TROPONIN T AND
HISTOLOGICAL
CHARACTERISTICS
OF RAT MYOCARDIAL
INFARCTION INDUCED
BY ISOPROTERENOL

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ABSTRACT

In our investigation, we used short-time model of myocardial infarction of rats induced by high dose of isoproterenol (ISP). We investigated cardiac troponin T blood level (cTnT) and histological characteristics of rat myocardium. ISP, single, intraperitoneal dose 250 mg/kg was given to male, adult, Wistar rats (n=12). Rats were distributed depending on their body weight in subgroups: ISP I (BW 260-280g) and ISP II (BW 250-400g). Control group (n=9) was treated with intraperitoneal dose of 0.95% NaCl. Cardiac TnT was measured by electrochemiluminescence (ECLA) sandwich immunoassay in rat serum 4 hours after ISP application. Rats’ hearts were dissected and examined by qualitative histological method (HE). Statistical significance was set at 0.05. There was significant difference in cTnT of ISP II (p<0.0001) vs. control and ISP I (p<0.05) vs. control. Significant difference was between ISP I and ISP II subgroups (p<0.001). The accent of histological changes of myocardium was on nuclei of cell. Cells showed acydophilic changes and nuclei disappearance as signs of coagulative necrosis development. Extensivity of histological changes were different between ISP I and ISP II subgroup. Used dose of ISP induced development of myocardial necrosis in rats. Subendocardial portion of myocardium was more vulnerability than subepicardial portion. Rats of ISP II had more extensive histological changes than these in ISP I. Administered doses of ISP enabled cTnT utilization as a marker of myocardial necrosis.

KEY WORDS: isoproterenol, cardiac troponin T, myocardial necrosis, histology
INTRODUCTION

We’re witnesses of permanent changes in diagnosing and therapy of acute myocardial infarction patients. Permanent changes are result of experimental investigations and better understanding of molecular mechanisms occurred during myocardial necrosis development. Myocardial infarction can be induced chemically and non-invasively in small laboratory animals like rats. Commonly used non-invasive techniques for induction of rat myocardial necrosis are those with use of catecholamines. For this purpose, isoproterenol-synthetic catecholamine is the most often used. It’s β-adrenergic receptors agonist and causes severe stress in myocardium resulting in infarct-like lesion. Rona and coworkers published the first results about ISP cardiotoxic effects (1). Rat myocardial changes induced by ISP are a similar to human myocardium changes during myocardial infarction (2). ISP produces a relative ischemia or hypoxia because of myocardial hyperactivity, coronary hypotension and cytosolic Ca\(^{2+}\) overload. There is evidence that ISP cardiotoxicity is result of catecholamine oxidation into aminochromes (3). Degree of pathomorphological changes depends on used ISP dose (4). Changes are present in subendocardial portion of myocardium, apex, left ventricle, papilar muscle and close to coronary artery. Troponins are highly sensitive and specific markers of cardiac cell damage (5). These contractile proteins are released from myocardium in proportion to the degree of tissue injury. Cardiac troponin T (cTnT) belongs to the proteins of contractile apparatus that are unique for cardiac muscle. Except in myocardial infarction diagnosing, troponins should be included in the evaluation of cardiotoxicity and cardioprotective properties of new drugs (6). There is so little information about cTnT blood level and histological finding of myocardium in ISP induced rat model of myocardial infarction. There isn’t well-standardized short time animal model of rat myocardial infarction induced by ISP for studying of: a) histological and cardiac marker changes during myocardial infarction, b) cardioprotective and c) cardiotoxic drugs investigation. In this short time rat model, we examine histological characteristics of cardiac muscle damage and TnT blood level in rats 4 hours after ISP administration.

