Comment

The broken promise of remote ischaemic conditioning

Intermittent ischaemia and reperfusion cycles, or ischaemic conditioning, has been viewed as a promising potential cardioprotective technique since its first description by Murry and colleagues in 1986.1 The most encouraging iteration of this technique to emerge over the past three decades has been the application of a modified blood pressure cuff placed on the upper arm to induce cycles of remote, transient ischaemia that relay protection to the heart via predominantly neurohormonal mechanisms.² Remote ischaemic conditioning showed early potential in reducing infarct size after myocardial infarction in animal models.^{3,4} The initial CONDI-1 trial, although small (n=333), suggested that remote ischaemic conditions used as an adjunct to the gold standard therapy of primary percutaneous coronary intervention (PPCI) for ST-elevation myocardial infarction (STEMI) improved the myocardial salvage index⁵ as well as left ventricular systolic function at 30 days in patients at risk of large infarcts.⁶ The results of the RIC-STEMI trial (n=258) echoed those of CONDI-1, with remote ischaemic conditioning showing additional benefits in reducing cardiac deaths and hospitalisations and increasing the overall mean improvement of ejection fraction after 1 year.⁷ Medium-term follow-up data (at a median 3.6 years) from the LIPSIA CONDITIONING study (n=696) showed that remote ischaemic conditioning combined with post-conditioning reduced new heart failure diagnoses compared with remote ischaemic conditioning alone.⁸ None of these trials showed a reduction in infarct size when remote ischaemic conditioning was applied; however, cardiac MRI data suggest the ability of remote ischaemic conditioning to induce positive ventricular remodelling in the absence of absolute infarct size reduction.⁹

In *The Lancet*, Derek Hausenloy and colleagues¹⁰ report the findings of the combined CONDI-2/ERIC-PPCI trial, a large and appropriately powered randomised controlled trial to address whether remote ischaemic conditioning as an adjunct to PPCI can improve the clinical outcomes of patients with STEMI when compared with PPCI alone. The trial was a pragmatic combination of two very similar protocol designs. Between 2013 and 2018, 5401 patients undergoing PPCI for STEMI across four European countries were randomly allocated (1:1) to PPCI alone (mean age 63.1 years [SD 12.2], 576 [22.4%] women) or PPCI plus remote ischaemic conditioning (63.9 years [12.1], 611 [24.0%] women). Remote ischaemic conditioning was administered either in the ambulance en route to the hospital or in the hospital, before or during the PPCI procedure. The combined primary endpoint, cardiac death or hospitalisation for heart failure within 1 year, did not differ between groups, occurring in 220 (8.6%) patients in the control group and 239 (9.4%) in the remote ischaemic conditioning group (hazard ratio [HR] 1.10 [95% Cl 0.91-1.32], p=0.32). This finding was consistent in prespecified subgroup analyses by age, presence of diabetes, pre-PPCI TIMI flow, ischaemia time, or infarct location. Furthermore, the 12-month incidence of major adverse cardiac and cerebrovascular events (a composite of all-cause death, reinfarction, unplanned revascularisation, and stroke) did not differ significantly between the groups (HR 1.09 [95% CI 0.90-1.32], p=0.38).

Hausenloy and colleagues should be praised for their robust and clinically relevant trial to assess the role of remote ischaemic conditioning in the current landscape of STEMI management. The findings show clearly that the use of remote ischaemic conditioning around the time of PPCI for STEMI adds no clinical benefit for outcomes assessed within 1 year of the procedure.¹⁰ A cautionary note pertains to the follow-up time of only 1 year: our current understanding of cardiac remodelling post-STEMI is that the clinical benefits derived from cardioprotective interventions might not manifest until after 2 years, or perhaps longer,¹¹ and as such the trial could have benefited from a longer follow-up time. The authors also acknowledge that the remote ischaemic conditioning protocol used (four 5-min cycles of upper arm cuff inflations and deflations applied before PPCI) might have been insufficient to induce an adequate neurohormonal response in some patients, although this protocol has been shown to be the most efficacious one in animal and subsequent clinical studies.12

The role of remote ischaemic conditioning in improving the lives of patients with STEMI has been thrown sharply into question. Despite the early promise of a number of preclinical and smaller proof-of-principle





Published Online September 6, 2019 http://dx.doi.org/10.1016/ S0140-6736(19)32047-1 See **Articles** page 1415 clinical trials, this large and adequately powered trial points to a lack of clinical efficacy for remote ischaemic conditioning in this setting. As such, it might be time to abandon this form of cardioprotection in favour of more effective therapies to extend and improve the lives of these patients.

Andrew Peter Vanezis

CK Hui Heart Centre, Royal Alexandra Hospital, Edmonton, AB T5H 3V9, Canada andrew.vanezis@nhs.net

I declare no competing interests.

Copyright @ 2019 The Author(s). Published by Elsevier Ltd. This is an Open Access article under the CC BY-NC-ND 4.0 license.

- 1 Murry CE, Jennings RB, Reimer KA. Preconditioning with ischemia: a delay of lethal cell injury in ischemic myocardium. *Circulation* 1986; **74**: 1124–36.
- 2 Davidson SM, Arjun S, Basalay MV, et al. The 10th Biennial Hatter Cardiovascular Institute workshop: cellular protection-evaluating new directions in the setting of myocardial infarction, ischaemic stroke, and cardio-oncology. *Basic Res Cardiol* 2018; **113:** 43.
- 3 Kerendi F, Kin H, Halkos ME, et al. Remote postconditioning. Brief renal ischemia and reperfusion applied before coronary artery reperfusion reduces myocardial infarct size via endogenous activation of adenosine receptors. Basic Res Cardiol 2005; 100: 404–12.
- 4 Andreka G, Vertesaljai M, Szantho G, et al. Remote ischaemic postconditioning protects the heart during acute myocardial infarction in pigs. *Heart* 2007; **93**: 749–52.

- Bøtker HE, Kharbanda R, Schmidt MR, et al. Remote ischaemic conditioning before hospital admission, as a complement to angioplasty, and effect on myocardial salvage in patients with acute myocardial infarction: a randomised trial. *Lancet* 2010; **375**: 727–34.
- 6 Munk K, Andersen NH, Schmidt MR, et al. Remote ischemic conditioning in patients with myocardial infarction treated with primary angioplasty: impact on left ventricular function assessed by comprehensive echocardiography and gated single-photon emission CT. *Circ Cardiovasc Imaging* 2010; **3:** 656–62.
- 7 Gaspar A, Lourenço AP, Pereira MÁ, et al. Randomized controlled trial of remote ischaemic conditioning in ST-elevation myocardial infarction as adjuvant to primary angioplasty (RIC-STEMI). Basic Res Cardiol 2018; 113: 14.
- 8 Eitel I, Stiermaier T, Rommel KP, et al. Cardioprotection by combined intrahospital remote ischaemic perconditioning and postconditioning in ST-elevation myocardial infarction: the randomized LIPSIA CONDITIONING trial. Eur Heart J 2015; 36: 3049–57.
- 9 Traverse JH, Swingen CM, Henry TD, et al. NHLBI-sponsored randomized trial of postconditioning during primary percutaneous coronary intervention for ST-elevation myocardial infarction. *Circ Res* 2019; 124: 769–78.
- 10 Hausenloy DJ, Kharbanda RK, Møller UK, et al. Effect of remote ischaemic conditioning on clinical outcomes in patients with acute myocardial infarction (CONDI-2/ERIC-PPCI): a single-blind randomised controlled trial. *Lancet* 2019; published online Sept 6. http://dx.doi.org/10.1016/ S0140-6736(19)32039-2.
- 11 Hassell ME, Vlastra W, Robbers L, et al. Long-term left ventricular remodelling after revascularisation for ST-segment elevation myocardial infarction as assessed by cardiac magnetic resonance imaging. *Open Heart* 2017; 4: e000569.
- 12 Johnsen J, Pryds K, Salman R, Løfgren B, Kristiansen SB, Bøtker HE. The remote ischemic preconditioning algorithm: effect of number of cycles, cycle duration and effector organ mass on efficacy of protection. *Basic Res Cardiol* 2016; **111:** 10.

HEALTH for heavy menstrual bleeding: real-world implications



Published Online September 12, 2019 http://dx.doi.org/10.1016/ S0140-6736(19)32086-0 See **Articles** page 1425

Heavy menstrual bleeding is common and can substantially affect quality of life.¹ Millions of women worldwide defer personal, professional, and family goals because of heavy menstrual bleeding.

In *The Lancet*, Kevin Cooper and colleagues² report a multicentre, parallel-group, open-label randomised trial of 660 women younger than 50 years that compared laparoscopic supracervical hysterectomy with endometrial ablation for the management of heavy menstrual bleeding. To our knowledge, the HEALTH trial is one of the largest randomised controlled trials to date in benign gynaecology. Society guidelines describe hysterectomy as a last resort in the context of providing less invasive options for gynaecological conditions.³ However, methodologically sound evaluation is required to test the status quo.

