REVIEW

Remote Ischemic Conditioning in Patients with Acute Coronary Syndromes: A Systematic Review with Meta-Analysis and Trial Sequential Analysis

This article was published in the following Dove Press journal: *Clinical Epidemiology*

Irene Sandven ^[b] Jan Eritsland² Michael Abdelnoor ^{[b]3,4}

¹Oslo Centre for Biostatistics and Epidemiology (OCBE), Oslo University Hospital, Oslo, Norway; ²Department of Cardiology, Oslo University Hospital, Oslo, Norway; ³Centre of Clinical Heart Research, Oslo University Hospital, Oslo, Norway; ⁴Epidemiology and Biological Statistics, Independent Health Research Unit, Oslo, Norway **Objective:** To evaluate the efficacy of remote ischemic conditioning (RIC) as compared to no conditioning on clinical endpoints in acute coronary syndromes (ACS) patients undergoing percutaneous coronary intervention (PCI).

Design: Systematic review of randomized clinical trials (RCTs).

Material and Methods: Literature was searched up to September 13, 2019, and we identified a total of 13 RCTs. The efficacy of RIC on incidence of clinical events during follow-up was quantified by the rate ratio (RR) with its 95% confidence interval (CI), and we used fixed and random effects models to synthetize the results. Small-study effect was evaluated, and controlled for by the trim-and-fill method. Heterogeneity between studies was examined by subgroup and meta-regression analyses. The risk of false-positive results in meta-analysis was evaluated by trial sequential analysis (TSA).

Results: Pooled analysis of 13 trials (7183 patients) showed that RIC compared to no conditioning revealed a non-significant risk reduction on endpoint mortality (RR=0.81, 95% CI: 0.56–1.17) during a median follow-up time of 1 year (range: 0.08–3.8) with low heterogeneity ($I^2=16\%$). Controlling for small-study effect showed no efficacy of RIC (adjusted RR: 1.03, 95% CI: 0.66–1.59). Pooled effect of RIC on the incidence of myocardial infarction (MI) from 11 trials (6996 patients) was non-significant too (RR=0.85, 95% CI: 0.62–1.18), with no observed heterogeneity ($I^2=0\%$) or small-study effect. A similar lack of efficacy was found in endpoint congestive heart failure (CHF) from 6 trials including 6098 patients (RR=0.71, 95% CI: 0.44–1.15), with moderate heterogeneity ($I^2=30\%$). TSAs showed that the pooled estimates from the cumulative meta-analyses were true negative with adequate power.

Conclusion: Evidence from this updated systematic review demonstrates no beneficial effect of RIC on the incidence of clinical endpoint mortality, MI and CHF during a median follow-up of 1 year in ACS patients undergoing PCI.

Keywords: remote ischemic conditioning, mortality, myocardial infarction, congestive heart failure, meta-analysis, trial sequential analysis

Introduction

Remote ischemic conditioning (RIC) is a non-invasive procedure providing temporal episodes of reversible ischemia through repeated inflations and deflations of a blood pressure limb cuff. RIC has been tried as a supplement to percutaneous intervention (PCI) in patients presenting with acute coronary syndromes (ACS) to protect against ischemia/reperfusion (I/R) injury, which refers to the damage caused by reperfusion of an organ exposed to a period of ischemia. In RIC the brief episodes of reversible

Correspondence: Irene Sandven Email uxsair@ous-hf.no



Clinical Epidemiology 2020:12 595-605

© 2020 Sandven et al. This work is published and licensed by Dove Medical Press Limited. The full terms of this license are available at https://www.dovepress.com/terms. work you hereby accept the Terms. Non-commercial uses of the work are permitted without any further permission for Dove Medical Press Limited, provided the work is properly attributed. For permission for commercial uses of the work are permitted without any further permission for Dove Medical Press Limited, provided the work is properly attributed. For permission for commercial uses of the work are peragraphs 4.2 and 5 of our Terms (https://www.dovepress.com/terms.php). ischemia with reperfusion in a vascular bed remote from the ischemic organ (eg the heart) have been shown to reduce I/R injury probably through neuronal or humoral mediators.¹ There are three temporal variants of RIC according to when the remote conditioning stimulus is applied; before (preconditioning), during (per-conditioning), or immediately after reperfusion (post-conditioning). In animal models, RIC has consistently been shown to reduce infarct size,² and subsequently a reduction of infarct size by RIC was reported in small-sized "proof-of-concept" clinical studies.³⁻⁶ Meta-analyses of clinical trials point towards an association between RIC and lower levels of cardiospecific biomarker release, and suggest a beneficial effect of RIC on major adverse cardiac events such as mortality, myocardial infarction (MI), and congestive heart failure (CHF) combined or separately.⁷⁻¹² A trend towards reduced clinical events at follow-up after RIC in ST-elevation myocardial infarction (STEMI) patients has been reported in two trials^{13,14} while no improvement by RIC was reported in one recent large multicenter trial¹⁵ providing additional clinical data.

Objective

Clinical trials assessing RIC-induced effect on clinical endpoints have been few. Therefore, we conducted an updated systematic review followed by meta-analysis and trial sequential analysis (TSA) to evaluate the efficacy of RIC as compared to no conditioning on clinical endpoints in patients with ACS undergoing PCI, and to help clarify the need for additional trials.

Materials and Methods

The review protocol has been registered at <u>https://www.</u> crd.york.ac.uk/PROSPERO/, ID: CRD42020147789.

Search

With guidance from a qualified medical librarian, we searched Ovid Medline and Embase until September 13, 2019 with no date or language restrictions. The population was limited to adult humans above 18 years, and the design to randomized clinical trials (RCTs). In Ovid Medline, the search was conducted using Medical Subject Headings (MeSH) and text words including, but not restricted to: remote and (ischemic or myocardial infarction) and (conditioning or preconditioning) or postconditioning) and (randomized controlled trial or controlled clinical trial). An additional search in Cochrane Database of Systematic Reviews was performed, and we searched for ongoing systematic reviews using Prospero. Unpublished clinical trials were searched for by consulting the Clinicaltrials.gov website. Reference lists of published meta-analyses were screened for any relevant studies not included in the original search.