MATERIAL AND METHODS

We used twelve, adult, male, Wistar rats as experimental group. They were raised and housed in air-conditioned, humidity-controlled cages. Rats had free access to water and commercial food during experimental period. Ethical Committee of our Institution approved the experiment. Rats were distributed depending on their body weight in subgroups: ISP I (BW 260-280g) and ISP II (BW 250-400g). Rats of both subgroups were treated with single intraperitoneal (i.p.) dose of ISP (250 mg/kg BW). Control group (n=9) was treated with intraperitoneal dose of 0.95% NaCl. Ketamin anesthesia (0.1 ml/100g BW) was performed in rats before of ISP administration. Isoproterenol hydrochloride was manufactured by Sigma Chemical Company, USA. Rats of ISP II died in different intervals from ISP application within 4 hours. Blood of these animals was taken by cardiac punction. Rats of ISP I survived 4 hours of experimental period and blood samples were drawn from tail wein. The blood was centrifuged for 10 minutes at 4000 r.p.m. The sera were frozen and stored at –20° until determination. The determination of cTnT was performed with Elecsys, electrochemiluminiscence (ECLIA) sandwich immunoassay, manufactured by Roche Diagnostics. We used analyzer Elecsys 2010, Roche. Values of cTnT are given in ng/ml. The chest cavities of the rats were opened to remove the heart shortly after blood samples were taken. Rats’ hearts were dissected for histological examination. Left ventricular tissue was placed in 10% buffered formalin solution, embedded in paraffin, sectioned at 5 μm intervals and stained with hematoxylin-eosin (HE). Histological analysis was obtained by using microscope Nikon type 400E with installed digital camera. Results of histological analysis are presented by using qualitative histological analysis. Cardiac TnT blood level data were analyzed by two-tail, unpaired Student’s t-test. Significance was set at 0.05. Results are reported as mean ± SD.

RESULTS

Control rats were treated with saline and 4 hours after that blood values of circulated cTnT were minimal.
Mean value was 0.01 ng/ml. Four hours of experimental period after ISP application, rats of ISP I subgroup were survived and ISP II died in different intervals. Compared to control rats, cTnT blood level of ISP treated rats was significantly higher (p<0.0008). Mean value was 55.97 ng/ml (Figure 1).

Significance of difference of cTnT level in ISP I vs control was p<0.05. ISP II had difference at higher level of significans p=0.0001. Mean values of cTnT in ISP I were 15.62 ± S.D.14.44 ng/ml and for ISP II subgroup TnT was 84.80 ± 26.23 ng/ml. There was significant difference between ISP I and ISP II subgroups ( p<0.001) (Figure 2.).

Figure 3. (A and B) presents normal myocardial tissue of control rats. The myocardium is composed of muscle fibers. Each fiber is enclosed by a sarcolemma. The nuclei are generally located in the central portion of the fiber. A net of reticular fibers and fine collagenous fibers surround each muscle fiber. The myocardium is richly supplied with small vascular channels forming an intramural circulation.

Subendocardial myocardium of ISP I rats is presented with areas of necrotic changed myocytes. Myocytes show variable degree of damage and accent of pathological changes are on nuclei. Cells are edematous, lost of the normal striation pattern and posses nuclei with inhomogeneous content (Figure 4.). Deposited material of hyperacidophilic features are presented close to sarcolemma (4.B). In necrotic myocytes myofibrillar lysis is complete. Deep subendocardial portion shows diffuse arranged focal necrosis of myocytes. Nuclei of these areas are in phases of picnosis or lysis or completely disappearance. Blood vessels near necrotic area contain eosinophilic amorphic material. Subepicardial myocardium of ISP I not deviate from normal myocardial tissue of control rats.

Rats of ISP II died in different intervals from ISP application within 4 hours. Large blood vessels involved in subendocardial tissue are strong dilated (Figure 5). Histological changes were more extensive compared to ISP I. Subendocardial portion is presented in form areas of massive necrosis, cellular arrangement disappeared, nuclei disappeared or in form of massive vacuola. In contrast to ISP I, rats of ISP II shows changes in subepicardial myocardium. Variable degree of nuclei cariolysis in subepicardial myocytes is presented.

**DISCUSSION**

In this study we used rat model of acute myocardial infarction induced by ISP. Based on Rona and coworkers experiences, it’s known that ISP has harmful influence on heart. O’Brien and associates (7) have shown that troponins are a powerful biomarker in laboratory animals for sensitive and specific detection of cardiac injury arising from various causes. From earlier ISP stud-
ies in rats, it is known that high doses of ISP (4-80 mg/kg) induce acute myocardial damage that has consequence in increasing of blood cTnT (8). We knew that cTnT determination with the cTnT assay, which was developed originally for human sera, was possible in the rat with the monoclonal antibodies used, because of cross-reaction between humans and rats cTnT (9). Very small amount of cTnT circulates as result of natural protein turnover. Minimal circulating cTnT values were obtained in our control rats. Detection of circulating cTnT depends on assay sensitivity. Control rats had TnT mean value 0.01 ng/ml. Intraperitoneal, single-ISP dose, administered to rats caused expectant, significant increasing cTnT. From damaged myocites, first is released free citosolic pool of cTnT. After that portion, cTnT bound in contractile apparatus is released by proteolytic degradation. We get significant difference in cTnT blood levels between ISP II and I subgroups (p<0.001) as consequence of difference in extensivity of histological changes. These results are in accordance

**FIGURE 4.** The structural changes of myocardium induced by ISP ISP I rat (A, B) Necrotic changed cardiomyocytes in subendocardial myocardium (C) Perivascular changes of myocytes are more extensive (D) Deep subendocardial portion of myocardium with focal necrosis. Edematous myocytes are presented with nuclei in phases of picnosis, lysis or completely disappearance (A & B X100, C, D X400).