The HEALTH trial found that at 15 months after randomisation, women allocated laparoscopic supracervical hysterectomy were more satisfied with their treatment (270 [97%] of 278 women vs 244 [87%] of 280 women; adjusted percentage difference 9.8, 95% CI 5.1–14.5; adjusted odds ratio 2.53, 95% CI 1.83-3.48; p<0.0001) and reported improved quality of life compared with those who were allocated endometrial ablation (180 [69%] of 262 women vs 146 [54%] of 268 women; adjusted percentage difference 13.3, 95% CI 3.8-22.8; adjusted OR 1.87, 95% CI 1.31-2.67; p=0.00058).² There were no differences detected in serious adverse events between the two groups; however, mean operating (laparoscopic supracervical times hysterectomy 114 min [SD 38] vs endometrial ablation 44 min [23]), median time to discharge from operation (laparascopic supracervical hysterectomy 21.5 h [IQR 17.0–26.1] vs endometrial ablation 3.2 h [2.1-5.1]), and median time to return to work (laparascopic supracervical hysterectomy 42 days [37-42] vs endometrial ablation 10 days [7-14]) were longer following laparoscopic supracervical hysterectomy.

The real-world implications and worldwide generalisability of this study are where we must now turn our attention. Although this trial specifically examined

Articles

Effect of remote ischaemic conditioning on clinical outcomes in patients with acute myocardial infarction (CONDI-2/ERIC-PPCI): a single-blind randomised controlled trial

Derek J Hausenloy*, Rajesh K Kharbanda*, Ulla Kristine Møller, Manish Ramlall, Jens Aarøe, Robert Butler, Heerajnarain Bulluck, Tim Clayton, Ali Dana, Matthew Dodd, Thomas Engstrom, Richard Evans, Jens Flensted Lassen, Erika Frischknecht Christensen, José Manuel Garcia-Ruiz, Diana A Gorog, Jakob Hjort, Richard F Houghton, Borja Ibanez, Rosemary Knight, Freddy K Lippert, Jacob T Lønborg, Michael Maeng, Dejan Milasinovic, Ranjit More, Jennifer M Nicholas, Lisette Okkels Jensen, Alexander Perkins, Nebojsa Radovanovic, Roby D Rakhit, Jan Ravkilde, Alisdair D Ryding, Michael R Schmidt, Ingunn Skogstad Riddervold, Henrik Toft Sørensen, Goran Stankovic, Madhusudhan Varma, Ian Webb, Christian Juhl Terkelsen, John P Greenwood, Derek M Yellon†, Hans Erik Bøtker†, on behalf of the CONDI-2/ERIC-PPCI Investigators‡

Summary

Background Remote ischaemic conditioning with transient ischaemia and reperfusion applied to the arm has been shown to reduce myocardial infarct size in patients with ST-elevation myocardial infarction (STEMI) undergoing primary percutaneous coronary intervention (PPCI). We investigated whether remote ischaemic conditioning could reduce the incidence of cardiac death and hospitalisation for heart failure at 12 months.

Methods We did an international investigator-initiated, prospective, single-blind, randomised controlled trial (CONDI-2/ERIC-PPCI) at 33 centres across the UK, Denmark, Spain, and Serbia. Patients (age >18 years) with suspected STEMI and who were eligible for PPCI were randomly allocated (1:1, stratified by centre with a permuted block method) to receive standard treatment (including a sham simulated remote ischaemic conditioning intervention at UK sites only) or remote ischaemic conditioning treatment (intermittent ischaemia and reperfusion applied to the arm through four cycles of 5-min inflation and 5-min deflation of an automated cuff device) before PPCI. Investigators responsible for data collection and outcome assessment were masked to treatment allocation. The primary combined endpoint was cardiac death or hospitalisation for heart failure at 12 months in the intention-to-treat population. This trial is registered with ClinicalTrials.gov (NCT02342522) and is completed.

Findings Between Nov 6, 2013, and March 31, 2018, 5401 patients were randomly allocated to either the control group (n=2701) or the remote ischaemic conditioning group (n=2700). After exclusion of patients upon hospital arrival or loss to follow-up, 2569 patients in the control group and 2546 in the intervention group were included in the intention-to-treat analysis. At 12 months post-PPCI, the Kaplan-Meier-estimated frequencies of cardiac death or hospitalisation for heart failure (the primary endpoint) were 220 ($8 \cdot 6\%$) patients in the control group and 239 ($9 \cdot 4\%$) in the remote ischaemic conditioning group (hazard ratio $1 \cdot 10$ [95% CI $0 \cdot 91 - 1 \cdot 32$], p= $0 \cdot 32$ for intervention versus control). No important unexpected adverse events or side effects of remote ischaemic conditioning were observed.

Interpretation Remote ischaemic conditioning does not improve clinical outcomes (cardiac death or hospitalisation for heart failure) at 12 months in patients with STEMI undergoing PPCI.

Funding British Heart Foundation, University College London Hospitals/University College London Biomedical Research Centre, Danish Innovation Foundation, Novo Nordisk Foundation, TrygFonden.

Copyright © 2019 The Author(s). Published by Elsevier Ltd. This is an Open Access article under the CC BY 4.0 license.

Introduction

Despite timely reperfusion with primary percutaneous coronary intervention (PPCI), morbidity and mortality following acute ST-elevation myocardial infarction (STEMI) remain substantial, and improvements in mortality and survival free of heart failure have plateaued.¹² New treatments are needed to reduce myocardial infarct size and preserve cardiac function to reduce risk of death and prevent onset of heart failure. In this regard, remote ischaemic conditioning, in which brief cycles of ischaemia

and reperfusion are applied to an organ or tissue (including a limb) away from the heart, has been shown to reduce myocardial infarct size in animal models.^{2,3}

In the clinical setting, the cardioprotective remote ischaemic conditioning stimulus can be applied using serial inflations and deflations of a pneumatic cuff placed on the upper arm or thigh to induce brief cycles of ischaemia and reperfusion.⁴ In most clinical studies in patients with STEMI, remote ischaemic conditioning has increased myocardial salvage and reduced myocardial





Lancet 2019; 394: 1415-24

Published Online September 6, 2019 http://dx.doi.org/10.1016/ S0140-6736(19)32039-2

See Comment page 1389

*Joint first authors

†Joint senior authors ‡Investigators are listed in

appendix 1 The Hatter Cardiovascular Institute (Prof D J Hausenloy PhD,

M Ramlall MBChB Prof D M Yellon DSc) and Roval Free Hospital London and Institute of Cardiovascular Science (R D Rakhit MD), University College London. London, UK; National Institute of Health Research Biomedical Research Centre at University College London Hospitals, Research & Development, London, UK (Prof D J Hausenloy); Cardiovascular & Metabolic Disorders Program, Duke-National University of Singapore Medical School. Singapore (Prof D J Hausenloy); National Heart Research Institute Singapore, National Heart Centre, Singapore (Prof D J Hausenloy); Yong Loo Lin School of Medicine, National University Singapore. Singapore (Prof D J Hausenloy); Centro de Biotecnologia FEMSA, Tecnologico de Monterrey, Monterrey, Mexico (Prof D J Hausenloy); Oxford Heart Centre, Oxford University Hospitals National Health Service Trust, Oxford, UK (R K Kharbanda PhD); Department of Cardiovascular Medicine, University of Oxford, Oxford, UK (R K Kharbanda): Department of Cardiology

(U K Møller PhD, Prof LE Lassen PhD. M Maeng PhD, M R Schmidt PhD, C I Terkelsen DMSci. Prof H E Bøtker DMSci) and Department of Clinical Epidemiology (Prof H T Sørensen DMSci), Aarhus University Hospital. Aarhus, Denmark: University Hospital Southampton National Health Service Foundation Trust Southampton, UK (M Ramlall); Department of Cardiology, Aalborg University Hospital. Aalborg, Denmark (| Aarøe MD, J Ravkilde MD); Department of Cardiology, University Hospitals of North Midlands, **Royal Stoke University** Hospital, Stoke-on-Trent, UK (R Butler MD): Department of Cardiology, Norfolk and Norwich University Hospital, Norwich, UK (H Bulluck PhD, A D Ryding PhD); Clinical Trials Unit and Department of Medical Statistics, London School of Hygiene & Tropical Medicine, London, UK (Prof T Clayton MSc, M Dodd MSc, R Evans BA, R Knight MSc. I M Nicholas PhD. A Perkins MSc); Portsmouth Hospitals National Health Service Trust, Portsmouth, UK (A Dana PhD); Department of Cardiology, Rigshospitalet, University of Copenhagen, Copenhagen, Denmark (Prof T Engstrom DMSci, JT Lønborg DMSci); Prehospital **Emergency Medical Services**, Central Denmark Region, Aarhus, Denmark (Prof E F Christensen MD, I S Riddervold PhD); Instituto de Investigación Sanitaria del Principado de Asturias, Hospital Universitario de Cabueñes, Oviedo, Spain (J M Garcia-Ruiz MD); Centro Nacional de Investigaciones Cardiovasculares. Madrid, Spain (J M Garcia-Ruiz, Prof B Ibanez PhD); Centro de Investigación Biomédica en Red de Enfermedades Cardiovasculares, Madrid, Spain (J M Garcia-Ruiz); Department of Cardiology. Lister Hospital, East and North Hertfordshire National Health Service Trust, Stevenage, UK (Prof D A Gorog PhD); National Heart and Lung Institute, Imperial College London, London, UK (Prof D A Goroa): Department of Clinical

