Biochemical signaling by remote ischemic conditioning of the arm versus thigh: Is one raise of the cuff enough?

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ABSTRACT
Remote Ischemic Conditioning (RIC), induced by brief cycles of ischemia and reperfusion, protects vital organs from a prolonged ischemic insult. While several biochemical mediators have been implicated in RIC's mechanism of action, it remains unclear whether the localization or "dose" of RIC affects the extent of protective signaling. In this randomized crossover study of healthy individuals, we tested whether the number of cycles of RIC and its localization (arm versus thigh) determines biochemical signaling and cytoprotection. Subjects received either arm or thigh RIC and then were crossed over to receive RIC in the other extremity. Blood flow, tissue perfusion, concentrations of the circulating protective mediator nitrite, and platelet mitochondrial function were measured after each RIC cycle. We found that plasma nitrite concentration peaked after the first RIC cycle and remained elevated throughout RIC. This plasma nitrite conferred cytoprotection in an in vitro myocyte model of hypoxia/reoxygenation. Notably, though plasma nitrite returned to baseline at 24 h, RIC conditioned plasma still mediated protection. Additionally, no difference in endpoints between RIC in thigh versus arm was found. These data demonstrate that localization and "dose" of RIC does not affect cytoprotection and further elucidate the mechanisms by which nitrite contributes to RIC-dependent protection.

1. Introduction
Remote ischemic conditioning (RIC) refers to brief sub-lethal ischemia applied to an area remote from the organ being targeted for protection from a future (preconditioning) or past (postconditioning) ischemic insult. In practical terms this is usually achieved in human subjects using a blood pressure cuff placed on an extremity and inflated above systolic blood pressure (often 200 mm Hg), rendering the limb ischemic for a period of 5 min prior to deflation (which permits reperfusion) [1, 2]. RIC was first shown to be cardioprotective in 1993 [3] and subsequent studies have confirmed that other organs such as brain [4, 5] and liver [6] can be similarly protected. RIC is attractive from a clinical perspective as it is minimally invasive compared to traditional ischemic preconditioning, which involves targeting sublethal ischemia to the organ intended for protection, and carries minimal risk and discomfort [7]. While RIC has shown promise in preliminary clinical studies with surrogate endpoints [1, 2, 8] and possible survival benefits [9, 10], other recent large randomized clinical trials have yielded disappointing neutral results [11–14].

The inconsistencies in the benefit of RIC between trials may derive from a lack of understanding of how best to "dose" RIC. The commonly used protocol of 5 min inflation-deflation of a blood pressure cuff placed on an upper extremity repeated for 3–4 cycles has been adopted in clinical studies [11–14] without formal testing of the optimal dose or location in humans. Thus, it is unknown whether some or all patients would benefit from more or less than 3–4 cycles of RIC. Further, it remains unknown whether RIC applied to an upper (arm) versus lower (thigh) extremity would alter the required dose or efficacy of RIC due to considerable differences in vascular and soft tissue mass in the arm versus leg.

One reason for the absence of a human dose titration of RIC is likely due to the lack of understanding of the precise mechanisms that underlie RIC-mediated protection [15–17]. Though a number of mechanisms have been proposed including neural and autonomic...
signaling [18,19], studies demonstrating that organ protection can be transferred from a RIC-treated animal to an untreated animal by the simple transfer of blood suggest that humoral factors are likely imperative to protection [20,21]. In this regard, Rassaf and colleagues recently demonstrated that nitrite, the one electron oxidation product of nitric oxide (NO), was significantly increased in the blood after RIC and necessary and sufficient to confer cardioprotection when applied to an ex vivo heart model of ischemia/reperfusion [20]. This production of nitrite was dependent on the reactive hyperemia (RH) that ensues during the reperfusion phase of RIC and due to oxidation of nitric oxide derived from vascular endothelial nitric oxide synthase [20]. Notably, prior studies have shown that nitrite mimics the effects of RIC and mediates cytoprotection in a number of ischemia/reperfusion models, and this is due to its inhibitory effect on mitochondrial respiration in the target organ [20,22–25]. Thus, the extent of RH and the resulting concentration of plasma nitrite generated represent a quantifiable parameter to measure the “dose” of RIC.