**FIGURE 5.** Histological characteristics of myocardium induced by ISP ISP II rat (A) dilated and injected large blood vessel (B) massive necrosis of myocytes; cellular arrangement disappeared (C) nuclei cariolyis of variable degree to completely disappearance.
to obtained results of qualitative histological analysis. Degree of myocardium pathomorphological changes depends on used ISP dose (10). Subcutaneous injection of ISP 10 mg/kg causes rat myocardial necrosis due to prolonged tachycardia. The proposed mechanism for ISP myocardial necrosis are myocardial hypoperfusion, glycogen depletion, electrolyte imbalance, lipid accumulation and free radical damage (11,12). According to Preus and coworkers, 6 hours after ISP application, cells showed acyrophilic changes and nuclei disappearance as signs of coagulative necrosis development (13). There aren’t published data about using higher doses of ISP and expressed myocardial changes in first hours after application. By using ISP 250 mg/kg dose, we noted extensive myocardial lesions 4 hours from drug application. Results of histological analysis gave us clear evidence of necrogenic effect of ISP on heart. Coagulative necrosis induced by ISP, which we found in our study, is the same in case of heart damage caused by ischaemia and presented in human myocardial infarction. Massive myocardial necrosis characterizes human myocardial infarction while catecholamine model of myocardial necrosis is presented in form of focal coagulative necrosis. Joseph and Balasz described focal and multiple areas of myofibrillar necrosis after ISP application to rats (14). Our results are in concordance with their findings. Histological examination of the myocardial tissue in all ISP treated rats showed necrotic areas in the subendocardial layer of the left ventricle. Poorer vascularisation and oxygen supply of subendocardial portion is possible cause of more vulnerability. By qualitative histological analysis, we noted the difference in extensivity of myocardial changes between ISP I and ISP II rats. Rats of ISP II had higher body weight (mean BW 325 g) than ISP I rats (mean BW 270g). ISP I subgroup have had changed subendocardial portion of myocardium, thus subepicardial portion didn’t show any changes. Lesions were presented in form of widespread focal necrotic areas. Cardiomyocytes were most frequently without nuclei, lost of the normal myofibrilar striation pattern. There were variable degrees of damage. Rats of ISP II had expressed myocytes changes in subepicardial and subendocardial portion of myocardium. Subendocardial alterations were more extensive than in ISP I subgroup. There were signs of massive necrosis with disappeared cellular arrangement and lost edematous and fragmented myocytes. Large blood vessel in myocardium of ISP II group were dilated greatly and injected which present the sign of shock development and myocardial hypoperfusion. Cells’ nuclei have disappeared or presented in form of massive vacuola. Sarcoplasma of cells were fragmented. ISP II rats have had altered subepicardial portion of myocardial tissue in comparing with ISP I. Subepicardial myocytes had variable degree of kariolysis. Sarcoplasma of isolated myocytes showed separation and vacuolization of myofibrils. Lipolysis due to the adrenergic action of ISP is a potential factor in ISP myocardial necrosis development. The heart supplies a significant portion of its fatty acid substrates as free fatty acids derived by lipolysis from adipose tissue. Although lipid availability is important for the heart, excess level of fatty acids in cardiomyocytes can be deleterious. According to Mohan, lipids mobilized from the adipose depot reach their highest level in blood one hour after ISP administration and are cleared by 4 hours (15). Lipolysis as leading critical biochemical event in induced ISP myocardial necrosis does not occur until 2 hours after ISP injection, and this time frame suggests a critical role for lipolysis. Possible mechanism of difference in extent of histological changes between ISP II and ISP I subgroup is lipolysis.

CONCLUSION

Dose of ISP (250 mg/kg) induced myocardial necrosis development in rats. Subendocardial myocardium showed more vulnerability compared to subepicardial portion. Rats of ISP II had more extensive myocardial changes than these in ISP I. Obtained results pointed at possible additive influence of lipolysis on cardiotoxic effects of ISP. Administered doses of ISP enabled cTnT utilization as a marker of myocardial necrosis. This animal model of myocardial infarction is suitable for cardiotoxic and possible cardioprotective substances investigation.
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