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Cardiac troponin in ischemic cardiomyocytes: Intracellular decrease before onset of cell death

Alexander S. Streng a,1 , Leo H.J. Jacobs a,1,2 , Robert W. Schwenk b,3 , Eline P.M. Cardinaels a , Steven J.R. Meex a , Jan F.C. Glatz b , Will K.W.H. Wodzig a , Marja P. van Dieijen-Visser a,*

a Department of Clinical Chemistry, Maastricht University Medical Centre, Maastricht, The Netherlands
b Department of Molecular Genetics and Cardiovascular Research Institute Maastricht, Maastricht University, Maastricht, The Netherlands

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Aim: Cardiac troponin I (cTnI) and T (cTnT) are the most important biomarkers in the diagnosis of acute myocardial infarction (AMI). Nevertheless, they can be elevated in the absence of AMI. It is unclear if such elevations represent irreversible cardiomyocyte damage or leakage from viable cardiomyocytes. Our objective is to evaluate whether cTn is released from viable cardiomyocytes in response to ischemia and to identify differences in the release of cTn and its molecular forms.

Methods and results: HL-1 cardiomyocytes (mouse) were subjected to ischemia (modeled by anoxia with glucose deprivation). The total contents and molecular forms of cTn were determined in culture media and cell lysates. Cell viability was assessed from the release of lactate dehydrogenase (LDH). Before the release of LDH, the intracellular cTn content in ischemic cells decreased significantly compared to control (52% for cTnI; 23% for cTnT) and was not matched by a cTn increase in the medium. cTnI decreased more rapidly than cTnT, resulting in an intracellular cTnT/cTnI ratio of 25.5 after 24 h of ischemia. Western blots revealed changes in the relative amounts of fragmented cTnI and cTnT in ischemic cells.

Conclusions: HL-1 cardiomyocytes subjected to simulated ischemia released cTnI and cTnT only in combination with the release of LDH. We find no evidence of cTn release from viable cardiomyocytes, but did observe a significant decrease in cTn content, before the onset of cell death. Intracellular decrease of cTn in viable cardiomyocytes can have important consequences for the interpretation of cTn values in clinical practice.

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Introduction

The cardiac troponin (cTn) complex is a heteromeric protein complex playing an important role in the regulation of cardiac muscle contraction and consists of three different subunits: cardiac troponin I (cTnI), T (cTnT) and C (cTnC). The cTn T–I–C-complex is predominantly structurally bound to the myofibrils with only a minor fraction of cTnT and cTnI (6–8%) present free and unbound in the cytoplasm as soluble intact protein (Bleier et al., 1998; Katus et al., 1991). Due to their cardiac specificity, cTnI and cTnT are accurate and sensitive markers of cardiac injury and the most important biochemical markers used in the diagnosis of acute myocardial infarction (AMI) (Morrow et al., 2007). Nevertheless, the cardiac troponins (cTns) have been reported to be elevated in the absence of AMI, situations where irreversible cardiomyocyte damage is unlikely to play an important role (Hamm et al., 2002; Kelley et al., 2009), such as seen in subjects after strenuous exercise (Fortescue et al., 2007; Michielsen et al., 2008; Mingels et al., 2009).

It has been hypothesized that the elevated levels of cTn seen after exercise are the result of a transient increase in the cardiomyocyte membrane permeability, resulting in the release of cTn from the cytosolic cTn pool of cardiomyocytes (Neumayr et al., 2002, 2005; Remppis et al., 1995; Shave et al., 2007). In contrast, irreversible cellular damage, as observed after AMI or myocardial ischemia, will result in the release of both cytosolic and structurally bound cTn (and its complexes) from disintegrating myofibrils. AMI patients undergoing rapid reperfusion demonstrate a first cTnT peak concentration within 24 h after onset of symptoms, which has been attributed to the fast release of cytosolic cTnT (Bleier et al., 1998; Wu et al., 1998). This peak is followed by a second and persistent cTnT elevation which remains present for 7–14 days and is generally thought to represent the...
relatively slow dissociation of cTnT from the sarcomeres and release from the cells during necrosis. This biphasic release pattern has not been shown for cTnI, which release is characterized by an initial rise, directly followed by a gradual decrease to undetectable levels.