Study Selection

Due to a therapeutic research question we included only RCTs comparing RIC with no conditioning in adult patients with ACS according to the specified research question presented in Table 1. Two investigators (IS and MA) independently evaluated the publications to assess whether they met the predefined inclusion criteria. We resolved any disagreement through discussion with a third author (JE).

Quality Assessment of the Trials

Two reviewers (IS, MA) independently assessed the quality of included studies following the Cochrane Collaboration's tool assessing risk of bias in RCTs;¹⁶ randomization and concealment of treatments allocation (selection bias), blinding of investigator to outcome (detection bias), dropout (attrition bias), adequacy of analysis according to intention to treat (other bias), and a beforehand power analysis for clinical endpoints.

Data Abstraction

Data regarding publication status (first author, publication year, and country where the study took place), patient-related characteristics of the total cohort (mean age, frequency of male

Table I Specification of the Research Question Applying a PICO (Population, Intervention, Comparison and Outcome) Model

Population	Intervention	Comparison	Outcomes
Adults (≥18 years) presenting with acute coronary syndromes (STEMI, NSTEMI or S/UA), treated with percutaneous coronary intervention	Remote ischemic conditioning; cycles of blood pressure cuff inflation and deflation applied before PCI (pre-conditioning), during PCI (per- conditioning) or after PCI (post-conditioning)	PCI with no conditioning	Mortality, myocardial infarction, congestive heart failure

Notes: Subgroups were stratified by population (STEMI versus NSTEMI or S/UA) and intervention (pre-, per-, post-conditioning).

Abbreviations: STEMI, ST-segment elevation myocardial infarction; NSTEMI, no ST-segment elevation myocardial infarction; S/UA, stable/unstable angina pectoris; PCI, percutaneous coronary intervention.

gender, hypertension, diabetes mellitus, and smokers), outcomes (total mortality, MI, and CHF), results (number of events, total number of patients in intervention and comparison group, and follow-up time), and study quality (risk of bias assessment) were extracted in duplicate on a standardized form according to the á priori protocol. Disagreements were resolved by discussion among the review authors and subsequent consensus.

Summary Measures

The pooled effect of intervention on each clinical endpoint was quantified by the rate ratio (RR) with its 95% confidence interval (CI), considering the person-year model to control for variability in the duration of follow-up. Both fixed and random effects model was considered, but in the presence of heterogeneity between trials the random effects model was preferred according to the DerSimonian and Laird method.¹⁷ Using this model the estimate of the pooled effect measure and its CI incorporate the additional variability due to inter-study variance (τ^2).

Sources of Heterogeneity, Evaluation and Quantification

Statistical heterogeneity among studies was assessed with Cochran's Q test. The magnitude of heterogeneity was evaluated by the I² statistics which describes the proportion of total variation due to heterogeneity rather than chance.¹⁸ When heterogeneity was present we used subgroup analyses and meta-regression to investigate possible sources of heterogeneity. We stratified our data according to type of ACS (STEMI versus non-STEMI or Stable/Unstable Angina), timing of RIC (pre-, per- or post-conditioning), and the following study characteristics; concealment of randomisation, blinding of the investigator on the outcome, analysis according to intention to treat strategy, and presence of a priori power analysis for the study. To further explore potential sources of heterogeneity we considered a random effect meta-regression analysis, where the outcome variable is the observed RR from every trial indicating treatment effect and the different studylevel and patient-level characteristics (covariates). Sources of heterogeneity were considered as important if the covariate decreased the between- study variance. The estimate of τ^2 in the presence of a covariate in comparison to that when the covariate is omitted, allows the proportion of the heterogeneity variance explained by the covariate to be calculated.¹⁹ Because we had less than ten trials for endpoint CHF, metaregression was not performed.²⁰

Evaluation of Small-Study Effect

In order to assess potential small-study effect we used the funnel plot, which is a good visual evaluation of sampling bias. To further assess potential bias we used two well-established tests of small-study effect; Begg and Mazumdar rank correlation,²¹ and Egger's test of asymmetry.²² For power considerations a test of funnel plot asymmetry was not applied for endpoint CHF.²³

Correction for Small-Study Effect

In the presence of small-study effect, we considered the trim-and-fill method²⁴ to adjust for small-study effect. This method is a kind of sensitivity analysis to assess the potential impact of missing studies.

Trial Sequential Analysis (TSA)

Evaluation of the meta-analyses by TSA was performed to eliminate false-positive results, and to help clarify the need for additional trials. This method was developed by Pogue and Yusuf^{25,26} and advocated the use of Lan-DeMets trial sequence monitoring boundaries (TSMB) for cumulative meta-analysis. More recently the method was used accounting for bias and observed heterogeneity in a retrospective cumulative meta-analysis.²⁷ We used TSA as it is implemented in Stata and with the ldbounds package in R-statistical software, and considered three types of information sizes calculated by the program: 1) Accrued Information Size (AIS), which is the total number of patients (N) in the metaanalysis. Type-I error, Type-II error and relative risk reduction (RRR) are entered by user, and we estimated power for the given RRR and sample size. 2) Low Bias Information Size (LBIS), where RRR is calculated from only those trials with low bias. Required sample is calculated for this RRR, Type-I error and Type-II error. 3) Low Bias Heterogeneity adjusted Information Size (LBHIS), which is calculated similar to LBIS and then adjusted for heterogeneity. We determined LBHIS when heterogeneity was over 30%. We set Type-I error at 5%, Type-II error at 20%, and for AIS the RRR at 10%. The fixed model was applied when there was low heterogeneity ($I^2 < 30\%$) and the random model when I^2 \geq 30%. We estimated the TSMB for AIS, LBIS and LBHIS to detect potentially spurious level.²⁸

Software for Meta-Analysis

Statistical analyses were performed with STATA 15.0,²⁹ and R–Package–Meta.³⁰ TSA was performed combining Stata commands with R packages.²⁸

We followed the preferred reporting items for systematic reviews and meta-analyses (PRISMA) guidelines for research reporting.³¹

Results

Trial Selection

After identifying 1220 references, 1184 were excluded due to irrelevant content and duplicate publications leaving 36 potentially eligible. Records were excluded mainly because they did not contain the right population (ACS patients), intervention (RIC versus no conditioning) or clinical endpoints. Finally, 12 studies^{3–6,13–15,32–36} met our inclusion criteria, whereof one with two intervention arms,³⁴ thus 13 trials were considered in the meta-analysis (Figure 1). No additional publications were identified in the forward citation searching or hand searching of reference lists.