Here we measure RH and plasma nitrite as physiological and biochemical signaling mediators of RIC and use these parameters to quantify the “dose” of RIC delivered using sequential cycles of arm versus thigh RIC. We sought to determine whether RIC applied to a larger endothelial bed (thigh) would generate more nitrite than a smaller bed (arm) and whether consecutive cycles of RIC would further increase production of protective plasma nitrite. We measured platelet mitochondrial respiration in each subject as a target of the action of nitrite as well as utilized an in vitro cultured myocyte model of hypoxia-reoxygenation to demonstrate cytoprotection. We discuss our results in the context of further understanding RIC-dependent biochemical signaling as well as determining how to optimize RIC dosing in the clinic.

2. Methods

2.1. Human subject selection

The study was approved by and in accordance with the University of Pittsburgh Institutional Review Board and informed consent was obtained from each subject. Inclusion criteria included any healthy individual over 18 years of age. Prisoners, pregnant females, history of cardiovascular disease, circulation abnormalities, renal disease, liver disease, and anyone currently taking any medication were excluded. The subjects were not allowed to eat, consume caffeine or alcohol, use nicotine or exercise for 12 h prior to the study.

2.2. Study design

The study was performed in the University of Pittsburgh Applied Physiology Lab. Subjects were randomized to receive RIC applied to the right forearm or right thigh during week 1, with the procedure repeated on the other extremity two weeks later (Fig. 1A). After a baseline blood pressure measurement, the subjects were allowed to get comfortably seated and a blood pressure cuff was placed either on the right forearm or right thigh. An 18 gauge intravenous (IV) catheter was placed in the left antecubital vein (ie contralateral side from the RIC extremity) and was used for all subsequent blood draws. The extremity being used for RIC was secured to the contralateral extremity using cushions and tape assuring the subject was comfortable. This was to minimize movement and permit simultaneous imaging of both ipsilateral and contralateral extremities by laser speckle contrast imaging (LSCI; Perimed AB, Jarfalla, Sweden) throughout the RIC protocol. The LSCI camera was positioned 20 cm over the dorsal surface of the feet or palmar surface of the hands for the thigh and arm occlusion, respectively. Tissue oximetry (StO2; Inspectra 650, Hutchinson Technologies, Hutchinson, Minnesota) pads where placed on both thenar eminences or both halluces for the thigh and arm occlusion, respectively. Stable baseline recordings were collected using the LSCI and StO2 devices for one minute prior to the first cuff occlusion during which time a baseline blood draw (14 ml) was collected (Fig. 1B). RIC was initiated by inflating the blood pressure cuff to 200 mmHg to cause vaso-occlusion. This was at least 50 mm Hg greater than all subjects’ systolic blood pressure, measured at the beginning of each study day. The blood pressure cuff was maintained at 200 mmHg for 5 min with continuous monitoring of pressure through an in-line manometer. After 5 min the cuff was rapidly released permitting reperfusion, which in all cases could be verified visually by witnessed hyperemia. Additional blood samples were obtained four minutes after cuff release. LSCI and StO2 recording continued until at least 5 min after the final cuff release. The subjects returned 24 h later for the final blood draw (Fig. 1B).

2.3. Whole blood and plasma nitrite measurements

For plasma and whole blood nitrite measurements, blood was collected in EDTA. Whole blood (1 ml) was immediately mixed with nitrite preservation solution (0.8 mol/L ferricyanide, 10 mmol/L N-ethylmaleimide, and 1% NP-40) at a 4:1 ratio and frozen for nitrite analysis as previously described [26]. The remaining blood (4 ml) was centrifuged within 5 min at 1000 × g, 5 min, 4 °C to obtain plasma. Plasma was immediately separated from the RBC pellet and frozen for nitrite analysis. Plasma and whole blood nitrite concentrations were measured by tri-iodide based reductive chemiluminescence as previously described [26].