In addition to different release kinetics between cardiomyocytes with increased membrane permeability (reversible damage) and irreversible damage, there may also be a difference in released molecular forms of cTn. The cTns are thought to be targets for proteases such as caspase (Communal et al., 2002; Lancel et al., 2005) and calpain (Barta et al., 2005; Ke et al., 2008; Kositprapa et al., 2000), which are being activated and released during cell death. Therefore, cTn is expected to be released partly fragmented from irreversibly damaged cells, whereas cytosolic cTn released after reversible damage will be predominantly intact. Moreover, the complexed (myofibrill bound) forms of cTn might be more susceptible to proteolytic degradation than the free forms, as illustrated by Communal et al., who found that caspase-3 cleaves cTnT when it is in complex with cTnI and cTnC, but not as free cTnT (Communal et al., 2002).

Hessel et al. showed the release of intact cTnI from viable cardiomyocytes by stimulation of stretch-responsive integrins (Hessel et al., 2008a), a model in which irreversible damage is unlikely to occur. Conversely, irreversible cardiomyocyte damage, induced by metabolic inhibition with sodium azide, induced the simultaneous release of intact and fragmented forms of both cTnI and cTnT (Hessel et al., 2008a,b; Li et al., 2004). In a recent study, Cardinaels et al. clearly showed time-related degradation of cTnI after myocardial infarction (Cardinaels et al., 2013). Intact cTnI rapidly disappears from the serum after the ischemic event and progressively smaller cTnI fragments appear in time. Degradation and changes in the molecular forms of cTnI may have consequences for the immunoreactivity of the antibodies used in the various clinical assays. Assays based on different antibodies may therefore generate different results when measuring cTn concentrations in the same serum sample, which complicates the clinical interpretation of those measurements.

In this in vitro study we investigated the release kinetics of cTnI and its molecular fragments in cell culture using true conditions of ischemia. This was achieved by subjecting HL-1 cardiomyocytes to a complete anoxic environment with glucose deprivation. The HL-1 atrial cardiomyocyte cell line can contract in solution, maintains a differentiated adult cardiac phenotype up to at least 240 passages and can be recovered from frozen stocks (Claycomb et al., 1998). The aims of our study are, firstly, to investigate whether cTnI can be released from ischemic, but viable, cardiomyocytes and secondly, to analyze the molecular forms of the cTns released from cells subjected to ischemia.

**Methods**

**Cell culture of HL-1 atrial cardiomyocytes**

HL-1 cardiomyocytes (derived from an AT-1 mouse atrial cardiomyocyte tumor lineage) were kindly provided by Dr. W. Claycomb (Louisiana State University, New Orleans, LA, USA), and cultured as described previously (Claycomb et al., 1998; Schwenk et al., 2010). Briefly, the cells were grown in Claycomb medium (Claycomb et al., 1998) supplemented with 10% Fetal Bovine Serum (FBS), 0.1 mM norepinephrine, 2 mM L-glutamine, 100 U/mL penicillin and 100 μg/mL streptomycin at 37 °C and at an atmosphere of 5% CO2 and 95% air. For routine passaging, the cells were maintained in T-75 flasks and split twice a week after reaching confluence. For each of our experimental conditions the cells were seeded onto 6-well culture plates at a density of 100,000 cells/cm² and grown for 24 h in supplemented Claycomb medium prior to the experimental treatment. The same batch of cells was used for all time points and for both the anoxia and control experiments. The medium used for the experimental treatment of the HL-1 cells consisted of Dulbecco’s modified eagle medium (DMEM), supplemented with 2 mM L-glutamine, 100 μM non-essential amino-acids (NEAA), 0.5% albumin, 100 U/mL penicillin, 100 μg/mL streptomycin; with or without 4.5 g/L glucose.

Ischemia was modeled by anoxia and glucose deprivation. At the start of our experiment (t = 0), Claycomb medium was replaced with supplemented DMEM without glucose and the cells were immediately placed in an anoxic environment (MACS VA500 microaerophilic work-station, Don Whitley Scientific, Shipley, UK). The atmosphere in the chamber consisted of 5% CO2, 5% H2 and residual N2. Anoxia treatment is performed for 1, 3, 5, 7, 9, 12 and 24 h.

As a control, non-treated HL-1 cells grown for 0, 1, 3, 5, 7, 9, 12 and 24 h in supplemented DMEM with glucose were grown alongside the ischemic cells. Fig. 1 depicts the workflow used in the cell culture experiments from routine cell passing to the storage of individual medium and lysate samples.