Trial Characteristics

The 13 trials included a total of 7183 ACS patients undergoing PCI (Table 2); the subtypes of ACS were STEMI in 9 trials and NSTEMI or S/UA in 4 trials. Median followup was 1 year (range: 0.08–3.8). The patients median age was 62.4 years (range: 51.6–67), 75.7% were men (range: 35.4–94.6), 52.1% with hypertension (range: 22.6–78.5), 27.9% with diabetes mellitus (range: 8.6–49.2), and 53.3% were smokers (range: 32.4–70). The trials were on populations from 3 geographical regions; Europe (6 trials), Asia (4 trials), and North America (3 trials).

RIC interventions were quite similar among all trials with 3–5 cycles of 5-minutes inflation and 5-minutes deflation of a blood pressure cuff to 200 mmHg sited on an upper arm or lower limb/thigh. RIC was performed as pre-conditioning in 6 trials,^{4–6,13–15} as per-conditioning in 4 trials,^{3,34,35} and as post-conditioning in 3 trials.^{32,33,36}

Clinical outcomes were considered as primary endpoints in 2 trials only,^{13,15} and reported as secondary/ additional endpoints in the rest of them. All trials reported on mortality, 11 on MI,^{3,5,6,13-15,32–34,36} and 6 on CHF.^{4,5,13-15,32}

Quality Assessment

Randomization and adequate concealment were considered adequate in 7 trials, blinding of the endpoint assessor to treatment allocation in all trials for endpoint mortality while for other endpoints misclassification might exist. No dropout was reported during follow-up in 9 trials, and intention-to-treat strategy was followed in 8 trials (Table 3). In summary, 5 of the included trials represented overall low risk of bias.^{3,13-15,36} Power estimation on clinical endpoint was present in two trials only.^{13,15}

Synthesis of Results from Individual Trials

Mortality was reported in 13 trials (7183 patients) and the pooled effect of RIC compared to no conditioning was RR=0.81 (95% CI: 0.56-1.17) with low heterogeneity $(I^2=16\%)$, indicating no statistically significant effect (Figure 2A). The funnel plot visually showed the possibility of small-study effect (Figure 3A) and asymmetry was indicated by the Egger's test (p=0.032), but not by the Begg's test (p=0.7603). The trim-and-fill simulation method suggested 5 studies as missing, and the adjusted point estimate was altered towards the null-effect (adjusted RR=1.03, 95% CI: 0.66–1.59). In stratification analysis the only significant subgroup difference found was between trials adequately powered for clinical outcomes and those not (p=0.0457), and in meta-regression this study-related characteristic was the only covariate significantly associated with intervention effect (p=0.0096).

TSA supported lack of evidence for an effect of RIC compared with no conditioning for endpoint mortality. Cumulative fixed-effect meta-analysis with Lan-DeMets bounds showed a true negative result for AIS of 7183 patients. The power of the meta-analysis for a 10% RRR and the conventional Type-I error of 5% was 99%. The LBIS required is lower (803patients) for a RRR of 18% and a power of 80% without changing the conclusion; RIC does not reduce the incidence of mortality in PCI treated ACS patients. Our meta-analysis is satisfactory powered for outcome mortality (Table 4A).

Myocardial Infarction (MI) during follow-up was reported in 11 trials (6996 patients) and the pooled effect of RIC compared to no conditioning was statistically non-significant (RR=0.85, 95% CI: 0.62–1.18) with no heterogeneity (I^2 =0%) (Figure 2B). The funnel plot visually indicated no small-study effect (Figure 3B), confirmed by the Egger's test (p=0.740) and the Begg's test (p=0.6971). Stratification analysis and metaregression were not indicated since there was no observed heterogeneity between the trials.

TSA supported lack of evidence for an effect of RIC compared with no conditioning on endpoint MI. Cumulative fixed-effect meta-analysis with Lan-DeMets bounds showed a true negative result for AIS of 6996 patients. The power of the meta-analysis for a 10% RRR and Type-I error of 5% was 99%. The LBIS required is

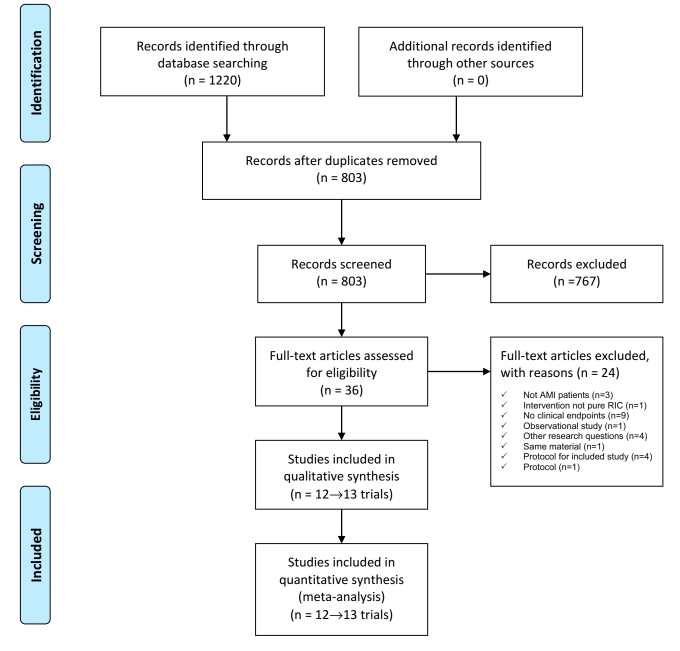


Figure I PRISMA flow diagram of study selection from the literature searches for the systematic review of randomized clinical trials (RCTs) investigating the efficacy of remote ischemic conditioning (RIC) in acute coronary syndromes (ACS) patients undergoing percutaneous coronary intervention (PCI).

Note: Copied from Moher D, Liberati A, Tetzlaff J, Altman DG, PRISMA Group. Preferred reporting items for systematic reviews and meta-analyses: the PRISMA statement. Ann Intern Med. 2009;151(4):264–269. doi:10.7326/0003-4819-151-4-200908180-00135.³¹

1407 without changing the conclusion for a RRR of 14% and a power of 80%; RIC does not reduce the incidence of MI in PCI treated ACS patients. Our meta-analysis is satisfactory powered for endpoint MI (Table 4A).