Medicine, Aarhus University,

Research in context

Evidence before this study

To improve clinical outcomes in patients with ST-elevation myocardial infarction (STEMI) treated by primary percutaneous coronary intervention (PPCI), new treatments are needed to protect the heart against acute ischaemia-reperfusion injury and thus reduce myocardial infarct size and prevent heart failure. Remote ischaemic conditioning with transient ischaemia and reperfusion of the arm or leg as an adjunct to PPCI has been shown in several small randomised controlled trials to reduce myocardial infarct size and increase myocardial salvage in patients with STEMI when compared with use of PPCI alone. However, whether remote ischaemic conditioning can improve clinical outcomes such as mortality and heart failure has not been confirmed. A PubMed search undertaken up to the start date of the trial (July 14, 2013) using the terms "remote ischemic conditioning; acute myocardial infarction; mortality; and heart failure" revealed no randomised controlled trials investigating the effect of remote ischaemic conditioning on clinical outcomes in patients with STEMI treated by PPCI.

Added value of this study

In this large, sufficiently powered, multicentre, randomised controlled trial of 5401 patients with STEMI, we investigated whether remote ischaemic conditioning applied as an adjunct to PPCI could reduce the incidence of cardiac death or hospitalisation for heart failure within 12 months post-PPCI, when compared with PPCI alone. Our findings showed that remote ischaemic conditioning did not improve clinical outcomes in terms of cardiac death or hospitalisation for heart failure at this timepoint. Furthermore, no improvements in any of the secondary endpoints were found, including major adverse

infarct size by 20-30% when applied before or during reperfusion.5-12 In addition, two follow-up studies and a prospective single-centre study have suggested that remote ischaemic conditioning might also improve clinical outcomes in patients with STEMI undergoing PPCI.¹³⁻¹⁵ However, a large, sufficiently powered, prospectively designed multicentre clinical outcome study has not yet been done. Therefore, in the CONDI-2/ ERIC-PPCI trial, we investigated the effect of remote ischaemic conditioning applied as an adjunct to PPCI on rates of cardiac death or hospitalisation for heart failure at 12 months in patients with STEMI. In this study, we combined the CONDI-2 and ERIC-PPCI trials and harmonised the protocols to increase sample size and ensure the study was sufficiently powered to detect an effect of remote ischaemic conditioning on clinical outcomes in patients with STEMI treated by PPCI.

Methods

Study design

We did an international, multicentre, single-blind, randomised controlled trial at 26 centres in the UK (the ERIC-PPCI component study), and at four hospitals in cardiovascular and cerebral events and myocardial infarct size evaluated by troponin T release. Furthermore, no effects of the intervention versus the control were seen among prespecified subgroup analyses by age, presence of diabetes, pre-PPCI coronary flow, ischaemia time, or infarct location.

Implications of all the available evidence

The CONDI-2/ERIC-PPCI trial has provided definitive evidence that remote ischaemic conditioning offers no benefits on either myocardial infarct size or clinical outcomes at 12 months in patients with STEMI treated with PPCI. Two follow-up randomised controlled trials and a prospective single-centre randomised controlled trial, published since the commencement of our trial, had suggested that remote ischaemic conditioning might improve clinical outcomes in patients with STEMI undergoing PPCI. The two follow-up trials were not prospectively designed or powered to detect a difference in clinical outcomes with remote ischaemic conditioning. The prospective single-centre trial might not have been sufficiently powered, and it had an extended follow-up period. Until now, remote ischaemic conditioning had been the most promising potential cardioprotective strategy for improving clinical outcomes following STEMI. Therefore, identification of novel cardioprotective targets and discovery of innovative approaches to cardioprotection are needed to improve clinical outcomes in patients with STEMI treated by PPCI. Such approaches might include combination multitarget therapy. Remote ischaemic conditioning might still be beneficial in other clinical settings of acute ischaemia-reperfusion injury, including renal transplantation, acute kidney injury, and stroke.

Denmark, two hospitals in Spain, and one hospital in Serbia (the CONDI-2 component study). The CONDI-2/ ERIC-PPCI study received ethical approval from regional and national health service research ethics committees and was conducted in accordance with the principles of good clinical practice. In the CONDI-2 component study, all participants provided written informed consent before randomisation. In the ERIC-PPCI component study, all patients provided initial verbal assent before randomisation, which was followed by written informed consent. The London School of Hygiene & Tropical Medicine Clinical Trials Unit (London, UK) coordinated the trial in collaboration with the Cardiology Trial Unit and Department of Clinical Epidemiology of Aarhus University Hospital (Aarhus, Denmark). Details of the trial design have been reported previously,16 and a copy of the protocol is available in appendix 2.

Participants

Patients with chest pain and suspected ST-segment elevation on electrocardiogram (ECG) were screened for possible inclusion. Patients were included if they were older than 18 years of age, had ST-segment elevation on ECG, and were eligible for PPCI. Exclusion criteria were previous coronary artery bypass graft surgery, myocardial infarct within the previous 30 days, left bundle branch block on ECG, treatment with therapeutic hypothermia, conditions precluding use of remote ischaemic conditioning (paresis of upper limb, or presence of an arteriovenous shunt), and life expectancy of less than 1 year due to a non-cardiac pathology.

Randomisation and masking

In both trials, patients were randomly allocated (1:1) to a remote ischaemic conditioning group or a control group by a designated study team member who was unmasked to treatment allocation. For ERIC-PPCI, randomisation was done via a secure website, Sealed Envelope (London, UK), and was stratified by centre with use of random permuted blocks (block sizes of 4 or 6). For CONDI-2, randomisation was done through a web-based clinical trial support system accessible 24 h a day (TrialPartner, Aarhus, Denmark) and stratified by centre using random permuted blocks (block sizes of 4, 6, or 8). In both trials, access to the randomisation website and list were strictly controlled at each site and limited to unmasked study team members. Study team members collecting the data and assessing outcomes were masked to the treatment allocation.

Procedures

An automated AutoRIC cuff device (CellAegis Devices, Toronto, ON, Canada) was used to deliver the remote ischaemic conditioning protocol, which comprised four alternating cycles of cuff inflation for 5 min to 200 mm Hg and deflation for 5 min. At the UK sites, the control group received a sham simulated remote ischaemic conditioning intervention. At the sites in Denmark, Spain, and Serbia, the control group received standard therapy. The remote ischaemic conditioning and control protocols were implemented before PPCI and did not delay the onset of PPCI. Where ambulance transit times allowed, remote ischaemic conditioning was administered in the ambulance (in Denmark and Spain). Where ambulance transit times were shorter, remote ischaemic conditioning was delivered immediately upon arrival at the PPCI centre (in Serbia and the UK).

Before PPCI, patients were treated with 300 mg aspirin orally or intravenously, 600 mg clopidogrel or 180 mg ticagrelor orally, and standard weight-adjusted heparin intravenously. The PPCI procedure followed guideline recommendations;⁷⁷ ad hoc thrombectomy and administration of glycoprotein IIb/IIIa inhibitors, bivalirudin, or cangrelor were per operator discretion.

Outcomes

The primary combined endpoint was cardiac death or hospitalisation for heart failure at 12 months postrandomisation in the intention-to-treat population. Hospitalisation for heart failure included heart failure both during the index hospitalisation and during rehospitalisation for heart failure. Hospitalisation was defined as treatment occurring during a hospital admission. Heart failure was judged to be present on the basis of at least one of the following symptoms and signs: new or worsening dyspnoea, orthopnoea, or paroxysmal nocturnal dyspnoea; increasing fatigue or worsening exercise tolerance; ejection fraction less than 45% on echocardiography; new pulmonary oedema seen on chest x-ray in the absence of a non-cardiac cause; crepitations believed to be due to pulmonary oedema; and use of loop diuretics to treat presumed pulmonary congestion.18 A blinded independent event validation committee reviewed all events according to standard operational procedures. Hospitalisation for heart failure was defined as an event that met the standard criteria18 (appendix 1 p 20).