2.4. Platelet isolation and mitochondrial analysis

At each time point blood (8 ml) was collected in citrate (CPT BD Vacutainer) and processed within one hour of collection. Platelet rich plasma and platelets were isolated by differential centrifugation as previously described [27,28] and outlined in Supplemental Methods. Oxygen consumption rate (OCR) was measured in isolated platelets (50×10^6/well) by Seahorse Extracellular Flux analysis as previously described [27,29]. A bioenergetic profile was generated by measuring basal OCR followed by OCR in the presence of oligomycin A (2.5 µmol/L), FCCP (0.7 µmol/L), and rotenone (15 µmol/L). Values reported are rotenone subtracted reflective of mitochondrial specific OCR. In separate experiments platelets were incubated with MitoSOX (5 µmol/L; 10 min; Invitrogen, Carlsbad, CA) and mitochondrial superoxide generation quantified as previously described [27,29].

2.5. Tissue oximetry

Two InSpectra Tissue Spectrometers Model 650 were used, each with its own probe as previously described [30]. The pads were placed over each thenar eminence of the subject’s hands during RIC performed on the forearm, or on each hallux, when RIC was performed on the thigh. Each machine was turned on at least 5 min prior to the beginning of RIC and a stable baseline verified for 1 min before experiment start. The devices recorded data throughout the study and were stopped > 5 min after the final vaso-occlusion. The StO2 data (expressed as a saturation percentage from 0 to 100) was analyzed for the baseline tissue oxygenation, the change from baseline to the RH peak after each vaso-occlusion, the time from cuff deflation to RH peak after each cycle, and the nadir five minutes after each VO cycle. Measurements were obtained simultaneously in the extremity ipsilateral and contralateral to the RIC. Since RH was not noted contralaterally, the “peak” and “nadir” corresponded in time to the ipsilateral side. All values are expressed as percent StO2 relative to baseline (100%). The StO2 measurements ranged from 5% to 98%, thus never reaching the limits of detection (0–100%).
Laser speckle contrast imaging (LSCI)

The PeriCam PSI LSCI system was used as previously described [31]. LSCI involves a near infrared (780 nm) laser, which is refracted based on movement, forming a random 'speckled' pattern. Changes in the speckled pattern correspond to red blood cell movement, thus allowing a relative measurement of blood flow rate [32]. The camera was placed 20 cm above the palms or ventral aspect of the feet. Baseline LSCI measurements were obtained for one minute prior to vaso-occlusion and recorded continuously until > 5 min after the last RIC cycle. Blood was obtained at baseline, 4 min into reperfusion (i.e. 1 min before next cycle) and at 24 h.

In vitro anoxia-reoxygenation model

H9c2 myocytes purchased from ATCC (Rockville, MD) were maintained in normoxia (21% O2, 5% CO2, 94% N2, 5 h), in modified Esumi buffer (137 mmol/L NaCl, 12 mmol/L KCl, 0.5 mmol/L MgCl2, 0.9 mmol/L CaCl2, 20 mmol/L HEPES; 20 mmol/L 2-Deoxy-d-Glucose (2-DG), pH 6.2) as described by previous publications [33–36]. Cells were then reoxygenated in the same buffer in normoxia (21% O2, 5% CO2) for 1 h. In some cases plasma (100 µl) from subjects who underwent RIC was added to the cells during the hypoxic period.

Lactate dehydrogenase (LDH) activity

LDH activity in the media was measured spectrophotometrically by measuring the decrease in NADH at 340 nm and was expressed as a percent of total LDH in the media and lysed cells.

Statistical analysis

Statistical analyses were performed using Prism 6.05 (GraphPad Software Inc., La Jolla, CA). All data are reported as mean ± standard deviation and analyzed using parametric analyses based on normal distribution. Comparisons between results over time and between groups using repeated measures 1-way ANOVA with post-hoc Sidak's comparison made between cycle 1 and 4 data. If the ANOVA was significant, paired t-tests were performed to compare baseline values to cycle 1–4 separately. All other comparisons between two groups utilized unpaired t-tests. In all cases p < 0.05 was considered significant and 0.05 < p < 0.10 was considered indicative of a trend.