Congestive Heart Failure (CHF) was reported in 6 trials (6098 patients) and the pooled effect of RIC compared to no conditioning was statistically non-significant (RR=0.71, 95% CI: 0.44–1.15) with moderate heterogeneity ($I^2=30\%$) (Figure 2C). The funnel plot visually indicated small-study effect

(Figure 3C), but asymmetry was not investigated further because of too few trials. Stratification analysis and meta-regression were not performed due to less than 10 trials. CHF was defined in 3 trials.^{13–15} The incidence of CHF in the largest trial with 5115 patients¹⁵ and the other 5 trials^{4,5,13,14,32} was close, and included in each others' 95% CIs.

TSA supported lack of evidence for an effect of RIC compared with no conditioning on endpoint CHF. Cumulative fixed-effect meta-analysis with Lan-DeMets

First Author	Year	Country	RIC/ controls (n/n)	Age ^a (mean)	Men (%)	Hypertension (%)	Diabetes (%)	Smokers (%)	Followup ^a (mean)	Patient Years ^b
Carrasco-Chinchilla ³⁶	2013	ES	118/114	64.6	68.I	75.6	42.1	66.0	1.0	232.00
Crimi ³	2013	ΙТ	48/48	58.4	87.5	52.1	11.5	47.9	1.0	96.00
Sloth ¹⁴	2014	DK	126/125	62.5	75.7	31.1	9	56.2	3.8	953.80
Yamanaka ⁴	2015	JP	47/47	67.0	74.5	63.8	33.0	53.3	0.08	7.52
Liu ⁵	2016	CN	59/60	62.4	79.0	42.9	20.2	42.9	1.0	119.00
Lavi ^{c34}	2016	Canada	119/118	63.8	72.6	70.0	32.9	-	1.7	402.90
Lavi ^{c34}	2016	Canada	119/118	64.4	73.8	69.6	30.0	-	1.7	402.90
Verouhis ³⁵	2016	SE	47/46	61.0	94.6	22.6	8.6	37.6	0.08	7.44
Zhou ⁶	2017	CN	72/58	59.7	35.4	78.5	49.2	56.9	0.5	65.00
Elbadawi ³³	2017	USA	30/30	51.6	83.3	33.3	41.7	70.0	0.5	30.00
Qian ³²	2018	CN	37/34	56.4	53.5	54.9	11.3	32.4	0.5	35.50
Gaspar ¹³	2018	РТ	231/217	60.0	80.1	49.1	27.9	58.9	2.1	940.80
Hausenloy ¹⁵	2019	UK, DK,	2546/2569	63.5	76.8	41.4	11.1	39.3	1.0	5115.00
		ES, RS								

Table 2 Characteristics of the 13 Trials Comparing Remote Ischemic Conditioning (RIC) versus No Conditioning (Controls) in Acute

 Coronary Syndromes (ACS) Patients Undergoing Percutaneous Coronary Intervention (PCI)

Notes: ^ayears, ^bpatients-years = (total number of patients × mean follow-up time in years), ^cstudy with two intervention arms (RIC with arm cuff and thigh cuff). Abbreviations: ES, Spain; IT, Italy; PT, Portugal; UK, United Kingdom; DK, Denmark; RS, Serbia; CN, China; SE, Sweden; JP, Japan.

Table 3 Review Authors' Quality Assessment of the 13 Trials Comparing Remote Ischemic Conditioning (RIC)	versus No
Conditioning in Acute Coronary Syndromes (ACS) Patients Undergoing Percutaneous Coronary Intervention (PCI)	

First Author	Selection Bia	Detection	Bias		Attrition Bias	Other bias	Overall Low Risk	Powered for Clinical	
	Random Sequence	Allocation Concealment	Blinding of Assessmer		ome	Incomplete Outcome	Intention to Treat	of Bias	Outcomes
	Generation		Mortality	мі	CHF	Data	Analysis		
Carrasco-Chinchilla ³⁶	Yes	Yes	Yes	Yes	-	No	Yes	Yes	No
Crimi ³	Yes	Yes	Yes	Yes	-	No	Yes	Yes	No
Sloth ¹⁴	Yes	Yes	Yes	Yes	Yes	No	Yes	Yes	No
Yamanaka ⁴	Yes	Unclear	Yes	-	Unclear	No	Yes	Nno	No
Liu ⁵	Yes	Yes	Yes	Yes	Yes	No	Unclear	No	No
Lavi ^{a34}	Yes	Unclear	Yes	Yes	-	Yes	Unclear	No	No
Lavi ^{a34}	Yes	Unclear	Yes	Yes	-	Yes	Unclear	No	No
Verouhis ³⁵	Yes	Yes	Yes	-	-	No	Unclear	No	No
Zhou ⁶	Unclear	Unclear	Yes	Yes	-	Yes	Unclear	No	No
Elbadawi ³³	Unclear	Unclear	Yes	Yes	-	Yes	Yes	No	No
Qian ³²	Unclear	Unclear	Yes	Yes	Unclear	No	Yes	No	No
Gaspar ¹³	Yes	Yes	Yes	Yes	Yes	No	Yes	Yes	Yes
Hausenloy ¹⁵	Yes	Yes	Yes	Yes	Yes	No	Yes	Yes	Yes

Notes: ^aStudy with two intervention arms (RIC with arm cuff and thigh cuff). **Abbreviations:** MI, myocardial infarction; CHF, congestive heart failure.

bounds showed a true negative result for AIS of 6098 patients. The power of the meta-analysis for a 10% RRR and Type-I error of 5% was 98%. The LBHIS required is 487 patients without changing the conclusion for a RRR of 26%, Type-I error of 5% and power of 80%; RIC does not reduce the incidence of CHF in PCI treated ACS patients.

Our meta-analysis is satisfactory powered for endpoint CHF (Table 4A).

When stratifying on the placement of the blood pressure cuff, upper arm or lower limb/thigh, there was no difference in pooled effect as far the 3 clinical endpoints considered.