Prespecified secondary endpoints, analysed in the intention-to-treat population, included combined cardiac death or hospitalisation for heart failure at 30 days; cardiac death and hospitalisation for heart failure (as individual endpoints) at 30 days and 12 months; major cardiovascular and cerebral adverse events (comprising all-cause death, reinfarction, repeat coronary revascularisation, and stroke; appendix 1 p 20) at 30 days and 12 months; incidence of implantable cardioverter-defibrillator insertion within 12 months; repeated episodes of hospitalisation for heart failure within 12 months; and myocardial infarct size (quantified by area under the curve [AUC] for high-sensitivity troponin T measured at 0–48 h after PPCI) in a subset of patients.

Statistical analysis

The primary outcome event rate was originally predicted to be 11.0% at 12 months. To show a 25% relative reduction in the remote ischaemic conditioning group (from 11.0% to 8.25%) with 80% power and 5% significance would have required 4300 patients (2000 patients in ERIC-PPCI and 2300 in CONDI-2), allowing for 15% attrition. A blinded review of accruing data revealed that the estimated event rate was lower than anticipated and the sample size was, therefore, increased to 5400 patients (2800 patients in ERIC-PPCI and 2600 in CONDI-2).

The primary and secondary time-to-event outcomes were compared between the remote ischaemic conditioning and control groups with use of a Cox regression model stratified by the two component studies on an intention-to-treat basis, and presented with Kaplan-Meier curves. To account for recurrent hospitalisation for heart failure and competing mortality risk, a negative binomial model was used to compare the total number of outcomes experienced by the two groups up to 12 months after randomisation. The proportion of patients with an implantable cardioverterdefibrillator, biventricular pacemaker, or single-chamber or dual-chamber pacemaker at 12 months was compared

Aarhus, Denmark (J Hjort MPH); Servicio de Atención Médica de Urgencia—Asturias, Oviedo, Spain (R F Houghton MD): Centro de Investigacion **Biomedica** En Red Cardiovascular, Madrid, Spain (Prof B Ibanez); IIS-Fundación Jiménez Díaz University Hospital, Madrid, Spain (Prof B Ibanez); Prehospital **Emergency Medical Services**, Capital Region of Denmark, Denmark (FK Lippert MD): Department of Cardiology (D Milasinovic MD), Cardiology Clinic (Prof N Radovanovic PhD Prof G Stankovic PhD), Emergency Centre (Prof N Radovanovic), and Department for Diagnostic and **Catheterization Laboratories** (Prof G Stankovic). Clinical Centre of Serbia, Faculty of Medicine, University of Belgrade, Belgrade, Serbia: Lancashire Cardiac Centre, Blackpool Teaching Hospitals National Health Service Foundation Trust, Blackpool, UK (R More MBChB): Department of Cardiology, Odense University Hospital, Odense, Denmark (Prof L O Jensen DMSci); The Heart Centre North Cumbria University Hospitals National Health Service Trust. Carlisle, UK (M Varma PhD): King's College Hospital, King's Health Partnership, London, UK (I Webb PhD); Leeds Institute of Cardiovascular and Metabolic Medicine, University of Leeds, Leeds, UK (Prof I P Greenwood PhD): and Leeds Teaching Hospitals National Health Service Trust. Leeds, UK (Prof J P Greenwood)

Correspondence to: Prof Derek J Hausenloy, Hatter Cardiovascular Institute, University College London, London WC1E 6HX, UK d.hausenloy@ucl.ac.uk

or

Prof Hans Erik Bøtker, Department of Cardiology, Aarhus University Hospital, 8200 Aarhus N, Denmark haboet@rm.dk

See Online for appendix 1

See Online for appendix 2 For the Sealed Envelope website see https://www. sealedenvelope.com/

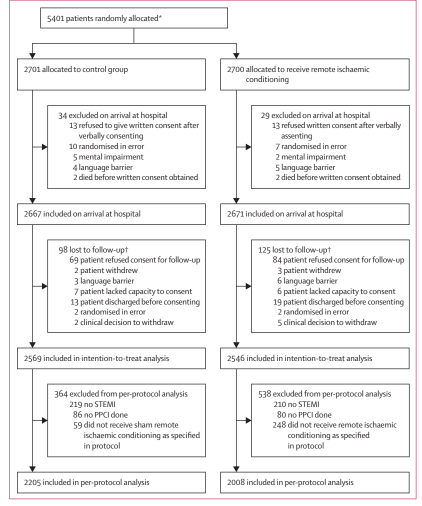


Figure 1: Trial profile

PPCI=primary percutaneous coronary intervention. STEMI=ST-elevation myocardial infarction. *Full screening log data were not available as not all sites were able to collect screening logs given the emergency context of patient enrolment and randomisation. †ERIC-PPCI had approval from the UK Confidentiality Group to collect data on patients who died before consent could be obtained (20 patients in the control group and 22 patients in the remote ischaemic conditioning group); thus, these patients were included in subsequent analyses.

using a generalised linear model for a binary outcome with a log link.

Prespecified analyses by subgroup (age [<55 years, 55 to <65 years, 65 to <75 years, or ≥75 years], diabetic status, infarct location [left anterior descending artery or other], pre-angioplasty TIMI flow grade [0–1 or 2–3], and time elapsed between first medical contact and PPCI [<90 min, 90 to <120 min, or 120 to 720 min]) were done on the primary outcome. The subgroups were analysed by including an interaction term between treatment group and subgroup in the Cox regression model.

Troponin T values were converted into a 48-h AUC summary measure for analysis, using multiple imputation by chained equations to account for missing data. As the distribution of the outcome variable was highly skewed, a log transformation was done. Differences were analysed by use of a linear regression model with the treatment effect presented as a ratio of the geometric means.

We also did a per-protocol analysis of all primary and secondary outcomes limited to patients who had a confirmed STEMI, received the full intervention as specified in the protocol, and had a completed PPCI.

Results were considered statistically significant if the two-sided p value was less than 0.05. Full details of the statistical methods are provided in the protocol and statistical analysis plan, which are available in appendix 2.

The CONDI-2/ERIC-PPCI trial is registered at ClinicalTrials.gov (number NCT02342522) and is a planned prespecified collaboration between the CONDI-2 and ERIC-PPCI studies (for further details see appendix 1 pp 20–21).

Role of the funding source

The sponsor had no role in the study design, data collection, data analysis, data interpretation, or writing of the report. The co-corresponding authors had full access to all the data in the study and had final responsibility for the decision to submit for publication.

Results

Between Nov 6, 2013, and March 31, 2018, 5401 patients were randomly allocated: 2701 to the control group and 2700 to the remote ischaemic conditioning group. 5115 patients (2569 in the control group and 2546 in the remote ischaemic conditioning group) were included in the intention-to-treat analysis (figure 1). The treatment groups were well balanced with respect to both patient baseline characteristics and PPCI details for the intention-to-treat analysis (table 1). The intervention was completed according to protocol for 2205 participants in the control group and 2008 in the remote ischaemic conditioning group, and the results were included in the per-protocol analysis. Reasons for incomplete interventions are provided in figure 1.

There was no evidence of difference between the control group (8.6% [n=220]) and the remote ischaemic conditioning group (9.4% [n=239]) with respect to the combined primary end point of cardiac death or hospitalisation for heart failure at 12 months (hazard ratio [HR] 1.10; 95% CI 0.91-1.32; p=0.32; table 2, figure 2). Similarly, the control and remote ischaemic conditioning groups did not differ with regard to the individual components of cardiac death or hospitalisation for heart failure at 12 months (table 2, figure 2). The primary and secondary outcomes in CONDI-2 and ERIC-PPCI were similar, with no effect of remote ischaemic conditioning seen in either study (appendix 1 pp 25–30, 32–37).

Prespecified subgroup analyses showed no difference in the effect of remote ischaemic conditioning on the primary outcome by age, presence of diabetes, pre-PPCI TIMI flow grade, time from first medical contact to PPCI, or infarct location (figure 3). The results of the perprotocol analysis were very similar to those of the

