Non-Ischemic Remote Cardioprotective Phenomena

Song Yang1 and Xiaoping Ren2,3*

1Hand and Microsurgical Center, Department of Orthopedics, The Second Affiliated Hospital of Harbin Medical University, China
2Clinical Translational Medicine Center, Harbin Medical University, Harbin, PR China
3Department of Pharmacology and Cell Biophysics, University of Cincinnati College of Medicine, Cincinnati, OH 45267-0575, USA.

Abstract

Ischemic preconditioning (IPC) is a powerful cardioprotective phenomenon that occurs in several species. IPC means that transient ischemia makes the cardioprotection in the subsequent period of ischemic thus it limits the scope of myocardial infarction. The reports of recent years have obtained the conclusion that this cardioprotection can be also induced by the non-ischemic preconditioning such as metabolic stimulation of myocardial cells and distention of the left ventricle. From the physiological point of view, the protective effect of non-ischemic preconditioning is associated with the higher proportion of myocardial cell metabolism. Remote ischemic preconditioning (RIC) is a therapeutic measure for cardioprotection against the deleterious effect of acute ischemia-reperfusion injury (IRI). And, it’s beneficial effects are also seen in other organs (lung, liver, kidney, brain) and tissues (skeletal muscle). Although the mechanism of RIC has not yet been completely determined, there are several probable hypotheses supporting the cardioprotection induced by RIC, which include the neuronal pathway, humoral pathway, and systemic response. Recently, some researchers have provided experimental evidence for the cardioprotection under non-ischemic trauma at a site from the heart produced by a transverse abdominal incision. This study demonstrated that protection could be elicited by non-ischemic stimulus which they term “remote preconditioning of trauma (RPCT)”. Because of remote preconditioning of trauma, which is produced by an abdominal incision only through the skin, so it means that it’s non-ischemic. Although remote ischemic stimulus has been shown to elicit cardioprotection against IRI, there are not many reports about the remote non-ischemic stimulus and its mechanism. In this article, we provide a review of cardioprotection, the potential mechanism of RPCT as well as its clinical prospects.

Keywords: Cardioprotection; Remote preconditioning of trauma; Animal model; Potential underlying mechanism; Clinical application

Introduction

With the first described by Murry et al. [1] in 1986, Ischemic preconditioning (IPC) has been known for past two decades. IPC is a powerful cardioprotective phenomenon that has been demonstrated that occurs in several species [2,3] and there is strong evidence that it also occurs in humans [4]. However, IPC demands an absolutely intervention to be applied to the heart directly which may not be necessary in all clinical events. So in 1993, Przeklenk et al. [5] showed the experimental evidence of remote preconditioning by demonstrating the protection in the left anterior coronary territory following a preconditioning stimulus in the circumflex coronary artery. Subsequently, some researchers termed this cardioprotective phenomenon “remote ischemic preconditioning (RIC)”. Kharbanda et al. [6] introduced that transient limb which was induced by short periods of ischemia by tourniquet or blood pressure cuff had proven to be an uncomplicated method of inducing RIC. Meanwhile, the potential cardioprotection has been demonstrated in animal models as well as patients in Botker's report [7]. The protective phenomenon has been translated to positive clinical trials [8]. RIC is a therapeutic measure for cardioprotection against the deleterious effect of acute ischemia-reperfusion injury (IRI). Moreover, its beneficial effects are also seen in other organs (kidney, brain) and tissues (skeletal muscle) [9-11]. As for the mechanism of the RIC, some researchers have shown that remote preconditioning by transient limb ischemia in rabbits and humans is associated with release of a blood-borne, hydrophobic and small circulating factor, the effect of which can be blocked by the opiate receptor blocker, naloxone [12]. Subsequently, the same experimental group established the signaling of PI3K/Akt/GSK3β in the cardioprotection by the same stimulus of RIC [13]. In addition, it has been demonstrated that bradykinin (BK), adenosine, opioids, and norepinephrine (NE) all play an important role in RIC [14-16]. Some studies show that both IPC and RIC are associated with the activation of BK2R [14,17]. In brief, a variety of intracellular signaling mediators have been implicated in the protective effect of RIC such as G-protein cell surface coupled receptors, PKC, reactive oxygen release of circulating humoral factor(s), and activation of a systemic protective effect (such as an anti-apoptotic or anti-inflammatory response).