Α

	Evno	rimental		Control				Weight	Woight
Study	Events		Events	Control Total	Rate Ratio	RR	95%-CI	-	Weight (random)
Carrasco-Chinchilla	2	118.00	0	114.00		- 4.83	[0.23; 99.54]	0.5%	1.4%
Crimi	0	48.00	1	48.00		0.33	[0.01; 7.98]	0.5%	1.3%
Sloth	5	478.80	15	475.00		0.33	[0.12; 0.90]	4.8%	10.6%
Yamanaka	0	3.76	3	3.76		0.14	[0.01; 2.00]	0.7%	1.8%
Liu	1	59.00	4	60.00		0.25	[0.03; 2.21]	1.0%	2.7%
Lavi	4	202.30	4	200.60		0.99	[0.25; 3.91]	2.6%	6.2%
Verouhis	1	3.76	2	3.68	+- <u></u>	0.49	[0.07; 3.35]	1.3%	3.4%
Lavi	3	202.30	4	200.60		0.74	[0.17; 3.28]	2.2%	5.4%
Zhou	0	36.00	0	29.00		0.81	[0.02; 39.52]	0.3%	0.9%
Elbadawi	0	15.00	0	15.00		1.00	[0.02; 47.28]	0.3%	0.9%
Qian	0	18.50	1	17.00		0.31	[0.01; 7.05]	0.5%	1.3%
Gaspar	17	485.10	21	455.10		0.76	[0.41; 1.42]	12.4%	20.9%
Hausenloy	122	2546.00	100	2569.00	+	1.23	[0.95; 1.59]	72.8%	43.1%
Fixed effect model Random effects model		4216.52		4190.74			[0.82; 1.27] [0.56; 1.17]	100.0%	 100.0%
Heterogeneity: $I^2 = 16\%$, τ	_	n = 0.29				0.01	[0.00, 1.11]		
1070, 1	0.0027	, p 0.20			0.1 0.51 2 10				

В

Study	Experi Events	imental Total	C Events	Control Total	Rate Ratio	RR	95%-CI	Weight (fixed)	Weight (random)
Carrasco-Chinchilla	4	118.0	1	114.0			.44; 34.05]	2.3%	2.3%
Crimi	0	48.0	1	48.0 -		0.33 [(0.01; 7.98]	1.1%	1.1%
Sloth	8	478.8	11	475.0		0.72 [(0.29; 1.78]	13.1%	13.1%
Liu	1	59.0	1	60.0		1.02 [0	.07; 15.88]	1.4%	1.4%
Lavi	2	202.3	1	200.6		1.98 [0	.18; 21.70]	1.9%	1.9%
Lavi	1	202.3	1	200.6	<u> </u>	0.99 [0	.06; 15.74]	1.4%	1.4%
Elbadawi	3	15.0	2	15.0		1.50 [0	0.29; 7.73]	4.0%	4.0%
Zhou	4	36.0	6	29.0		0.54 (0.17; 1.72]	7.8%	7.8%
Gaspar	5	485.1	7	455.1		0.67 [0	0.21; 2.10]	8.2%	8.2%
Qian	1	18.5	2	17.0		0.46 (0.05; 4.62]	2.0%	2.0%
Hausenloy	38	2546.0	43	2569.0		0.89 [0	0.58; 1.37]	56.9%	56.9%
Fixed effect model		4209.0		4183.3	4	0.85 [0	.62; 1.18]	100.0%	
Random effects mode Heterogeneity: $I^2 = 0\%$, τ^2		93					.62; 1.18]		100.0%
······	-, p				0.1 0.51 2 10				

С

Study	Experimer Events To	ntal otal Events	Control Total	Rate Ratio	RR	95%-CI	Weight (fixed)	Weight (random)
Sloth Yamanaka Liu Qian Gaspar Hausenloy	1 59	.76 3 .00 2 .50 2 .10 17	00.00		0.57 0.33 0.51 0.46 0.44 1.06	[0.06; 1.93] [0.05; 5.46] [0.05; 4.62] [0.19; 1.01]	2.3% 1.1% 0.6% 0.6% 5.0% 90.3%	12.3% 6.7% 3.9% 4.1% 21.2% 51.9%
Fixed effect model Random effects model Heterogeneity: $J^2 = 30\%$, τ			3579.86	0.1 0.5 1 2		[0.82; 1.18] [0.44; 1.15]	100.0% 	 100.0%

Figure 2 Forest plot for meta-analysis of randomized clinical trials (RCTs) comparing remote ischemic conditioning (RIC) with no conditioning in acute coronary syndromes (ACS) patients for endpoint; (A) mortality (B) myocardial infarction (C) congestive heart failure.

Notes: Total; amount of person-years, Lavi; study with two intervention arms (RIC with arm cuff and thigh cuff)

Abbreviations: RR, rate ratio; Cl, confidence interval.

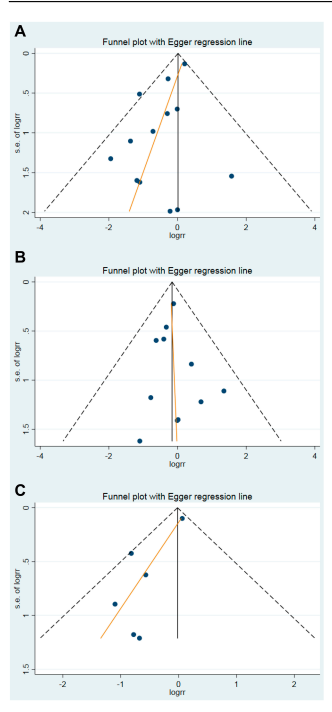


Figure 3 Funnel plot of the effect of remote ischemic conditioning (RIC) for endpoint (A) mortality (B) myocardial infarction (C) congestive heart failure. Abbreviations: logrr, log rate ratio; s.e, standard error.