**Sample collection and cTn measurements**

After each treatment period, the culture medium was collected and centrifuged for 5 min at 500 RCF and the supernatant was stored at −80 °C. Immediately after medium removal the cells were washed twice with cold (4 °C) phosphate buffered saline (PBS) and 600 μL cold (4 °C) RIPA lysis buffer, containing 150 mM NaCl, 50 mM Tris–HCl, 1% (v/v) Igepal CA-630, 0.5% (w/v) sodium deoxycholate, 0.1% (w/v) SDS, and 4% (v/v) complete protease inhibitor cocktail and 5% (v/v) PhosSTOP phosphatase inhibitor cocktail was added to each well. The cells were lysed for 2 h with mild agitation at 4 °C. Resulting cell lysates were stored at −80 °C until analysis.

Troponin concentrations were measured in DMEM and RIPA buffer aliquots. cTn concentrations were measured using the 4th generation cTn immunoassay on the Elecsys 2010 instrument (Roche Diagnostics, Mannheim, Germany), with a limit of detection (LOD) of ~0.01 μg/L, a 10% CV of 0.03 μg/L and a linear measuring range of 0.010–25.00 μg/L, according to the product insert. cTn concentrations were measured using the AxSYM® Troponin-I ADV assay (Abbott Diagnostics, Wiesbaden, Germany), with an LOD of 0.02 μg/L, a 10% CV of 0.16 μg/L and a linear measuring range of 0.02–22.78 μg/L.

**Cell death**

The release of lactate dehydrogenase (LDH) into the culture medium was used to quantify cell death. The cytosolic protein LDH is released from cells exhibiting a loss of membrane integrity as seen during primary and secondary necrosis and its release is commonly used to quantify cell death (Hessel et al., 2008b; Seymour et al., 2003). LDH activity was determined with the LD-P lactate dehydrogenase assay on the Synchon LX20 pro clinical system (Beckman Coulter Inc., Palo Alto, CA, USA), with a linear measuring range of 20–2500 IU/L and a 5.3% CV of 22.5 IU/L. The release of LDH is expressed as the LDH activity in the culture medium as a percentage of the total LDH in the well:

\[
\text{LDH}_{\text{medium}}/\left(\text{LDH}_{\text{medium}} + \text{LDH}_{\text{lysates}}\right) \times 100\%.
\]

In addition to the LDH release, the amount of viable cells was also estimated by measuring the total protein content in the collected cell lysates with the BCA (bicinchoninic acid) Protein Assay Reagent (Thermo Fisher Scientific, Inc., MA, USA), according to the manufacturer’s instructions.

**Validation of protein measurements**

The clinical assays used to measure protein concentrations are optimized for the use in human serum. In order to exclude any matrix-effects from influencing our measurements, we validated their use in DMEM and RIPA buffer. Fig. 2 shows LDH (a), cTnI (b) and cTnT (c) concentration curves of untreated cell lysates serially diluted in either
DMEM or RIPA buffer. From the figure follows that the LDH, cTnI and cTnT assays used in this study give linear results over their entire measuring ranges in our used buffers and that the DMEM and RIPA curves are superimposed, indicating that measurements in both buffers are comparable to one another. Samples with concentrations outside of this range were diluted 10 times in the appropriate buffer before measurement. The Roche cTnT immunoassay shows a slight increase in measured cTnT concentration in the medium as compared to the RIPA buffer, resulting in a measurement that is on average 14% higher in the medium (Fig. 2c).

Detection of cTnI and cTnT degradation products

Before protein separation, the cell lysate samples were diluted to a cTnT concentration of 1 μg/L or to a cTnI concentration of 7.5 μg/L (unless stated otherwise). The cell culture lysates were each mixed with 4× XT Sample Buffer (Bio-Rad, 161-0791) and 2× Reducing Agent (Bio-Rad, 161-0792) and heated for 5 min at 95 °C. Subsequently, 25 μL of the samples, containing equal amounts of cTnT and cTnI, was separated on a 12% Criterion XT SDS-PAGE gel (Bio-Rad, 345-0118). Proteins were then transferred to a nitrocellulose membrane (Bio-Rad, 162-0115, 0.45 μm). Further treatment of the blots was done with the SNAP i.d. protein detection system (Millipore) according to the manufacturer’s instructions. Briefly, blocking was performed for 10 s with 0.5% (w/v) Blocking buffer (Bio-Rad, 170-6404). The incubation with the primary antibody (specified below) was performed for 10 min. After washing (washed 4× with PBS with 0.1% (v/v) Tween-20 (PBST)), the blot was incubated with secondary antibody (specified below) and washed 4× with PBST. The following primary and secondary antibodies were used:

- All primary antibodies were mouse monoclonal antibodies (mAbs) ordered from HyTest LTD, Turku, Finland. For cTnT the primary anti-cTnT solution was a mixture of the following mAbs: 9G6 (epitope: 1–60), 1c11 (epitope: 171–190), 7F4 (epitope: 67–86), and 7A9 (epitope: 171–190) at concentrations of 2 μg/mL. For cTnI, the primary anti-cTnI was a mixture of 3 anti-cTnI mAbs: 19C7 (epitope: 41–49), MF4 (epitope: 190–196) and 84 (epitope: 117–126) at concentrations of 4 μg/mL. The secondary antibody solution consisted of 0.4 μg/mL goat anti-mouse IgG conjugated with peroxidase (Dako, P0447) in PBST. Membranes were incubated for 5 min in Super Signal West Femto Substrate (Thermo Scientific, 34096) and exposure was detected using the ChemiDoc XRS scanner (Bio-Rad).

Data analysis and statistics

Experiments were performed in triplicate and data are presented as mean ± S.E.M. Comparisons are made with Student’s paired t-test with p < 0.05 considered statistically significant. Protein release (for cTnT, cTnI and LDH) is defined as the percentage of total protein that is present in the medium: Release = [Pm] / [Pm + Pl] * 100%; Pl = protein amount in the medium and PIM is the protein amount in the cell lysate. The recovery of each of the proteins is defined as the amount of protein that can be recovered at each time point, relative to the amount in the control: (Recovery = [Pm] / [Pm + Pl]experimental ) / ([Pm] / [Pm]control) * 100%. The relative amount of intact and fragmented cTnI and cTnT was determined by densitometry, using the Quantity One software (Bio-Rad, Version 4.6.5.).

Results

Protein release after ischemic damage

Cell viability was assessed by measuring the release of LDH into the medium. In response to ischemia, LDH is gradually being released into the culture medium starting between 3 and 5 h of treatment (Fig. 3a and b). During this period, intracellular cTnI and cTnT contents decreased by 52 ± 7% and 23 ± 7%, respectively, compared to control (Fig. 3c). This intracellular decrease was not matched by an increase in the medium (Fig. 3d). We were only able to detect cTnI in the medium after the release of LDH. This means that there is a considerable loss in the total amount of cTnI, illustrated by a recovery decrease during the first 3 h of treatment (Fig. 3e). The slight increase in protein recoveries after 24 h is most likely caused by a decrease of the net cellular protein content in the control cells (Fig. 3a and c) due to substrate depletion of the culture medium. Throughout the treatment, all of the LDH could be recovered (average recovery: 107 ± 5%). As an alternative measure of cell viability, the total protein content of the cell lysates was assessed and showed an excellent correlation with the LDH content (Spearman’s rho = 0.843, p < 0.001; results not shown).

Fig. 1. Flow diagram depicting the workflow of the cell culture experiments. Cells were continuously routinely passaged in supplemented Claycomb medium. The experiment started with the seeding of 100,000 cells/cm² to multiple 6-well plates. The workflow was followed as indicated until medium and lysate samples were stored at −80 °C. These samples were further processed as described in the method section.
Western blots of the cell lysates were performed to investigate whether a change in the molecular forms of cTn could be found. A mixture of different mAbs was used to create optimal coverage of both proteins. The results show two separate bands for cTnI: intact cTnI migrating to 29 kDa and a cTnI fragment migrating to 25 kDa (Fig. 5a). About 91% of the total signal on this blot originates from the 29 kDa band and 9% from the 25 kDa band. After 5 h of anoxia the relative intensities of the bands began to change with increasing lengths of treatment. After 12 h, the relative signal of the 25 kDa band had increased to 34% and the 29 kDa band decreased to 66%. After 24 h, cTnI content has dropped to a level where it was no longer possible for them to be visualized.

For cTnT, a similar effect was seen, although less pronounced. In the control samples and during the first hours of ischemia, 98–99% of the total signal originated from the 37 kDa (intact cTnT) band (Fig. 5b). Upon progressive lengths of ischemia, the relative amount of the 37 kDa band decreased and a cTnT fragment band at 27 kDa started appearing. After 24 h the intensity of the fragment band increased to 7.4% and the intact cTnT band decreased accordingly.

Western blots from the culture media were also performed, however, no fragmentation was observed and only a very faint, single, intact, cTnI and cTnT band was visible throughout the experiment (results not shown).