| | Control group (n=2569) | Remote ischaemic conditioning group (n=2546) | |
|--|---------------------------|--|--|
| Baseline characteristics | | | |
| Age, years | 63·1 (12·2) | 63.9 (12.1) | |
| Sex | | | |
| Female | 576/2569 (22.4%) | 611/2546 (24.0%) | |
| Male | 1993/2569 (77.6%) | 1935/2546 (76.0%) | |
| Body-mass index (kg/m²) | 27.5 (4.7) | 27.5 (4.9) | |
| Current smoker | 1015/2483 (40.9%) | 993/2449 (40·5%) | |
| Comorbidities | | | |
| Hypertension | 1020/2544 (40·1%) | 1100/2518 (43.7%) | |
| Previous acute myocardial infarction | 253/2553 (9.9%) | 265/2530 (10.5%) | |
| Hypercholesterolaemia | 682/2503 (27-2%) | 696/2490 (28%) | |
| Medically treated diabetes | 265/2557 (10.4%) | 302/2534 (11·9%) | |
| Family history of ischaemic heart disease | 819/2387 (34·3%) | 853/2337 (36.5%) | |
| Medications at admission fo | or PPCI | | |
| Sulfonylurea | 62/2519 (2.5%) | 61/2496 (2.4%) | |
| Metformin | 196/2518 (7.8%) | 221/2499 (8.8%) | |
| Insulin | 74/2548 (2.9%) | 81/2533 (3.2%) | |
| Clopidogrel | 85/2548 (3.3%) | 98/2528 (3.9%) | |
| Ticagrelor | 52/2549 (2.0%) | 62/2527 (2.5%) | |
| Prasugrel | 4/2546 (0.2%) | 3/2528 (0.1%) | |
| Statins | 652/2547 (25.6%) | 665/2526 (26-3%) | |
| Aspirin | 456/2548 (17.9%) | 485/2528 (19·2%) | |
| β blockers | 384/2539 (15·1%) | 407/2525 (16·1%) | |
| Angiotensin-converting enzyme inhibitors or angiotensin receptor blockers | 456/2541 (17·9%) | 466/2525 (18·5%) | |
| Blood pressure at randomis | ation, mm Hg | | |
| Systolic | 131 (24) | 131 (24) | |
| Diastolic | 77 (15) | 76 (15) | |
| Killip class at randomisatior | 1 | | |
| I | 2462/2568 (95.9%) | 2435/2546 (95.6%) | |
| II | 75/2568 (2.9%) | 75/2546 (2·9%) | |
| Ш | 14/2568 (0.5%) | 11/2546 (0.4%) | |
| IV (including cardiogenic shock) | 17/2568 (0.7%) | 25/2546 (1.0%) | |
| Procedural details | | | |
| Number of complete cycles administered | of remote ischaemic co | nditioning | |
| 1 | NA | 37/2246 (1.6%) | |
| 2 | NA | 40/2246 (1.8%) | |
| 3 | NA | 142/2246 (6·3%) | |
| 4 | NA | 2027/2246 (90·2%) | |
| TIMI flow grade at admissic | in | | |
| 0 | 1513/2227 (67.9%) | 1478/2203 (67.1%) | |
| 1 | 154/2227 (6.9%) | 145/2203 (6.6%) | |
| 2 | 218/2227 (9.8%) | 225/2203 (10·2%) | |
| 3 | 342/2227 (15.4%) | 355/2203 (16·1%) | |
| 3 | | 355/2203 (16 ntinues in next co | |

| | Control group (n=2569) | Remote ischaemic conditioning grou (n=2546) | | | | |
|---|----------------------------|---|--|--|--|--|
| (Continued from previous c | olumn) | | | | | |
| Symptom to balloon time, min | 177 (128–279) | 178 (130–278) | | | | |
| First medical contact to balloon time, min | 102 (82–126) | 103 (83–128) | | | | |
| Infarct-related coronary artery | | | | | | |
| Left anterior descending | 974/2258 (43·1%) | 911/2226 (40·9% | | | | |
| Circumflex | 297/2258 (13·2%) | 298/2226 (13·4%) | | | | |
| Right coronary | 985/2258 (43·6%) | 1014/2226 (45.6% | | | | |
| Other | 2/2258 (0.1%) | 3/2226 (0.1%) | | | | |
| Number of vessels with clin | ically significant disease | 2 | | | | |
| 0 | 1/2258 (<0.1%) | 2/2227 (0.1%) | | | | |
| 1 | 1314/2258 (58-2%) | 1274/2227 (57·2%) | | | | |
| 2 | 659/2258 (29·2%) | 641/2227 (28.8% | | | | |
| 3 | 284/2258 (12.6%) | 310/2227 (13·9%) | | | | |
| Stenting of culprit lesion by PPCI | 2104/2258 (93·2%) | 2080/2226 (93·4% | | | | |
| Aspiration thrombectomy | 560/2242 (24·8%) | 553/2209 (24.8% | | | | |
| Supplementary staged PPCI | 291/2258 (12.9%) | 261/2227 (11.7%) | | | | |
| Supplementary staged coronary artery bypass grafting | 52/2258 (2·3%) | 62/2227 (2.8%) | | | | |
| TIMI flow grade after procee | dure | | | | | |
| 0 | 27/2223 (1.2%) | 26/2200 (1.2%) | | | | |
| 1 | 20/2223 (0.9%) | 9/2200 (0.4%) | | | | |
| 2 | 112/2223 (5.0%) | 86/2200 (3.9%) | | | | |
| 3 | 2064/2223 (92.8%) | 2079/2200 (94.5% | | | | |
| Medications given in relation | in to PPCI | | | | | |
| Opioids* | 549/1054 (52·1%) | 522/1040 (50-2%) | | | | |
| Heparin | 1904/2255 (84·4%) | 1893/2224 (85.1%) | | | | |
| Aspirin | 2147/2255 (95·2%) | 2129/2221 (95·9%) | | | | |
| Clopidogrel | 610/2256 (27.0%) | 573/2226 (25.7%) | | | | |
| Ticagrelor | 1551/2257 (68.7%) | 1554/2227 (69.8%) | | | | |
| Prasugrel | 103/2256 (4.6%) | 97/2226 (4·4%) | | | | |
| Nitrates | 1758/2222 (79·1%) | 1693/2185 (77·5%) | | | | |
| Glycoprotein Ilb/IIIa inhibitor | 450/2250 (20.0%) | 404/2223 (18·2%) | | | | |
| Bivalirudin | 505/2250 (22.4%) | 489/2224 (22·0%) | | | | |
| Cangrelor* | 134/1088 (12·3%) | 127/1085 (11.7%) | | | | |
| Data are n/N (%), mean (SD), o provided in appendix 1 (pp 37- intervention. NA=not applicab | 39). PPCI=primary percut | taneous coronary | | | | |
| Table 1: Baseline characteris intention-to-treat populat | • | etails for the | | | | |

intention-to-treat analysis for the primary outcome (appendix 1 p 24), with the primary combined endpoint within 12 months occurring in 179 (9.0%) participants in the remote ischaemic conditioning group versus 178 (8.1%) in the control group (HR 1.11 [95% CI 0.90-1.36], p=0.35).

With respect to major cardiovascular and cerebral adverse events within 12 months of follow-up, no

| | | Outcomes* | | Treatment effect† (95% CI) | p value |
|---|--|---------------------------|---|-------------------------------|---------|
| | | Control group (n=2569) | Remote ischaemic conditioning group (n=2546) | | |
| т | ime-to-event outcomes | | | | · |
| h | ombined cardiac death or ospitalisation for heart failure within 2 months (primary outcome) | 220 (8.6%) | 239 (9·4%) | 1.10 (0.91–1.32) | 0.32 |
| | Cardiac death within 12 months | 69 (2·7%) | 77 (3·1%) | 1.13 (0.82–1.56) | 0.46 |
| | Hospitalisation for heart failure within 12 months | 182 (7.1%) | 192 (7.6%) | 1.06 (0.87–1.30) | 0.55 |
| | 1ajor cardiovascular and cerebral dverse events‡ within 12 months | 197 (7.8%) | 212 (8.4%) | 1.09 (0.90–1.32) | 0.38 |
| | All-cause death within 12 months | 100 (3.9%) | 122 (4.8%) | 1·24 (0·95–1·61) | 0.11 |
| | Reinfarction within 12 months | 43 (1·7%) | 38 (1.5%) | 0.89 (0.58–1.38) | 0.61 |
| | Unplanned revascularisation within 12 months | 61 (2·4%) | 61 (2.5%) | 1.02 (0.71–1.45) | 0.93 |
| | Stroke within 12 months | 21 (0.8%) | 23 (0.9%) | 1.11 (0.61–2.00) | 0.73 |
| | ombined hospitalisation for heart ailure or cardiac death within 30 days | 185 (7·2%) | 204 (8.0%) | 1.11 (0.91–1.36) | 0.29 |
| | Cardiac death within 30 days | 47 (1.8%) | 59 (2·3%) | 1.27 (0.87–1.86) | 0.22 |
| | Hospitalisation for heart failure within 30 days | 162 (6·3%) | 171 (6.8%) | 1.06 (0.86–1.32) | 0.57 |
| | 1ajor cardiovascular and cerebral dverse events‡ within 30 days | 97 (3.8%) | 109 (4·3%) | 1.14 (0.87–1.50) | 0.35 |
| | All-cause death within 30 days | 52 (2.0%) | 63 (2.5%) | 1.23 (0.85–1.77) | 0.27 |
| | Reinfarction within 30 days | 21 (0.8%) | 19 (0.8%) | 0.92 (0.49–1.70) | 0.78 |
| | Unplanned revascularisation within 30 days | 29 (1.1%) | 29 (1·2%) | 1.02 (0.61–1.70) | 0.95 |
| | Stroke within 30 days | 7 (0·3%) | 10 (0.4%) | 1.44 (0.55–3.80) | 0.46 |
| | nplantable cardioverter-defibrillator nplantation within 12 months | 28 (1.1%) | 37 (1.5%) | 1.34 (0.82–2.18) | 0.24 |
| E | vent frequency outcomes | | | | |
| h | lumber of repeat episodes of ospitalisation for heart failure within 2 months | 0.09 (0.36) | 0.09 (0.34) | 1.00 (0.81–1.24) | 1.00 |
| h | lumber of repeat episodes of ospitalisation for heart failure or ardiac death within 12 months | 0.11 (0.43) | 0.12 (0.41) | 1.03 (0.85–1.25) | 0.77 |
| | | | | | |

*Data are Kaplan-Meier estimates of the n (%) of patients with the outcome at the specified timepoint (for time-to-event outcomes), or mean (SD; for event frequency outcomes). †Data are hazard ratios (for time-to-event outcomes) or ratio of means (for event frequency outcomes), for treatment group versus control group. ‡Composite of all-cause death, reinfarction, unplanned revascularisation, and stroke.