Subgroup of STEMI Trials

Our results were robust in the subgroup of STEMI trials. For endpoint mortality from 9 trials (6347 STEMI patients) the pooled effect estimate was larger but still non-significant (RR=0.66, 95% CI: 0.39–1.11), and a true negative result indicated with sufficient

 Table 4 Trial Sequential Analysis (TSA) for Power Estimation

A. All Studies					
Accrued Information S	Size (AIS))			
Endpoint	RRR (%)	Type-l error (%)	Power (%)	AIS (N)	
Mortality	10	5	99	7183	
Myocardial infarction	10	5	99	6996	
Congestive heart failure	10	5	98	6098	
Low Bias Information	Size (LBI	S)			
Endpoint	RRR	Type-I error	Power	LBIS	
	(%)	(%)	(%)	(N)	
Mortality	18	5	80	803	
Myocardial infarction	14	5	80	1407	
, Congestive heart failure	26	5	80	340	
Low Bias Heterogenei	ty-Adjust	ed Information S	Size (LBHI	5)	
Endpoint	RRR	Type-I error	Power	LBHIS	
	(%)	(%)	(%)	(N)	
Congestive heart failure	26	5	80	487	
B. Subgroup of Trials	vith STE	MI Patients			
Accrued Information S	Size (AIS)				
Endpoint	RRR	Type-I error	Power	AIS (N)	
	(%)	(%)	(%)		
Mortality	10	5	99	6347	
Myocardial infarction	10	5	99	6160	
Congestive heart failure	10	5	98	6098	
Low Bias Information	Size (LBI	S)			
Endpoint	RRR	Type-I error	Power	LBIS	
	(%)	(%)	(%)	(N)	
Mortality	24	5	80	437	
Myocardial infarction	17	5	80	854	
Congestive heart failure	26	5	80	340	
Low Bias Heterogenei	ty-Adjust	ed Information S	Size (LBHI	5)	
Endpoint	RRR	Type-I error	Power	LBHIS	
•	(%)	(%)	(%)	(N)	
	(/0)				
Mortality	24	5	80	716	

Abbreviations: RRR, relative risk reduction; N, patients.

power (Table 4B). For endpoint MI from 7 trials (6160 STEMI patients) the pooled effect estimate was unchanged (RR=0.84, 95% CI: 0.59–1.19) and a true negative result indicated with sufficient power (Table 4B). For endpoint CHF there was no change as the patient population was STEMI in all 6 trials.

Sandven et al

Discussion

Summary of Evidence

Evidence from this updated systematic review suggests no protective effect of RIC on the incidence of clinical adverse events during follow-up in patients with ACS undergoing PCI. TSAs performed on meta-analyses conducted demonstrate true negative intervention effects on the endpoints mortality, MI and CHF with adequate power.

Strengths and Limitations

We performed a comprehensive literature search to prevent missing relevant trials.³⁷ Trial selection and data extraction were done by two authors to minimize transcription errors, and the components used for quality assessment are validated and reported to be associated with bias.³⁸ We have respected the important principles for meta-analysis methodology regarding eligibility criteria for the individual trials and analysis methods, pinpointing small-study effect and correcting for it when necessary.^{19,20} In addition to standard methods in meta-analysis, TSA was performed to avoid false-positive conclusions.^{27,28} Few trials provided data on development of CHF during follow-up, they used different definitions, and the probability of misclassification of this endpoint cannot be excluded. In the large trial of Hausenloy et al¹⁵ a blinded independent validation committee reviewed all events according to standard operation procedures. In this case the nature of the misclassification is non-differential which dilutes the efficacy of the intervention,³⁹ and the true effect of RIC on CHF might be stronger than the one estimated by this study. The major limitation of our study was the small-study effect and underpowered trials to evaluate clinical endpoints.

The Problem of Small-Study Effect for Endpoint Mortality

The majority of trials included in our meta-analysis were small and power deficient for investigating clinical outcomes, except Hausenloy et al¹⁵ and Gaspar et al.¹³ In this situation we found no treatment effect when stratifying on trials adequately powered versus power deficient. The impact of underpowered trials on meta-analysis results has been investigated,⁴⁰ indicating better intervention effects in underpowered than in adequately powered trials. Another reason might be that smaller trials are conducted and analyzed with less methodological rigor than larger trials, and overestimation of the intervention effects has been highlighted in lower quality trials as compared to trials of higher quality.⁴¹ Controlling for small-study effect by the trim-and-fill method confirmed the non-efficacy of RIC for endpoint mortality.

Evaluation of the Risk of False-Positive Result in Our Present Meta-Analyses

Often meta-analyses give a false-positive result, especially when updated with the publication of a new trial.^{27,28} When we evaluated the power of our meta-analyses for the 3 endpoints considered, the TSAs indicated presence of excellent statistical power for a 10% risk reduction and a 5% Type-I error as we included in the cumulative meta-analyses 7183, 6996, and 6098 patients, respectively. The newest trial¹⁵ assessed to be of high quality contributed with 5115 patients in the meta-analyses, and decisive for the conclusion of no effect of RIC on endpoint mortality, MI and CHF.

The Potential of False-Positive Results in Previous Meta-Analyses

Previous meta-analyses have reported cumulative pooled beneficial treatment effects of RIC on clinical endpoint mortality,^{8,10,12} MI,^{8,11,12} CHF,^{8,11} and different composite endpoints.^{7–9,11,12} Unfortunately none of them considered a power analysis of the cumulative effect in the meta-analyses, and their estimates might be false positive presumably due to small-study effect.

Discrepancies in Efficacy of RIC in ACS Between Surrogate Endpoints (SEPs) and Clinical Outcomes

Eleven of the 13 trials included in our study had SEPs as primary outcomes. Myocardial damage was commonly measured by biomarkers of infarct size (eg troponin and creatine kinase) and other outcomes such as resolutions of STsegment elevation and change in left ventricular ejection fraction (LVEF), and their results pointed to possible effect of RIC. Unfortunately, experience in clinical research underlines the failure of SEPs to be a valid measure of clinically important outcomes in different fields of clinical research.⁴² SEPs can be used in Phase 2 trials to determine whether an intervention is biologically active and whether to perform large trials with clinically important outcomes. RCTs with SEPs are comparable to experiments in animal model and must be used with great caution if at all in making clinical decisions. In our situation there was concordance between animal models and efficacy on SEPs as both of them considered measures of infarct size as an endpoint. However, trials adequately powered for meaningful clinical outcomes as incidence of mortality, MI and CHF are essential for evidence-based decisions in everyday clinical practice. The need for RCTs investigating clinical endpoints was highlighted in a recent meta-analysis reporting marginal efficacy of RIC during PCI in STEMI patients on SEPs such as infarct size and change in LVEF.⁴³

Implication for Research

The findings from the recent large high-quality RCT¹⁵ and the present systematic review suggest that further trials are not necessary to be conclusive regarding the efficacy of RIC on major clinical events as mortality, MI and CHF.

