Remote ischaemic preconditioning: the current best hope for improved myocardial protection in cardiac surgery?

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Ischaemic preconditioning has been recognised as a major cardioprotective phenomenon for many years.1 2 Cycles of non-lethal ischaemia and reperfusion applied to the heart before a potentially lethal ischaemic insult have the ability to reduce infarct size by >50%. More recently, it became apparent that the protection generated by this classical form of direct ischaemic preconditioning could be replicated when the non-lethal ischaemia was applied to one segment of the heart and the lethal ischaemia applied to a separate segment.3 Thereafter, it became established that the same protection could also occur even if the preconditioning ischaemic stimulus was applied completely distant from the target organ requiring protection—that is, transient ischaemia of a remote organ or limb could still generate protection for the organ being subsequently challenged by lethal ischaemia.4 There is now clinical evidence, suggesting that this remarkable remote ischaemic preconditioning (RIPC) phenomenon may represent a simple, inexpensive, easily applied method of increasing cardioprotection during an array of interventional procedures that require a period of cardiac ischaemia to allow repair or intervention. Moreover, as it is now recognised that such protection may be achieved by starting the cyclical remote ischaemia and reperfusion after the period of injurious cardiac ischaemia has started—so-called remote post- or peri-conditioning—the possibility arises of enhancing protection in other situations, including transplantation.5

Several clinical reports of RIPC in cardiovascular surgery have now been published. In children undergoing congenital heart defect repairs using cardiopulmonary bypass, lower limb RIPC has been shown to reduce troponin release and inotrope requirements.6 In adults undergoing coronary artery bypass (CABG) surgery, intermittent upper limb ischaemia has been followed by reductions in postoperative release of lactate dehydrogenase7 and troponin T.8 In abdominal aortic aneurysm surgery, RIPC, induced by unilateral iliac artery clamping has reduced troponin release and renal injury.9

In most of these studies the release of troponin is used as a marker of the quantum of injury suffered by the myocardium. Post-cardiac surgery troponin levels have been used to compare different myocardial protection strategies and provide an indicator of long-term outcome.10-12 Which troponin metric—isolated values at specific time points or area under the curve (AUC) release—provides the most prognostically important information is not yet known. Additionally, whether troponin release in the first few hours after surgery reflects true infarction or a change in sarcosomal integrity or permeability has been questioned.13

In this issue of Heart Venugopal et al.14 provide further evidence that RIPC may improve myocardial protection in humans (see page 1567). This single-centre randomised trial studied RIPC or a placebo intervention in 45 patients undergoing CABG with or without concomitant aortic valve replacement (AVR) as an adjunct to antegrade ± retrograde blood cardioplegia myocardial protection. Patients with diabetes, renal, hepatic or pulmonary dysfunction were excluded as were those with unstable angina or myocardial infarction within 4 weeks of surgery.

The remote preconditioning stimulus comprised three 5 min cycles of forearm ischaemia, induced by inflating a blood pressure cuff on the upper arm to 200 mm Hg, with an intervening 5 min reperfusion. The control group had a deflated cuff placed on the upper arm for 30 min. On parametric analysis, RIPC was found to reduce the area under the curve (AUC) serum cardiac troponin T (cTnT) release by >40%. The magnitude of the effect was similar to that seen in a cohort of patients undergoing intermittent ischaemic arrest as a mode of myocardial protection, reported by the same group previously.8 Unfortunately, clinical outcomes are not reported.

On the basis of this and other work, there is now a need (a) to determine the efficacy of RIPC in promoting protection in other forms of cardiac surgery; (b) to ascertain whether the changes in troponin release are reproducible in other studies; (c) to establish if these changes are reflected in improved clinical outcomes and that RIPC independently reduces risk.15 However, before these studies are designed and started the design and analysis of this study require some further comment.