However, does remote non-ischemic preconditioning occur in animal or humans? This is an essential question from the clinical viewpoint meanwhile it could have the therapeutic implications [18,19]. Cardiac ischemia/reperfusion injury (IRI) contributes significantly to morbidity and mortality all over the world. Various researches for protecting the heart against myocardial infarction (MI) and IRI have been developed. In 2004, Ren et al. observed that infarct size after in vivo ischemia/reperfusion was altered by nonischemic surgical trauma. The transverse abdominal incision may result in the decreased infarct size in a TNFα-independent manner [20]. This study suggested that the cardioprotection could be induced by non-ischemic stimulus at a distance from the heart. To describe the phenomenon of this cardioprotection, they term it “remote preconditioning of trauma” (RPCT). In 2009, subsequently, the same group showed that the cardioprotective effect of RPCT is initiated by skin nociception, and requires neurogenic signaling involving spinal nerves and activation of cardiac sensory and

*Corresponding author: Xiaoping Ren, Department of Orthopedics, The Second Affiliated Hospital, Harbin Medical University, 194 Xuefu Rd, Nangiang District, Harbin, China, Tel: 451-136-446-065-83; E-mail: renxpj@uc.edu

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sympathetic nerves [21]. These results demonstrated that bradykinin-independent activation and repression, respectively, of PKCe and PKCd in myocardium after RPCT, and their conclusion shows that the cardioprotection is associated with the K⁺ leak. The experiment suggests that nontraumatic nociceptive preconditioning represents a novel therapeutic strategy for cardioprotection with great potential clinical utility. Furthermore, in 2011, Gross et al. has showed that this protective phenomenon of heart in canine hearts [22]. Although it is found in canine hearts, it is significant for clinical setting and has implicated the cytochrome P450 epoxidegenase pathway as a mediatory factor. This result suggests that bradykinin (BK) and the epoxyeicosatrienoic acids (EETs) share cardioprotective properties in a large animal model of RPCT. Recently, as a completely new concept, because RPCT could be mimicked non-invasively by topical application of capsaicin to induce the cardioprotection suggesting an innovative application without ischemia or trauma [23], since RPCT has been accepted by the researchers. At present, Ren's conference abstract expands the concept of RPCT again. Their experimental group tested the hypothesis that remote electrical stimulation (ES) of skin nociceptors could produce the effect similar to that of remote preconditioning of trauma. Mice I/R model were established in vivo [24]. ES was used 15 minutes prior to I/R or at the beginning of reperfusion. The results of research show that the skin nociceptor stimulation at specific points exactly reduce infarct size (85% reduction), and they also find that cardioprotective effect of ES was abolished by both β-AR antagonism and in BK2R mice. Western blot results show PKC isoform translocation is altered after ES and pharmacologic blockade of PKC prevented cardioprotection due to ES. Therefore, they draw the conclusion 1) ES elicited a powerful cardioprotection via a neural mechanism. 2) The protective effect of ES against MI involves β-AR, BK2R signaling and PKC modulation. In other experiment, the evidence was found that transcutaneous electrical nerve stimulation had been used widely clinically for nociceptive suppression by means of repeated C-fiber stimulation [25]. From remote cardioprotection described to the establishment of RPCT, there have been gone for past two decades. A variety of experimental models were set out to support the experiments for the powerful evidence. In the original study by Kharbanda and colleagues, transient ischemia of hind limb using a tourniquet was shown to reduce subsequent myocardial infarction (MI) size in pigs [6]. Subsequently, in experiment of introducing the RPCT and demonstrating the mechanism of RPCT by Ren and Jones group, a minimally traumatic mouse model was used to ascertain the effect of remote nonischemic surgical trauma upon I/R injury [20]. The abdominal incision used as the stimulus for RPCT was a 2-cm transverse incision located on the abdominal midline. The incision extended through the skin and muscle and into the peritoneum and was immediately closed after 3-0 polypropylene sutures. Katherine's report showed that the rabbits were allocated several groups. Remote ischemic preconditioning was induced through four cycles of 5 minutes of hind limb ischemia (via a tourniquet). Some groups applied the 5 ml of 0.1% capsaicin topical analgesic cream and pharmacologic blockade of PKC prevented cardioprotection due to ES. The delay of lethal cell injury in ischemic myocardium. Circulation 74: 1124-1136.

Conclusion

The discovery of remote cardioprotection has offered a significant therapeutic strategy for protecting the heart against the myocardial ischemia-reperfusion injury. The experimental research of past decade (especially RPCT) have encouraged scientists to do further exploration about the complex mechanism underlying the protective effect. Considering that the remote cardioprotection has been induced in many species of animals, the researchers should think about that how to combine the remote cardioprotection with the clinical work. This is the most important question from a clinical viewpoint, and could have therapeutic implications. In summary, remote cardioprotection especially remote non-ischemic preconditioning including RTCP is a pretty new field and could have clinical implications.

References

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