Implication for Practice

According to criteria for grading the quality of evidence and the strengths of recommendations^{44,45} this systematic review of RCTs provides high-level evidence of a nonpreventive effect of RIC against mortality, MI and CHF in ACS patients undergoing PCI. There seem to be no benefits on clinical endpoints during a median follow-up of 1 year (range: 0.08–3.8), and the use of RIC as a supplement to PCI can presently not be recommended.

Conclusion

This systematic review with meta-analysis and TSA suggests that there is no beneficial effect of RIC as a supplement to PCI in the treatment of ACS patients with respect to the incidence of mortality, MI and CHF during a median follow-up time of 1 year.

Author Contributions

All review authors contributed to conception and design, acquisition of data, or analysis and interpretation of data, drafting or revising the article critically for important intellectual content, gave final approval of the version to be published, and agree to be accountable for all aspects of the work.

Disclosure

The authors report no conflicts of interest in this work.

References

- Heusch G, Bøtker HE, Przyklenk K, Redington A, Yellon D. Remote Ischemic Conditioning. J Am Coll Cardiol. 2015;65(2):177–195. doi:10.1016/j.jacc.2014.10.031
- Bromage DI, Pickard JMJ, Rossello X, et al. Remote ischemic conditioning reduces infarct size in animal in vivo models of ischaemia-reperfusion injury: a systematic review and meta-analysis. *Cardiovasc Res.* 2017;113(3):288–297. doi:10.1093/cvr/cvw219

- 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 (10):1055–1063. doi:10.1016/j.jcin.2013.05.011
- Yamanaka T, Kawai Y, Miyoshi T, et al. Remote ischemic preconditioning reduces contrast-induced acute kidney injury in patients with ST-elevation myocardial infarction: a randomized controlled trial. *Int J Cardiol.* 2015;178:136–141. doi:10.1016/j.ijcard.2014.10.135
- Liu Z, Zhao L, Hong D, Gao J. Remote ischaemic preconditioning reduces myocardial ischaemic reperfusion injury in patients with ST-elevation myocardial infarction undergoing primary percutaneous coronary intervention. *Acta Cardiol.* 2016;71(5):596–603. doi:10.10 80/AC.71.5.3167504
- Zhou FZ, Song W, Yin LH, et al. Effects of remote ischemic preconditioning on myocardial injury and endothelial function and prognosis after percutaneous coronary intervention in patients with acute coronary syndrome. *Eur Rev Med Pharmacol Sci.* 2017;21 (20):4642–4648.
- Gong R, Wu YQ. Remote ischemic conditioning during primary percutaneous coronary intervention in patients with ST-segment elevation myocardial infarction: a systematic review and meta-analysis. *J Cardiothorac Surg.* 2019;14(1):14. doi:10.1186/ s13019-019-0834-x
- Liu H, Fu L, Sun X, Peng W, Chen Z, Li Y. Remote ischemic conditioning improves myocardial parameters and clinical outcomes during primary percutaneous coronary intervention: a meta-analysis of randomized controlled trials. *Oncotarget*. 2017;9(9):8653–8664. doi:10.18632/oncotarget.23818
- Elbadawi A, Ha LD, Abuzaid AS, Crimi G, Azzouz MS. Metaanalysis of randomized trials on remote ischemic conditioning during primary percutaneous coronary intervention in patients with ST-segment elevation myocardial infarction. *Am J Cardiol.* 2017;119(6):832–838. doi:10.1016/j.amjcard.2016.11.036
- Man C, Gong D, Zhou Y, Fan Y. Meta-analysis of remote ischemic conditioning in patients with acute myocardial infarction. *Sci Rep.* 2017;8;7:43529. doi:10.1038/srep43529
- McLeod SL, Iansavichene A, Cheskes S. Remote ischemic preconditioning to reduce reperfusion injury during acute ST-Segment-Elevation Myocardial Infarction: A systematic review and meta-analysis. J Am Heart Assoc. 2017;6(5):pii: e005522. doi:10.1161/JAHA.117.005522
- Le Page S, Bejan-Angoulvant T, Angoulvant D, Prunier F. Remote ischemic conditioning and cardioprotection: a systematic review and meta-analysis of randomized clinical trials. *Basic Res Cardiol.* 2015;110(2):11. doi:10.1007/s00395-015-0467-8
- 13. 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(3):14. doi:10.1007/s00395-018-0672-3
- 14. 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(3):168–175. doi:10.1093/eurheartj/eht369
- 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 randomized controlled trial. *Lancet.* 2019;395(10207):1415–1424. doi:10.1016/S0140-6736(19)32039-2
- Higgins JP, Altman DG, Gøtzsche PC, et al. The Cochrane Collaboration's tool for assessing risk of bias in randomized trials. *BMJ*. 2011;343(oct18 2):d5928. doi:10.1136/bmj.d5928
- DerSimonian R, Laird N. Meta-analysis in clinical trials. Control Clin Trials. 1986;Sep;7(3):177–188. doi:10.1016/0197-2456(86)900 46-2