Results

Ten healthy subjects (eight males and two females) were enrolled and completed the study. Subject demographics are outlined in Table 1. All subjects underwent RIC administered to the thigh and arm (Fig. 1). The only adverse effect reported in the context of the study was mild to moderate pain during ischemia and early reperfusion (during RH), and this was experienced by all participants. All subjects reported that pain during thigh RIC was greater than arm RIC and that the perception of pain diminished with each subsequent cycle. None of the participants requested that the procedure be stopped due to the pain severity and all subjects reported resolution of pain shortly after the last RIC cycle.

| Table 1 | Subject characteristics (n=10). |
|---------|-------------------------------|
| Age     | 34.5 (26.5–38.75)             |
| Gender, male | 8 (80%)                |
| SBP     | 118 (114–122)               |
| DBP     | 78 (68–80)                  |
| Lactate | 0.6 (0.58–0.65)             |
| Hemoglobin | 16.7 (14.83–17.15)      |
| Hematocrit | 44.5 (40.75–47)          |

Values represent median (interquartile range) or n.
3.1. RIC causes reactive hyperemia measured by blood flow and tissue oxygen saturation

We first compared blood flow and tissue oxygen saturation after each cycle of RIC administered to the arm versus the thigh. Consistent with prior studies, RH was observed after each RIC cycle in both the arm and thigh (when measured by LSCI) and tissue oxygen saturation (measured by StO2) was also significantly increased. Notably, RH was observed earlier after cuff deflation (initiation of reperfusion) when measured by LSCI compared to StO2 in both extremities (all \( p < 0.01 \); Fig. 2A–B). Additionally, comparison of RIC in each extremity showed that RH peaked significantly earlier following arm RIC compared to thigh when measured by either LSCI or StO2 (Fig. 2A–B). Despite an earlier peak with arm RIC, blood flow measured by LSCI was comparable in the forearm (278 ± 85% of baseline) versus the thigh (300 ± 92% of baseline; Fig. 2C) at the peak RH (after cycle 1). Tissue oxygen delivery measured by StO2 after RIC in the forearm (116 ± 6% of baseline) was slightly lower than that generated by RIC delivered to the thigh (127 ± 11% of baseline; \( p = 0.013 \); Fig. 2D). The blood flow and StO2 nadirs generally showed non-significant trends towards being increased after the first RIC cycle with little variation over subsequent cycles (Supplemental Fig. 1A–B). In the case of the StO2 nadir after thigh RIC, these differences were small (7% absolute increase) but significant (Supplemental Fig. 1C–D). Blood flow and StO2 measured in the extremity contralateral to that used for RIC did not change throughout the period of observation (data not shown).

3.2. Circulating nitrite levels peak after the first cycle of RIC

To determine whether RH observed after RIC resulted in the production of nitrite, we measured plasma nitrite at baseline and after 4 cycles of RIC. Plasma nitrite concentration at the end of 4 cycles of RIC was increased after arm or thigh RIC in eight out of ten subjects (Fig. 3A–B), with this elevation reaching significance after RIC in the arm (Fig. 3A). Notably, RIC (4 cycles) was ineffective in elevating plasma nitrite in the arm or thigh of the same two subjects. We next determined the effect of each cycle of RIC on the concentration of nitrite in platelet rich plasma (PRP). All ten subjects showed an increase in PRP nitrite after the first RIC cycle and at this peak, both arm (0.39 ± 0.06 µmol/L) and thigh (0.20 ± 0.06 µmol/L) PRP nitrite was significantly augmented compared to baseline (0.22 ± 0.04 and 0.24 ± 0.03 µmol/L respectively; Fig. 3C). Nitrite levels remained significantly elevated for each subsequent cycle, but decreased by ~10% with each cycle of RIC (Fig. 3C). The peak nitrite concentration and its rate of decrease did not significantly differ in the arm compared to thigh. In both cases, concentration was not significantly different from baseline 24 h after RIC (Fig. 3C). No significant change in whole blood nitrite was detected (data not shown).