Table 2: Primary and secondary outcomes in the intention-to-treat population

difference was observed between the remote ischaemic conditioning group and the control group, for either the combined outcome or the individual components (table 2).

No differences between the control and remote ischaemic conditions group were observed with regard to cardiac death or hospitalisation for heart failure (both combined and individually) within 30 days, major cardiovascular and cerebral adverse events (or the individual components of this outcome) after 30 days, or incidence of implantable cardioverter-defibrillator implantation within 12 months (table 2). Numbers of repeat episodes of hospitalisation for heart failure, or repeat episodes of hospitalisation for heart failure plus cardiac death were also similar between groups (table 2). Secondary endpoints were very similar between the CONDI-2 and ERIC-PPCI trials (appendix 1 pp 45, 57).

The AUC for high-sensitivity troponin T did not differ between the control group (median $91.9 \text{ ng} \times h/mL$ [IQR 37.6-185.6], n=1330) and the remote ischaemic conditioning group (110.3 ng \times h/mL [48.0-218.5], n=1332; ratio of means 1.05 [95% CI 0.92-1.18], p=0.48; appendix 1 p 66).

No unexpected adverse events related to the trial treatment were reported. Skin petechiae and transient pain or paraesthesia were considered expected adverse events of remote ischaemic conditioning. In the remote ischaemic conditioning group, skin petechiae were reported for 72 (2.8%) patients and transient pain or paraesthesia was reported for 147 (5.8%) patients. There were no withdrawals due to adverse events.

Discussion

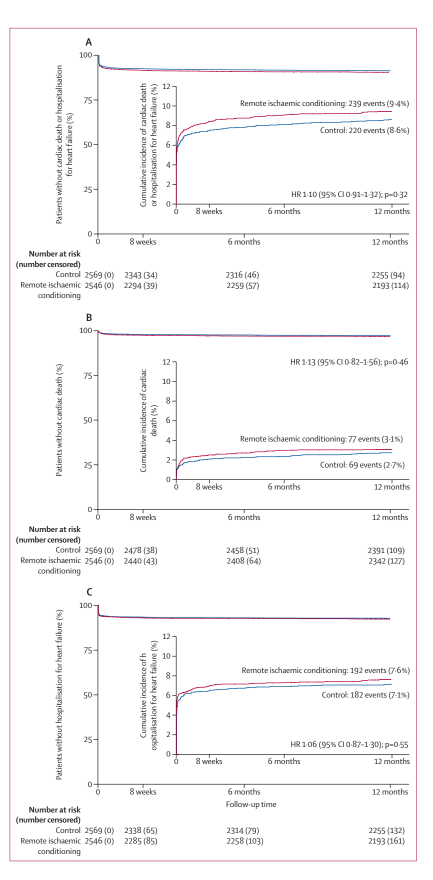
In this large, appropriately powered, prospective, international, multicentre trial, we found no clinically meaningful beneficial effects of remote ischaemic conditioning administered as an adjunct to PPCI on clinical outcomes (cardiac death or hospitalisation for heart failure) at 12 months in patients with STEMI when compared with PPCI alone. Furthermore, remote ischaemic conditioning had no effect on major secondary endpoints, including myocardial infarct size assessed by cardiac biomarkers.

These findings are in direct contrast to previously published clinical studies of remote ischaemic conditioning in patients with STEMI that reported increased myocardial salvage (on nuclear myocardial perfusion imaging and cardiac MRI) and reductions in myocardial infarct size (assessed by cardiac biomarkers and cardiac imaging),⁵⁻¹² although not all studies have been positive for these endpoints.¹² In our initial proofof-concept CONDI-1 trial, no statistically significant reduction in myocardial infarct size (assessed by cardiac biomarkers) was observed despite an increase in myocardial salvage.5 Measurements of cardiac biomarkers might not be a sufficiently sensitive marker for detecting reductions in myocardial infarct size with cardioprotective therapies, and myocardial salvage might be a more sensitive marker to assess cardioprotection. In our cardiac biomarker analysis of a subset of 2662 patients, we found no effect of remote ischaemic conditioning on myocardial infarct size assessed by troponin T release, confirming no biological effects of remote ischaemic conditioning, a finding which is consistent with the observed absence of effect on clinical outcomes at 12 months. The lack of effect observed in our study raises the possibility that previously published, smaller studies, which used similar remote ischaemic conditioning protocols, were subject to type 1 errors.5-12

Although myocardial infarct size is a known independent determinant of clinical outcomes post-PPCI in patients with STEMI, it is unclear whether a reduction in myocardial infarct size by a cardioprotective intervention applied as an adjunct to PPCI can be translated into improved clinical outcomes. The RIC-STEMI trial by Gaspar and colleagues15 also showed no reduction in myocardial infarct size (based on 48 h troponin I AUC), but still found improved clinical outcomes with remote ischaemic conditioning in terms of fewer cardiac deaths or hospitalisations for heart failure after a median followup time of 2.1 years (HR 0.35 [95% CI 0.15-0.78]). The unexpected and discordant effects of remote ischaemic conditioning on myocardial infarct size and clinical outcomes in the RIC-STEMI trial¹⁵ might have been due to a type 1 error, as only 516 patients with STEMI were randomly allocated, and the numbers of cardiac mortality events (three [1%] in the remote ischaemic conditioning group vs 11 [5%] in the control group) and hospitalisation for heart failure events (eight [3%] in the remote ischaemic conditioning group vs 17 [8%] in the control group) were small. An alternative explanation could be that the primary effect of remote ischaemic conditioning was on post-STEMI left ventricular remodelling rather than acute myocardial infarct sizealthough this mechanism would contradict experimental animal studies, which have clearly shown a beneficial effect of remote ischaemic conditioning on reducing acute myocardial infarct size. Notably, a clinical study investigating the cardioprotective effects of ischaemic postconditioning in patients with STEMI treated by PPCI showed no effect on myocardial infarct size, but still found reduced severity of adverse left ventricular remodelling at 1 year of follow-up, especially in patients with microvascular obstruction.19

Follow-up (median 3.8 years) of participants in the initial CONDI-1 trial⁵ (n=333 patients with STEMI) showed that increased myocardial salvage with remote ischaemic conditioning was associated with reduced frequencies of major cardiovascular and cerebral adverse events (17 [13.5%] patients) compared with the control group (32 [25.6%] patients).13 The LIPSIA CONDITIONING trial¹⁰ (n=696 patients with STEMI) showed improved myocardial salvage in patients who received PPCI combined with remote ischaemic conditioning and ischaemic postconditioning compared with either the control group (patients who received PPCI alone) or patients receiving ischaemic postconditioning with PPCI. Follow-up of LIPSIA CONDITIONING trial participants at a median of 3.6 years after the index event also revealed that major adverse cardiac events (cardiac death, reinfarction, and

Figure 2: 12-month cumulative incidence of combined cardiac death or hospitalisation for heart failure (A), cardiac death (B), and hospitalisation for heart failure (C) in the intention-to-treat population HR=hazard ratio.