- Higgins JP, Thompson SG, Deeks JJ, Altman DG. Measuring inconsistency in meta-analysis. *BMJ*. 2003;327(7414):557–560. doi:10.1136/ bmj.327.7414.557
- Egger M, Smith GD, Altman DG, editors. *Systematic Reviews in Health Care: Meta-Analysis in Context*. London: BMJ Publishing Group; 1995.
- Borenstein M, Hegges LV, Higgins JPT, Rothstein H. Introduction to Meta-Analysis. Third ed. United Kingdom: John Wiley & Sons; 2011.
- Begg CB, Mazumdar M. Operating characteristics of a rank correlation test for publication bias. *Biometrics*. 1994;50(4):1088–1101. doi:10.2307/2533446
- Egger M, Davy Smith G, Schneider M, Minder C. Bias in meta-analysis detected by a simple, graphical test. *BMJ*. 1997;315 (7109):629–634. doi:10.1136/bmj.315.7109.629
- Sterne JA, Sutton AJ, Ioannidis JP, et al. Recommendations for examining and interpreting funnel plot asymmetry in meta-analyses of randomised controlled trials. *BMJ*. 2011;343(jul22 1):d4002. doi:10.1136/bmj.d4002
- 24. Duval S, Tweedie R. Trim and fill: A simple funnel-plot-based method of testing and adjusting for publication bias in meta-analysis. *Biometrics*. 2000;56(2):455–463. doi:10.1111/j.0006-341X.2000.00455.x
- Pogue JM, Yusuf S. Cumulative evidence from randomized trials: utilizing sequential monitoring boundaries for cumulative meta-analysis. *Control Clin Trials*. 1997;18(6):580–593. doi:10.1016/ S0197-2456(97)00051-2
- Pogue JM, Yusuf S. Overcoming the limitations of current meta-analysis of randomized controlled trials. *Lancet*. 1998;351 (9095):47–52. doi:10.1016/S0140-6736(97)08461-4
- Wetterslew J, Thorlund K, Brok J, Gluud C. Trial sequential analysis may establish when firm evidence is reached in cumulative meta-analysis. *J Clin Epidemiol.* 2008;61(1):64–75. doi:10.1016/j. jclinepi.2007.03.013
- Miladinovic B, Hozo I, Djulbegovic B. Trial sequential boundaries for cumulative meta-analysis. *Stata J.* 2013;13(1):77–91. doi:10.11 77/1536867X1301300106
- StataCorp LLC. College Station, TX, USA. Retrieved from: http:// www.stata.com. Accessed May 20, 2020.
- Schwarzer G. meta: an R package for meta-analysis. *R News*. 2007;7 (3):40–45.
- 31. Moher D, Liberati A, Tetzlaff J, Altman DG, PRISMA Group. Preferred reporting items for systematic reviews and meta-analyses: the PRISMA statement. *Ann Intern Med.* 2009;151(4):264–269. doi:10.7326/0003-4819-151-4-200908180-00135
- 32. Qian YX, Dai KS, Zhao LL, Yang XJ. Effects of remote ischemic post-conditioning on platelet activation of AMI patients. *Exp Ther Med.* 2018;16(2):1273–1277. doi:10.3892/etm.2018.6280
- 33. Elbadawi A, Awad O, Raymond R, Badran H, Mostafa AE, Saad M. Impact of remote ischemic postconditioning during primary percutaneous coronary intervention on left ventricular remodeling after anterior wall ST-segment elevation myocardial infarction: A single-center experience. *Int J Angiol.* 2017;26(4):241–248. doi:10.1055/s-0037-1601870

- 34. Lavi S, Abu-Romeh N, Wall S, Alemayehus M, Lavi R. Long-term outcome following remote ischemic postconditioning during percutaneous coronary interventions-the RIP-PCI trial long-term follow-up. *Clin Cardiol.* 2017;40(5):268–274. doi:10.1002/clc.22668
- 35. 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. doi:10.1016/j. ahj.2016.08.004
- 36. Carrasco-Chinchilla F, Muñoz-García AJ, Domínguez-Franco A, et al. Remote ischaemic postconditioning: does it protect against ischaemic damage in percutaneous coronary revascularisation? Randomised placebo-controlled clinical trial. *Heart.* 2013;99 (19):1431–1437. doi:10.1136/heartjnl-2013-304172
- 37. Egger M, Juni P, Bartlett C, Holenstein F, Sterne J. How important are comprehensive literature searches and the assessment of trial quality in systematic reviews? Empirical study. *Health Technol Assess.* 2003;7(1):1–76.
- Schulz KF, Chalmers I, Hayes RJ, Altman DG. Empirical evidence of bias. Dimensions of methodological quality associated with estimates of treatment effects in controlled trials. *JAMA*. 1995;273(5):408–412. doi:10.1001/jama.1995.03520290060030
- Kleinbaum DG, Kupper LL, Morgenstern H. *Epidemiologic Research. Principles and Quantitative Methods*. USA: Van Nostrand Reinhold Company Inc; 1982.
- 40. Turner RM, Bird SM, Higgins JP. The impact of study size on meta-analyses: examination of underpowered studies in Cochrane reviews. *PLoS One.* 2013;8(3):e59202. doi:10.1371/journal.pone.00 59202
- 41. Hempel S, Miles J, Suttorp MJ et al. Methods research report. Detection of associations between trial quality and effect sizes. agency for healthcare research and quality; January 2012. Retrieved from: www.ahrq.gov. Accessed May 20, 2020. Publication No. 12-EHC010-EF
- 42. Fleming TR, DeMets DL. Surrogate endpoints in clinical trials: are we being misled? Ann Intern Med. 1996;125(7):605–613. doi:10.7326/0003-4819-125-7-199610010-00011
- 43. Haller PM, Vargas KG, Haller MC, et al. Remote ischaemic conditioning for myocardial infarction or elective PCI: systematic review and meta-analyses of randomised trials. *Eur Heart J Acute Cardiovasc Care*. 2020;9(1_suppl):82–92. doi:10.1177/20488726 18784150
- 44. Howick J, Chalmers I, Glasziou P, et al. The 2011 oxford CEBM evidence levels of evidence: introductory document. oxford centre for evidence-based medicine. Retrieved from: http://www.cebm.net/ index.aspx?o=5653. Accessed May 20, 2020.
- Atkins D, Best D, Briss PA, et al. GRADE working group. Grading quality of evidence and strength of recommendations. *BMJ*. 2004;328 (7454):1490–1494.

Clinical Epidemiology

Publish your work in this journal

Clinical Epidemiology is an international, peer-reviewed, open access, online journal focusing on disease and drug epidemiology, identification of risk factors and screening procedures to develop optimal preventative initiatives and programs. Specific topics include: diagnosis, prognosis, treatment, screening, prevention, risk factor modification,

Submit your manuscript here: https://www.dovepress.com/clinical-epidemiology-journal

Dovepress

systematic reviews, risk & safety of medical interventions, epidemiology & biostatistical methods, and evaluation of guidelines, translational medicine, health policies & economic evaluations. The manuscript management system is completely online and includes a very quick and fair peer-review system, which is all easy to use.