First, blinding of treatment allocation was applied to patients and surgeons only; anaesthetists (who administer agents capable of preconditioning or affecting myocardial protection) and investigators were not blinded. Similar proportions of patients received isoflurane or sevoflurane for anaesthetic maintenance but dosages were not reported. As such volatile anaesthetic agents may induce a dose-dependent conditioning effect,16 a potential for inadvertent bias arises. Second, the study was small and contained only half of the estimated number of patients to detect the initially expected difference in AUC cTnT of 15 μg/L ± 1.72 h (standard deviation 25 μg/1.72 h) quoted in the statistical methodology. Statistical significance was actually attained with a mean difference and sample size and this is attributable to the lower than expected variance seen in the RIPC group. Third, the study also included patients requiring AVR; whether RIPC was effective in the patients undergoing CABG alone is not reported. The larger number of combined AVR/CABG cases contributed to a longer mean bypass time in the control group and bypass time was an independent predictor of greater troponin release. Despite this potentially confounding effect, an intergroup statistically significant difference was maintained after correction for bypass time using a generalised linear model. Fourth, many of the important variables in the study—for example, bypass time, cross-clamp time and AUC cTnT had unequal variances yet were analysed parametrically unlike the authors’ previous report. Lastly, although the drug history is reported, whether potentially relevant
drugs—for example, atorvastatin, potassium channel blockers,17 18 were administered in the 24 h preoperatively is not clear.

Nevertheless, the effect of RIPC on troponin release was large and the data are very encouraging. In particular, the troponin effect was observed despite the use of halogenated anaesthetic gases in the majority of patients. Several studies have demonstrated that such volatile anaesthetic agents may reduce evidence of myocardial injury during CABG through what is thought to be a preconditioning mechanism.19–21 Thus, this study is important as it suggests that the effect of RIPC is, at least additive to any potential of RIPC could be realised. If the findings of this study are corroborated by larger trials, the great potential of RIPC could be realised.

Theoretically, RIPC could be used in the brainstem dead donor to generate cardioprotection. However, although the preconditioning site would remain innervated, central neural connection is lost. Moreover, in this circumstance, reperfusion does not occur in the preconditioned environment unless the stimulus is repeated in the recipient. This is indeed the case, a donor heart, denervated at transplantation may still be protected from a postimplantation ischaemic insult by an RIPC stimulus in the recipient.22 However, whether such conditioning could protect during the retrieval, transport and implant period of transplantation is not clear.

The data accruing thus far for RIPC in both the medical and surgical cardiological arenas are promising and at present it appears to be one of the most important potential myocardial protective adjuncts so far identified.23–26 Let us be sure to investigate its role, comprehensively, throughout cardiac surgery, in large studies with clinical end points.

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REFERENCES


Education in Heart: 10th anniversary

Education in Heart (EIH) first appeared in this journal 10 years ago, since when it has been one of our flagship sections, comprising articles that have regularly been among the most widely accessed of Heart’s published output. Throughout this period it has been edited semi-autonomously—and very successfully—by Peter Mills, who is now standing down, the baton passing to Jeroen Bax who takes over this month. EIH’s 10th anniversary is a suitable time to look back at its achievements and those of its departing editor whose key contribution has been to identify an international cadre of opinion leaders able to produce material that goes beyond the traditional review process into areas specifically designed to educate. This has required not only subspecialty expertise on the part of the authors but also considerable editorial skill, ensuring that the contemporary content of EIH is presented in clear, unambiguous prose, with careful structuring, relevant illustration and annotated references to direct the reader to key source material. To underpin EIH’s educational agenda Peter Mills has commissioned all articles against national and European curricula to ensure the requirements of our readership are fully met, with proper regard to both mainstream and Cinderella subject matter and to the need for updates, particularly in areas where new discoveries are proceeding most rapidly. The formal stamp of educational approval has been delivered through submission of all articles to the European Board for Accreditation in Cardiology (EBAC) in order to obtain European continuing medical education (CME) accreditation, with multiple-choice questions available at a dedicated web address. EIH is also scheduled to play a central role in the European revalidation initiative currently under development. We now welcome Jeroen Bax as the new Editor of EIH and acknowledge the work of Peter Mills in its evolution these last 10 years.

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