| | Events, n/N (%)* | | Hazard ratio (95% CI) | Interaction p value |
|----------------------------|------------------|-------------------------------|--|------------------------|
| | Control | Remote ischaemic conditioning | | |
| Age (years) | | | | |
| <55 | 24/692 (3.5) | 22/628 (3.5) | 1.00 (0.56–1.78) | 0.70 |
| 55-<65 | 40/761 (5·3) | 48/688 (7.0) | 1.33 (0.87–2.02) | |
| 65-<75 | 58/624 (9·4) | 65/724 (9.0) | 0.97 (0.68–1.38) | |
| ≥75 | 79/472 (16·9) | 85/484 (17.9) | 1·05 (0·77–1·43) | |
| Medically treated diabete | s | | | |
| No | 179/2292 (7.9) | 193/2232 (8.7) | 1.11 (0.91–1.36) | 0.60 |
| Yes | 38/265 (14.6) | 43/302 (14·5) | 0.98 (0.63-1.51) | |
| TIMI flow grade at admiss | ion | | | |
| 0–1 | 156/1696 (9·2) | 169/1657 (10·3) | 1.11 (0.89–1.38) | 0.92 |
| 2-3 | 36/632 (5·7) | 42/647 (6.5) | 1.14 (0.73–1.78) | |
| Infarct location | | | | |
| Left anterior descending | 132/1001 (13.3) | 147/939 (15.7) | 1.21 (0.96–1.53) | 0.33 |
| Other | 65/1319 (5.0) | 66/1351 (4.9) | 0.98 (0.70–1.39) | |
| First medical contact to b | alloon time, min | | | |
| <90 | 46/742 (6·2) | 56/726 (7.7) | 1.24 (0.84–1.84) | 0.67 |
| 90 to <120 | 63/795 (7.9) | 60/733 (8-2) | 1·01 (0·71–1·44) | |
| 120 to 720 | 73/672 (10.9) | 75/697 (10.8) | 1.01 (0.73-1.39) | |
| Overall | 220/2569 (8.6) | 239/2546 (9.4) | 1.10 (0.91–1.32) | |
| | | | 0.5 1.0 1.5 2.0 | |
| | | | Favours remote Favours ischaemic conditioning control | |

Figure 3: Forest plot for prespecified subgroup analyses of the primary endpoint in the intention-to-treat population *Kaplan-Meier estimates of the number of patients with hospitalisation for heart failure or cardiac death within 12 months.

new congestive heart failure) were reduced in the group that received combined remote ischaemic conditioning and ischaemic postconditioning (23 [10·2%] patients) compared with the control group (37 [16·9%] patients) an effect that was mainly driven by a reduced rate of heart failure.¹⁴ However, these studies were not prospectively designed or powered to assess clinical outcomes of remote ischaemic conditioning following STEMI. The present study is almost ten-times larger than any previous study and was adequately powered to address the efficacy of remote ischaemic conditioning on clinical outcomes.

The reasons for the failure of remote ischaemic conditioning to reduce myocardial infarct size and improve clinical outcomes in our study are not clear. One potential reason could relate to the remote ischaemic conditioning protocol itself. Although four 5-min cycles of upper arm cuff inflations and deflations applied before PPCI has been shown to increase the myocardial salvage index⁵ and reduce myocardial infarct size in previous clinical studies,11 whether this protocol is the most effective cardioprotective intervention in terms of cycle numbers and duration is not known. One experimental study in mice suggested that two cycles of remote ischaemic conditioning were ineffective, whereas four cycles of remote ischaemic conditioning reduced myocardial infarct size to a similar extent to that achieved with six or eight cycles.20 However, such phase 2 doseresponse studies in humans are absent. The RIC-STEMI study,15 which showed no reduction in myocardial infarct size despite improved clinical outcomes, used a remote ischaemic conditioning stimulus comprising three cycles of inflation and deflation of a pneumatic cuff placed on the thigh, in contrast to our remote ischaemic conditioning stimulus that consisted of four cycles of inflation and deflation of a pneumatic cuff placed on the upper arm. The stimulus used in the RIC-STEMI trial might be expected to be more efficacious because of the greater ischaemic tissue burden, but this explanation would not account for the absence of effect on myocardial infarct size in the RIC-STEMI trial. Furthermore, a previous experimental study in mice showed that bilateral hindlimb remote ischaemic conditioning was no more efficacious at reducing myocardial infarct size than was unilateral limb remote ischaemic conditioning.²⁰

The timing of the remote ischaemic conditioning protocol in relation to reperfusion by PPCI might also be important. Previous clinical studies have shown that remote ischaemic conditioning is effective at reducing myocardial infarct size when administered before PPCI (either in transit to the PPCI centre⁵ or on arrival at the hospital¹¹), during reperfusion by PPCI, and even at onset of reperfusion.⁷ In our study, clinical outcomes did not differ on the basis of whether the remote ischaemic conditioning protocol was completed during transportation to the PPCI centre or on admission to hospital, where the remote ischaemic conditioning protocol was continued during reperfusion in most patients (appendix 1 pp 40–41). Furthermore, no differences in clinical outcomes were observed whether the full four cycles of the remote ischaemic conditioning protocol was completed before reperfusion or not (appendix 1 pp 40–41).

Comedications and comorbidities might affect the cardioprotective efficacy of remote ischaemic conditioning. In experimental studies, $P2Y_{12}$ receptor antagonists confounded the cardioprotective effects of ischaemic conditioning,^{21,22} although no evidence for remote ischaemic conditioning is available. In our trial, the $P2Y_{12}$ receptor antagonist ticagrelor was administered to 69.8% of patients receiving remote ischaemic conditioning. We found no difference in clinical outcomes between the control group and the remote ischaemic conditioning group with respect to ticagrelor administration (appendix 1 p 41).

Experimental data have shown that age and presence of comorbidities, including diabetes, might attenuate the cardioprotective effects of ischaemic preconditioning and postconditioning.²³ Whether these factors affected the cardioprotective efficacy of remote ischaemic conditioning in our study remains unclear; our prespecified subgroup analyses showed no differences in clinical outcomes between the control group and the remote ischaemic conditioning group with increasing age or presence of diabetes (figure 3).

The efficacy of cardioprotective interventions applied at reperfusion in patients with STEMI is closely related to myocardial infarct size and pre-PPCI TIMI flow, with the greatest benefit reported for patients with complete occlusion (pre-PPCI TIMI flow \leq 1) due to infarcts in the left anterior descending artery.⁵ However, our prespecified subgroup analyses showed no differences in clinical outcomes according to pre-PPCI TIMI flow and infarct location (figure 3).

Remote ischaemic conditioning has shown mixed effects on outcomes in other clinical settings including cardiac surgery. Three large, prospective, multicentre, randomised controlled trials failed to find any beneficial effects of remote ischaemic conditioning on post-surgical clinical outcomes,²⁴⁻²⁶ possibly because of the confounding effects of propofol anaesthesia. Smaller studies of remote ischaemic conditioning in patients with stroke, renal transplantation, or undergoing elective PCI have suggested potential benefits.²⁷⁻²⁹ A clinical study in patients with STEMI treated by PPCI investigated the effects of remote ischaemic conditioning applied daily for 1 month (termed chronic remote ischaemic conditioning).³⁰ This intervention showed no beneficial effects on post-infarct left ventricular remodelling, although it was only started on day 3 post-PPCI in that study, which might be too late to observe any beneficial effects.

A potential limitation of our study is the short follow-up time of 12 months post-STEMI for the primary outcome of cardiac death and hospitalisation for heart failure, which might have been too short to observe any effect of remote ischaemic conditioning on clinical outcomes. In summary, the findings from our trial provide conclusive evidence that remote ischaemic conditioning offers no benefits on either myocardial infarct size or clinical outcomes at 12 months in patients with STEMI treated by PPCI. Unfortunately, remote ischaemic conditioning had been the most promising cardioprotective strategy for improving clinical outcomes following STEMI, and few other prospective options exist. As such, further studies are needed to identify novel cardioprotective targets and innovative approaches to cardioprotection such as combination multitarget therapy.³¹

Contributors

DJH, HEB, RKK, and DMY wrote the first draft of the manuscript. DJH, HEB, DMY, RKK, and TC designed the study. All authors participated in data collection. JMN, MD, HEB, RKK, DJH, DMY, UKM, MM, AP, and TC analysed the data. All authors vouch for the data and the analysis, contributed to writing the paper, and participated in the decision to publish the paper.

Declaration of interests

RKK, MRS, and HEB are shareholders in CellAegis. DAG reports institutional grants from Bristol-Myers Squibb/Pfizer and Bayer. TE reports personal fees from Bayer, Novo, Bristol-Myers Squibb, and Abbott outside the submitted work. LOJ has received unrestricted grants from Biotronik, Biosensors, and Boston Scientific to her institution outside the submitted work. All other authors declare no competing interests.

Data sharing

Requests for data collected for the study can be made to the co-corresponding authors and will be considered on an individual basis. Additional, related documents are immediately available (eg, study protocol, statistical analysis plan, informed consent form) and can be requested from the co-corresponding authors.