3.3. RIC inhibits platelet mitochondrial respiration

Mitochondria have been identified as a cytoprotective target of both nitrite and RIC. To determine whether RIC modulates mitochondrial function, we next measured the platelet mitochondrial respiration rate in each subject after each cycle of RIC. RIC performed in both the arm and thigh significantly inhibited platelet basal and maximal uncoupled mitochondrial respiration, but had no effect on mitochondrial proton leak (Fig. 4A). RIC performed in the arm decreased basal platelet mitochondrial OCR from baseline values of 109.3 ± 19.7 pmol/min to 72.7 ± 13.5 pmol/min by cycle 3 representing a maximal inhibition of 34%. Interestingly, significant inhibition was still present 24 h after RIC administered to the thigh resulting in a similar pattern of inhibition of basal OCR with maximal inhibition reaching a slightly greater level than arm (41%; Fig. 4B). Maximal uncoupled OCR was also inhibited to a similar extent by both arm and thigh RIC (Fig. 4C). However, no significant change was observed in mitochondrial proton

Fig. 2. RIC to the arm or thigh causes reactive hyperemia (RH) and increased tissue oxygen delivery by StO2. The time to (A) peak blood flow (measured by LSCI) and (B) peak tissue oxygen saturation after RIC applied to either the arm (black bars) or thigh (white bars). Peak (C) blood flow and (D) tissue oxygenation after each cycle of RIC (C1–C4) applied to the arm (closed circles) or thigh (open squares). All measurements are normalized to the baseline (BL) for the respective appendage. (A–B) *\( p < 0.01 \) versus arm measurement for the same cycle. (C–D) \( * p < 0.01 \) versus baseline measurement. §\( p < 0.05 \) for thigh versus arm. N=10. All data are mean ± SEM.
**Fig. 3.** RIC increases plasma nitrite concentration. (A–B) Plasma nitrite concentrations for each subject at baseline and after 4 cycles of RIC applied to the (A) arm or (B) thigh. (C) Nitrite concentration in the platelet rich plasma at baseline (BL) and after each cycle of RIC (C1-C4) applied to the arm (black bars) or thigh (white bars) as well as at 24 h. Data are mean ± SEM. *p < 0.01 and #p < 0.05 compared to baseline for each appendage. N=10.

**Fig. 4.** RIC inhibits platelet mitochondrial respiration. (A) Representative Seahorse XF Analysis trace measuring oxygen consumption rate (OCR) in platelets from one subject at baseline (gray circles), after 4 cycles of RIC applied to the arm (open squares), and 24 h later (black triangles). For each trace, after a baseline reading, oligomycin A (2.5 µmol/L), FCCP (0.7 µmol/L) and rotenone (15 µmol/L) were added as noted by the arrows. (B–C) Quantitation of (B) basal and (C) maximal respiration rates calculated from traces similar to that shown in (A). (D) Mitochondrial superoxide production by platelets at baseline and after 4 cycles of RIC applied to the arm or thigh. (B–D) are means ± SEM. *p < 0.01 compared to baseline for the respective appendage. N=10.
versus thigh. Consistent with RH, we found that blood flow increases after each RIC cycle, and we extended previous studies to show that this results in a smaller but significant increase in tissue oxygen saturation compared to baseline in the ipsilateral extremity. On a biochemical level, circulating nitrite concentration increased to the greatest extent after the first cycle of RIC but remained elevated after subsequent cycles. Nitrite production was concomitant with a progressive inhibition of platelet mitochondrial respiration and a trend to increased superoxide production. Plasma taken from subjects after RIC protected myocytes from cell death in an in vitro model of hypoxia-reoxygenation and the protection for plasma from cycles 1–3 was dependent on nitrite.

