Error							

Roc (Receiver Operating Curve) Analysis (Honley&Mc Nell - Youden index J) AUC: Area under the ROC curve SE: Standard Error

Table III: Neural network models for prediction of AMI.

Model	Variable Import	Sample (Holdout)	Predicted AMI			
	Independent Variable	Normalized Importance	Training (%100)	Absent	Present	Percent Correct
1	Delta hs Trop I (0-1)h ±20%	75.4%	AMI Absent	94	0	100.0%
	Delta hs Trop I (0-3)h ±20%	72.1%	AMI Present	20	0	0.0%
	Delta hs Trop I (1-3)h ±20%	100.0%	Overall Percent	100.0%	0.0%	82.5%
2	Delta Cut off 1h-0h %	39.0%	AMI Absent	87	7	92.6%
	Delta Cut off 3h-0h %	38.6%	AMI Present	5	15	75.0%
	Delta Cut off 3h-1h %	100.0%	Overall Percent	80.7%	19.3%	89.5%
Neural netwo Dependent	ork (Multilayer Perceptron, Radial Basis /ariable: AMI	Function), Hidden layer a	ctivation function: Hyperboli	c tangent, Outpu	t layer activation	function: identity,

In addition, in the neural network modelling (Model 1) made, based on the change in delta HS troponin I levels of > 20%, none of the patients with AMI could accurately be predicted. However, with a model created according to the cut-off values (Model 2), patients with AMI could be predicted with 75% accuracy and 89.5% in total. In both models, normalised importance was found at the maximum delta value between 1-3h (Table III, Figure 1).



Figure 1: Model 2 of the neural network.

DISCUSSION

AMI is classified as type 1, if it develops due to atherosclerotic heart disease; and type 2, if it develops due to the mismatch between oxygen supply and demand.² The distinction of these two is clinically difficult. While an invasive approach is not required for most patients in type 2 MI, an invasive approach is required primarily in type 1 MI. At this point, the amount of increase in high sensitivity troponins is important. If the increase in troponin value is excessive, first of all, invasive approach should be considered.⁹

While it is emphasised in the guidelines that at least two serial measurements should be made.⁸ It is stated that troponin measurement intervals can be increased to 3 hours, and up to 6 hours in suspected patients.⁸ However, with the advancing technology and the increasing use of highly sensitive troponin, studies are being conducted to further shorten these periods.¹⁰⁻¹⁴

Given that cellular ischemia results in cellular dysfunction within 10 minutes of the onset of an AMI, early and accurate diagnosis has a high clinical importance. Accurate diagnosis enables rapid identification of patients who merit an emergency invasive revascularisation procedure, performed with the intention of preserving myocardial function and avoiding subsequent problems such as sudden cardiac death or congestive heart failure. Early identification also could decrease the ED length of stay, and thus improve ED throughput times and efficiency.¹⁰

Although there are delta troponin studies up to 30 minutes

in the literature, 2015 ESC manual, under certain conditions, contains information about rapid discharge regarding 1 hour delta troponin levels.^{1,15} In literature in similar studies, patients below the level of detection value were examined to quickly exclude the diagnosis of AMI.¹⁶⁻²⁰ The authors investigated for the faster diagnosis of patients with HS troponin I>99th URL at presentation.

For the diagnosis of AMI, authors compared the percentage change in HS troponin I at the 1st hour with the percentage change at the 3rd hour (diagnostic performance). In patients diagnosed with AMI determination was done as follows: 16 patients (80%) with delta HS troponin I (0-1h) >20% (p <0.001 odds ratio: 7.8 (2.4-25.1)), delta HS troponin I (0-3h) >20% in 17 patients [85%, p <0.001, odds ratio: 9.5 (2.6-34.9)].

However, the highest odds ratio was found as delta HS troponin I (1-3h) >20% (p <0.001 odds ratio: 13.3 (3.6-49.2)). The reason of this might be the variation in the perception of symptoms by persons, which can be very subjective. Therefore, the symptom duration of the patients was not explained in this study. Troponin value starts to increase with the onset of symptoms. However, due to troponin kinetics, when sampling too early or late, the delta value may not be detected.²

Delta troponin values in the range of 1-3h in the presently reported two case groups corresponded to the rising period in the troponin kinetic curve.

According to the cut-off value, the maximum diagnostic accuracy is delta HS between 1-3 hours troponin I >47.2% had a sensitivity (80.0%) and specificity (89.4%) with AUC of 0.838 \pm 0.066 (p < 0.001). Secondly, delta HS between 0-3 hours troponin I >148.3% with sensitivity of 80.0%, specificity of 85.1%, AUC 0.809 ± 0.074 and p < 0.001. Finally, delta HS troponin I >66.7% has a sensitivity of 75.0% and specificity of 79.8% at 0-1-hour interval, AUC 0.755 ± 0.072 (p <0.001). In the study conducted by Keller *et al.*¹¹ in patients with HS troponin I level >99th URL at the time of admission; when the third hour cut-off value was >250%, they determined the specificity as 99.6, sensitivity 32.6, and PPV 95.8.¹¹ Although the diagnostic power was greatest at the delta HS troponin I (1-3h) change; the delta HS troponin I (0-1h) change and the delta HS troponin I (0-3h) changes are very close in terms of diagnostic power. For this reason, one should consider acting earlier, rather than waiting 3 hours for the second troponin sample, as stated in the literature in patients with a high probability of ACS clinically.⁸ At this point, the patient's history, age and risk factors are very important. However, do not forget the other reasons that increase troponin values in low changes or the factors that cause type 2 MI.⁹ When a statistically neural network study was done, the model established based on >20% change in troponin value could not detect any of the patients with AMI. The second model, built on the basis of cut-off values,

detected 75% of patients diagnosed with AMI and achieved an accuracy rate of 89.5% in all patients.

Although the low number of patients with AMI diagnosis is considered as the main limitation, the time between the onset of symptoms and admission to the hospital should be clearly demonstrated in order for such studies to yield better results. The authors did not explore the symptom duration in the study because of possible recall bias.

A relative value was used as the delta troponin value. In studies comparing absolute and relative delta troponin in the literature, although both were effective in the diagnosis of AMI, absolute u was found to be statistically superior.^{14,21} Using relative and absolute delta troponin values together in this study could have shown a more detailed approach.

CONCLUSION

In AMI, the initiation time of treatment is closely related to mortality and morbidity. The delta HS troponin I 0-1h will be as useful as delta HS troponin I 0-3h in diagnosing NSTEMI. It is noted that a change of >20% in the HS troponin value specified in the manuals will not be sufficient in future models of artificial neural networks, but the models to be established with a higher rate of change will be effective in confirming the diagnosis of AMI.

ETHICAL APPROVAL:

This research was carried out after obtaining permission from the Ethics Committee of the Firat University, Elazig, Turkey.

PATIENTS' CONSENT:

Consents were obtained from all patients.

CONFLICT OF INTEREST:

Authors declared no conflict of interest.

AUTHORS' CONTRIBUTION:

BC: Analysis and interpretation of data and critical revision. SS: Study conception and design, acquisition of data and drafting of manuscript.

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