Acknowledgments

The ERIC-PPCI trial was funded by a British Heart Foundation clinical study grant (grant number CS/14/3/31002) and a University College London Hospitals/University College London Biomedical Research Centre clinical research grant. The CONDI-2 trial was funded by Danish Innovation Foundation grants (grant numbers 11-108354 and 11-115818), Novo Nordisk Foundation (grant number NNF13OC0007447), and TrygFonden (grant number 109624). DJH was supported by the British Heart Foundation (grant number FS/10/039/28270), the National Institute for Health Research (NIHR) Biomedical Research Centre at University College London Hospitals, the Duke-National University Singapore Medical School, the Singapore Ministry of Health's National Medical Research Council under its Clinician Scientist-Senior Investigator scheme (grant number NMRC/CSA-SI/0011/2017) and its Collaborative Centre Grant scheme (grant number NMRC/CGAug16C006), and the Singapore Ministry of Education Academic Research Fund Tier 2 (grant number MOE2016-T2-2-021). HEB was supported by the Novo Nordisk Foundation (grant numbers NNF14OC0013337, NNF15OC0016674). RKK is supported by the Oxford NIHR Biomedical Centre. The research was also supported by the NIHR infrastructure at Leeds. The views expressed are those of the author(s) and not necessarily those of the National Health Service, the NIHR, or the Department of Health. This article is based on the work of COST Action EU-CARDIOPROTECTION (CA16225) and supported by COST (European Cooperation in Science and Technology). We thank all study personnel for their invaluable assistance.

References

- Szummer K, Wallentin L, Lindhagen L, et al. Improved outcomes in patients with ST-elevation myocardial infarction during the last 20 years are related to implementation of evidence-based treatments: experiences from the SWEDEHEART registry 1995–2014. Eur Heart J 2017; 38: 3056–65.
- Heusch G, Bøtker HE, Przyklenk K, Redington A, Yellon D. Remote ischemic conditioning. J Am Coll Cardiol 2015; 65: 177–95.

- 3 Schmidt MR, Smerup M, Konstantinov IE, et al. Intermittent peripheral tissue ischemia during coronary ischemia reduces myocardial infarction through a KATP-dependent mechanism: first demonstration of remote ischemic perconditioning. *Am J Physiol Heart Circ Physiol* 2007; 292: H1883–90.
- 4 Kharbanda RK, Mortensen UM, White PA, et al. Transient limb ischemia induces remote ischemic preconditioning in vivo. *Circulation* 2002; **106**: 2881–83.
- 5 Bøtker HE, Kharbanda R, Schmidt MR, et al. Remote ischaemic conditioning before hospital admission, as a complement to angioplasty, and effect on myocardial salvage in patients with acute myocardial infarction: a randomised trial. *Lancet* 2010; 375: 727–34.
- 6 Rentoukas I, Giannopoulos G, Kaoukis A, et al. Cardioprotective role of remote ischemic periconditioning in primary percutaneous coronary intervention: enhancement by opioid action. *JACC Cardiovasc Interv* 2010; 3: 49–55.
- 7 Crimi G, Pica S, Raineri C, et al. Remote ischemic post-conditioning of the lower limb during primary percutaneous coronary intervention safely reduces enzymatic infarct size in anterior myocardial infarction: a randomized controlled trial. *JACC Cardiovasc Interv* 2013; 6: 1055–63.
- 8 Prunier F, Angoulvant D, Saint Etienne C, et al. The RIPOST-MI study, assessing remote ischemic perconditioning alone or in combination with local ischemic postconditioning in ST-segment elevation myocardial infarction. *Basic Res Cardiol* 2014; 109: 400.
- 9 Yellon DM, Ackbarkhan AK, Balgobin V, et al. Remote ischemic conditioning reduces myocardial infarct size in STEMI patients treated by thrombolysis. J Am Coll Cardiol 2015; 65: 2764–65.
- 10 Eitel I, Stiermaier T, Rommel KP, et al. Cardioprotection by combined intrahospital remote ischaemic perconditioning and postconditioning in ST-elevation myocardial infarction: the randomized LIPSIA CONDITIONING trial. *Eur Heart J* 2015; 36: 3049–57.
- 11 White SK, Frohlich GM, Sado DM, et al. Remote ischemic conditioning reduces myocardial infarct size and edema in patients with ST-segment elevation myocardial infarction. *JACC Cardiovasc Interv* 2015; 8: 178–88.
- 12 Verouhis D, Sörensson P, Gourine A, et al. Effect of remote ischemic conditioning on infarct size in patients with anterior ST-elevation myocardial infarction. Am Heart J 2016; 181: 66–73.
- 13 Sloth AD, Schmidt MR, Munk K, et al. Improved long-term clinical outcomes in patients with ST-elevation myocardial infarction undergoing remote ischaemic conditioning as an adjunct to primary percutaneous coronary intervention. *Eur Heart J* 2014; 35: 168–75.
- 14 Stiermaier T, Jensen JO, Rommel KP, et al. Combined intrahospital remote ischemic perconditioning and postconditioning improves clinical outcome in ST-elevation myocardial infarction. *Circ Res* 2019; 124: 1482–91.
- 15 Gaspar A, Lourenço AP, Pereira MA, et al. Randomized controlled trial of remote ischaemic conditioning in ST-elevation myocardial infarction as adjuvant to primary angioplasty (RIC-STEMI). *Basic Res Cardiol* 2018; 113: 14.
- 16 Hausenloy DJ, Kharbanda R, Rahbek Schmidt M, et al. Effect of remote ischaemic conditioning on clinical outcomes in patients presenting with an ST-segment elevation myocardial infarction undergoing primary percutaneous coronary intervention. *Eur Heart J* 2015; **36**: 1846–48.
- 17 Ibanez B, James S, Agewall S, et al. 2017 ESC Guidelines for the management of acute myocardial infarction in patients presenting with ST-segment elevation: The Task Force for the management of acute myocardial infarction in patients presenting with ST-segment elevation of the European Society of Cardiology (ESC). *Eur Heart J* 2018; **39**: 119–77.

- 18 Hicks KA, Tcheng JE, Bozkurt B, et al. 2014 ACC/AHA key data elements and definitions for cardiovascular endpoint events in clinical trials: a report of the American College of Cardiology/ American Heart Association task force on clinical data standards (writing committee to develop cardiovascular endpoints data standards). J Am Coll Cardiol 2015; 66: 403–69.
- 19 Traverse JH, Swingen CM, Henry TD, et al. NHLBI-sponsored randomized trial of postconditioning during primary percutaneous coronary intervention for ST-elevation myocardial infarction. *Circ Res* 2019; **124**: 769–78.
- 20 Johnsen J, Pryds K, Salman R, Løfgren B, Kristiansen SB, Bøtker HE. The remote ischemic preconditioning algorithm: effect of number of cycles, cycle duration and effector organ mass on efficacy of protection. *Basic Res Cardiol* 2016; 111: 10.
- 21 Cohen MV, Downey JM. The impact of irreproducibility and competing protection from P2Y₁₂ antagonists on the discovery of cardioprotective interventions. *Basic Res Cardiol* 2017; **112**: 64.
- 22 Yang XM, Liu Y, Cui L, et al. Platelet P2Y₁₂ blockers confer direct postconditioning-like protection in reperfused rabbit hearts. *J Cardiovasc Pharmacol Ther* 2013; 18: 251–62.
- 23 Ferdinandy P, Hausenloy DJ, Heusch G, Baxter GF, Schulz R. Interaction of risk factors, comorbidities, and comedications with ischemia/reperfusion injury and cardioprotection by preconditioning, postconditioning, and remote conditioning. *Pharmacol Rev* 2014; 66: 1142–74.
- 24 Hong DM, Lee EH, Kim HJ, et al. Does remote ischaemic preconditioning with postconditioning improve clinical outcomes of patients undergoing cardiac surgery? Remote Ischaemic Preconditioning with Postconditioning Outcome Trial. *Eur Heart J* 2014; 35: 176–83.
- 25 Meybohm P, Bein B, Brosteanu O, et al. A multicenter trial of remote ischemic preconditioning for heart surgery. N Engl J Med 2015; 373: 1397–407.
- 26 Hausenloy DJ, Candilio L, Evans R, et al. Remote ischemic preconditioning and outcomes of cardiac surgery. N Engl J Med 2015; 373: 1408–17.
- 27 Hougaard KD, Hjort N, Zeidler D, et al. Remote ischemic perconditioning as an adjunct therapy to thrombolysis in patients with acute ischemic stroke: a randomized trial. *Stroke* 2014; 45: 159–67.
- 28 Davies WR, Brown AJ, Watson W, et al. Remote ischemic preconditioning improves outcome at 6 years after elective percutaneous coronary intervention: the CRISP stent trial long-term follow-up. *Circ Cardiovasc Interv* 2013; 6: 246–51.
- 29 MacAllister R, Clayton T, Knight R, et al. REmote preconditioning for Protection Against Ischaemia–Reperfusion in renal transplantation (REPAIR): a multicentre, multinational, double-blind, factorial designed randomised controlled trial. *Efficacy Mech Eval* 2015; 2: eme02030.
- 30 Vanezis AP, Arnold JR, Rodrigo G, et al. Daily remote ischaemic conditioning following acute myocardial infarction: a randomised controlled trial. *Heart* 2018; **104**: 1955–62.
- 31 Davidson SM, Ferdinandy P, Andreadou I, et al. Multitarget strategies to reduce myocardial ischemia/reperfusion injury: JACC review topic of the week. J Am Coll Cardiol 2019; 73: 89–99.