Our data confirm prior studies noting that RIC results in RH [20,37] and extend these results to demonstrate that increased blood flow results in a significant augmentation of tissue oxygen saturation. Notably, the increases in tissue oxygen saturation peaked later and were 10-fold lower in magnitude than the increase in blood flow. This may be a reflection of increased tissue oxygen consumption after ischemia which would limit the measurement of oxygen saturation or may reflect a more limited blood supply to the regions where we assessed tissue oximetry. It also must be noted that the duration of peak RH blood flow was brief with peaks lasting only several seconds and return to baseline (pre-cuff occlusion) blood flows within 120 s such that oxygen delivery is likely more reflective of the area under the blood flow curve rather than just the peak. The lack of increased blood flow in the contralateral extremity or changes in the nadir blood flow before the subsequent RIC cycle are consistent with this and imply that RIC is not causing a prolonged change in systemic blood flow, but rather having a discrete blood flow effect limited to the tissue bed undergoing ischemia and reperfusion.

It is interesting that we failed to see further augmentation of RH with multiple cycles of RIC or when RIC was applied to the thigh, which encompasses a much larger tissue/endothelial bed than the arm. These findings are consistent with those of Johnson et al. who showed in a murine model of RIC that increasing cycle number or performing RIC on two hindlimbs instead of one did not confer greater protection [38]. In our study, while blood flow did peak significantly earlier in the forearm than thigh, this may merely be a reflection of the distance from the cuff to the site of LSCI data collection. With respect to multiple cycles of RIC, it is possible that the 5 min cycle length achieves maximal dilation in healthy volunteers. Consistent with this, Raff and colleagues saw little increase in forearm blood flow (FBF) measured by plethysmography between 2 and 5 min of occlusion but noted a difference between 1 and 2 min of occlusion [39]. Notably, these authors did report an increase in FBF when the blood pressure cuff was applied to the wrist (ie forearm) versus upper arm [39], though differences in blood flow measurement methodology could account for this.

We measured RIC-induced increases in plasma nitrite of the same magnitude as previously reported by Rassaf and colleagues in healthy subjects [37]. Our results are consistent with NOS-dependent production of NO, which is oxidized to nitrite during RH [39,40]. However, this is the first report that demonstrates that circulating nitrite concentration peaks after the first cycle of RIC and decreases with subsequent cycles. This is likely due to ischemic consumption of nitrite. Nitrite is a recognized endocrine reservoir of NO that can be reduced to bioavailable NO in conditions of hypoxia and acidosis, such as those present during ischemia [19,23,41,42]. Hypoxic metabolism of nitrite results in the generation of a number of NO-dependent species including iron-nitrosylated and S-nitrosated proteins [42], which are markers of NO production and can be responsible for downstream signaling. Prior studies demonstrate that RIC-dependent NO generation is maximal after 1 min of ischemia [39]. Thus our 5 min ischemic period potentially results in a net consumption rather than augmented

3.4. Plasma from the first cycle of RIC confers the greatest cytoprotection ex vivo

We next sought to determine the cytoprotective potency of each cycle of RIC utilizing an in vitro hypoxia-reoxygenation model (Fig. 5). In this model, cultured myocytes subjected to hypoxia (1% O2; 5 h) and subsequent reoxygenation (21% O2; 1 h) showed ~50% cell death. Treatment of the cells with PRP from each cycle of RIC (from either arm or thigh) as well as PRP collected 24 h after RIC significantly attenuated cell death. However, PRP from the first cycle of RIC showed the greatest cytoprotection (Fig. 5). To determine whether the nitrite generated by RIC was responsible for the cytoprotection, PRP samples were treated with acidified sulfanilamide to bind bioavailable nitrite. Scavenging of nitrite significantly inhibited the cytoprotective effect of PRP from the first through the fourth RIC cycle, but had no effect on cytoprotection from the 24 h sample (Fig. 5).