Discussion

In this study, we explored the release of cTnT and cTnI in an in vitro model in which HL-1 cardiomyocytes were subjected to ischemia and glucose deprivation. cTnI and cTnT were being released from the cardiomyocytes starting between 3 and 5 h after the initiation of treatment. This release was associated with a concomitant release of LDH, implying that the release of troponin only occurred after the onset of cell-death. No cTn release was observed during the first 3 h. Interestingly, we did observe a significant decrease of the cTnI and cTnT contents (52% and 23%, respectively) during this period which was not matched by an increase in the medium and was more pronounced for cTnI than for cTnT.

The release characteristics of cTnI and cTnT from the cells in response to ischemia seemed comparable to those seen after AMI. Following an AMI, the cTnI and cTnT concentrations begin to rise 3–6 h after the onset of AMI and peak after 12–24 h (Collinson, 1998; Januzzi, 2011). Similarly, the HL-1 cells began releasing cTnI and cTnT between 3 and 5 h of ischemia. Within our model, we found a more pronounced increase in cTnI medium levels in the early hours after the onset of ischemia than for cTnT (Fig. 3d and f). cTnI was released alongside LDH, while cTnT release appears to occur 1 or 2 h later. A possible explanation for this could lie in the fact that cTnI is located outermost on the thin filament (Toyo-Oka and Ross, 1981), which may facilitate a more readily access to proteases such as caspase (Communal et al., 2002) and calpain (Barta et al., 2005; Ke et al., 2008; Kositprapa et al., 2000), whereas cTnT is structurally bound to tropomyosin with high affinity (Feng et al., 2009; Perry, 1998).

In the case of LDH, every decrease in the lysates resulted in an increase in the medium. In the end, an average of 107 ± 5% of total LDH was recovered. However, before the cTns were released into the medium, a significant decrease in their cellular contents was detected. This decrease was not matched by an increase in the medium and is not observed in control cells. We have not been able to explain this decrease, but an analytical effect is unlikely. In this report, the presented data was not corrected for matrix effects of the assays. Even though a slight measurement difference between culture medium and lysis buffer was shown for the Roche cTnT assay (Fig. 2c), correcting for this deviation would reduce the recovery of cTnT even further (61% recovery instead of 67% after 24 h of treatment, results not shown). Several explanations can be offered for this seemingly selective decrease in troponin content. The
most prominent one is the complete degradation of these proteins by the proteasome in a process called autophagy. Autophagy pathways are highly upregulated in cardiomyocytes upon ischemia as part of a cardioprotective mechanism (Sciarretta et al., 2013). This mechanism reduces necrosis and apoptosis and could therefore explain why our cells are able to survive for a seemingly long period without glucose and oxygen. The intracellular degradation of troponin is further shown by western blots of cTnI and cTnT which showed an increase in degradation products in response to the ischemia treatment. In addition, the decreased measured concentration might also be the result of other modifications, such as phosphorylation (Layland et al., 2005; Streng et al., 2013) and ubiquitination (Kedar et al., 2004; Witt et al., 2005), which might have an influence on antibody affinity. Irrespective of the exact underlying mechanism for the intracellular decrease, the decreased recoveries suggest that the released cTns represent only a relatively small part of the total amount of cTn that was originally present in the cardiomyocytes. This finding is a
A study by Hessle et al. (2008b) found recoveries of 34% for cTnI and 39% for cTnT in response to 24 h of metabolic inhibition using sodium azide. More importantly, this phenomenon might also occur in vivo, where the reduced recoveries may be even more pronounced. Early reports have shown that at 72 h after AMI only 4–5% of cTnT is recovered (in comparison to cytoplasmic enzymes like LDH) (Kragten et al., 1996).

In this study, we used an immortalized atrial cardiomyocyte cell line. By using these cells, we were able to perform a multitude of experiments to validate and fine-tune our model. Our results have been very reproducible and treatment and control conditions could be carefully controlled. However, we recognize that primary cardiomyocytes would have resembled the in vivo situation more closely and that our cells are atrial instead of ventricular. On the other hand, HL-1 cardiomyocytes are well defined and extensively characterized. They have been used in numerous studies since their creation and have been shown to resemble primary cardiomyocytes very closely (White et al., 2004). Despite these limitations, the implications of molecular changes on the immunoreactivity of cTn in the clinical immunoassays should not be underestimated.

Results from our study suggest that in vivo, only part of the released cTn is measured in the serum. Future experiments need to be performed to unravel the precise mechanisms behind this “disappearance” of cTn from viable cardiomyocytes.

Conflict of interest statement

The authors declare that they have no conflict of interest.

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The authors are grateful to Kevin Gerritsen for valuable experimental contributions and to Dr. Douwe de Boer for insightful discussions.

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