4. Discussion

In this study we compared the physiological and biochemical effects of 1–4 cycles of RIC applied to the arm versus thigh of healthy subjects. Our major finding was that physiological (RH and tissue oxygen saturation) and biochemical (plasma nitrite concentration) effects of RIC were not significantly different when RIC was applied to the arm
production of nitrite. This finding leads us to wonder whether shorter ischemic duration or fewer cycles would actually increase nitrite generation at least in subjects with “healthy” endothelium.

The consumption of nitrite has potential consequences for the number of RIC cycles required to mediate cytoprotection. Though it did not reach statistical significance, plasma collected after the first cycle of RIC showed the greatest level of cytoprotection in our in vitro model of hypoxia-reoxygenation. Interestingly, though plasma from subsequent cycles showed significant cytoprotection, this cytoprotection appeared to be less dependent on nitrite (evidenced by the decreased effect of acidified sulfanilamide). We and others have previously shown that acute nitrite-dependent cytoprotection after ischemia/reperfusion is dependent on the S-nitrosation and inhibition of complex I, which decreases respiratory rate [22,25,43]. Our data for the four cycles of RIC are consistent with this mechanism as we show that platelet mitochondrial respiration is progressively more inhibited with each RIC cycle. While we did not measure S-nitrosation in this study, the consumption of nitrite is consistent with the production of S-nitrosothiols.

The data demonstrating that plasma collected 24 h after RIC also mediated cytoprotection is consistent with the paradigm that RIC mediates two temporal windows of protection [44]. The acute window (minutes to hours after RIC is administered) is traditionally thought to be due to biochemical mediators, while delayed cytoprotection (hours to days after RIC) is usually attributed to changes in gene or protein expression [44]. Our data suggests the presence of plasma nitrite is required for acute protection (cycles 1–4), while nitrite does not have to be present to mediate protection 24 h after RIC is administered. We have previously shown that nitrite induces delayed protection through the production of mitochondrial ROS and subsequent upregulation of AMP Kinase, and we observe a trend to increased mitochondrial ROS production in this study. However, the protective effect observed with the plasma at 24 h could also be mediated by other factors including the release of high mobility group box-1 [45] or adenosine [46,47].

Our data have potential implications for the interpretation of the recent neutral RIC clinical trials [13,14]. We recently showed that subjects with comorbid endothelial dysfunction show diminished RH [31] and it is known that these subjects show minimal increases in nitrite after RIC [37]. Thus an alternative explanation for the failure to translate RIC to humans may be a failure to appreciate dosing requirements in the setting of health vs. disease. However, our data suggesting progressive mitochondrial inhibition with multiple cycles which persists at 24 h may suggest a more delayed effect could show efficacy in these subjects. Further investigation is required to elucidate the exact mechanism of RIC-dependent mitochondrial inhibition and its effect on delayed protection.

A limitation of our study is the inclusion of only healthy subjects as well as only Caucasian subjects. Prior studies suggest that blood flow regulation may be variable between subjects of different race [48–50]. It is therefore important to replicate these results in other healthy and diseased populations to fully understand the effects of RIC which will be needed to achieve the optimal dose. Additionally, our study has shown that while nitrate may stimulate signaling mediating delayed protection, its actual presence at 24 h is potentially not required. Future studies should expand on this pathway. Notwithstanding the limitations we have demonstrated that RIC increases blood flow and tissue oxygen saturation after RIC. These increases correspond with augmented plasma nitrite which is associated with mitochondrial inhibition and cytoprotection. These effects of RIC are similar regardless of extremity used and suggest that while nitrate-mediated protection is maximized after the initial cycle of RIC, significant protection is retained with up to four cycles.

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**Author contributions**

CD, FXG, JCR, and SS conceived, designed and directed the study. CD and SS analyzed and interpreted the data and drafted the manuscript. MT executed and analyzed physiological data. CC, GH and NK executed measurement of nitrite and mitochondrial data.

**Competing financial interests**

None to report.

**Appendix A. Supplementary material**

Supplementary data associated with this article can be found in the online version at http://dx.doi.org/10.1016/j.redox.2017.03.010.

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