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Prostaglandin contribution to postexercise hyperemia is dependent on tissue oxygenation during rhythmic and isometric contractions

Rehan T. Junejo1 | Clare J. Ray2 | Janice M. Marshall2

1School of Sport, Exercise & Rehabilitation Sciences, College of Life & Environmental Sciences, Birmingham, UK
2Institute of Clinical Sciences, College of Medical and Dental Sciences, University of Birmingham, Birmingham, UK

Correspondence
Janice M. Marshall, Institute of Clinical Sciences, College of Medical and Dental Sciences, University of Birmingham, Birmingham B15 2TT, UK.
Email: j.m.marshall@bham.ac.uk

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Abstract
The role of prostaglandins (PGs) in exercise hyperemia is controversial. We tested their contributions in moderate intensity forearm exercise, whether their release is oxygen (O2)-dependent or affected by aging. A total of 12 young (21 ± 1 years) and 11 older (66 ± 2 years) recreationally active men performed rhythmic and isometric handgrip contractions at 60% maximum voluntary contraction for 3 min during air breathing after placebo, after cyclooxygenase (COX) inhibition with aspirin, while breathing 40% O2 and during their combination (aspirin + 40% O2). Forearm blood flow (FBF) was recorded with venous occlusion plethysmography (forearm vascular conductance (FVC): FBF/mean arterial pressure). Venous efflux of PGI2 and PGE2 were assessed by immunoassay. Postcontraction increases in FVC were similar for rhythmic and isometric contractions in young and older men, and accompanied by similar increases in efflux of PGI2 and PGE2. Aspirin attenuated the efflux of PGI2 by 75%–85%, PGE2 by 50%–70%, (p < .05 within group; p > .05 young versus. older), and postcontraction increases in FVC by 22%–27% and 17%–21% in young and older men, respectively (p < .05 within group and young versus. older). In both age groups, 40% O2 and aspirin + 40% O2 caused similar inhibition of the increases in FVC and efflux of PGs as aspirin alone (p < .05 within group). These results indicate that PGs make substantial contributions to the postcontraction hyperemia of rhythmic and isometric contractions at moderate intensities in recreationally active young and older men. Given PGI2 is mainly released by endothelium and PGE2 by muscle fibers, we propose PG generation is dependent on the contraction-induced falls in O2 at these sites.

KEYWORDS
exercise-hyperemia, oxygen-dependent, prostaglandins
1 | INTRODUCTION

There is substantial evidence that prostaglandins (PGs) contribute to exercise hyperemia, but there is also conflicting evidence. For example inhibition of cyclooxygenase (COX) attenuated postcontraction hyperemia associated with rhythmic and isometric contractions of forearm and leg (Cowley, Stainer, Rowley, & Wilcox, 1985; Duffy, New, Tran, Harper, & Meredith, 1999; Kilbom & Wennmalm, 1976; Win & Marshall, 2005). However, it was separately reported that COX inhibition had no effect on hyperemia during rhythmic contraction in forearm (Shoemaker, Naylor, Pozeg, and Hughson (1996); Mortensen, González-Alonso, Damsgaard, Saltin, & Hellsten, 2007), and that combined inhibition of COX and nitric oxide (NO) synthase (NOS) was required to attenuate the hyperemia (Boushel et al., 2002; Mortensen et al., 2007). These findings led to the suggestion that PGs and NO contribute synergistically, rather than independently, to exercise hyperemia (Boushel et al., 2002; Mortensen et al., 2007). In contrast, the observation that the attenuating effect of COX inhibition on hyperemia during rhythmic contraction was transient, whereas that of NOS inhibition was sustained led to the proposal that the contribution of PGs to exercise hyperemia is independent of NO, and can be compensated for by other dilator/s (Schrage, Joyner, & Dinenno, 2004).

A possible explanation for these disparities is that they reflect differences between studies in exercise intensity and a possible fall in partial pressure of O₂ (PO₂) within muscles. For, those which suggested a relatively minor contribution of PGs to exercise hyperemia used exercise intensities of ≤20% maximum (Mortensen et al., 2007; Schrage et al., 2004; Shoemaker et al., 1996), whereas those suggesting a substantial contribution used intensities of ≥60% maximum (Kilbom & Wennmalm, 1976; Win & Marshall, 2005). In line with this idea, PGE₂ release into muscle interstitium during isometric contraction was enhanced by arterial occlusion, which would have greatly reduced tissue PO₂ (Symons, Theodosy, Longhurst, & Stebbins, 1991). Moreover, the release of PGI₂ into venous efflux and PGI₂ and PGE₂ into muscle interstitium during rhythmic exercise was directly related to O₂ consumption (VO₂) and exercise intensity (Karamouzis, Karamouzis, & Vamvakoudis, 2001; Zoladz, Majerczak, Duda, & Chlopiczki, 2009). Furthermore, the postcontraction hyperemia of isometric handgrip contraction at 60% maximum voluntary contraction (MVC) was similarly attenuated by breathing 40% O₂ or COX inhibition, whereas combined COX inhibition and 40% O₂ had no greater effect (Win & Marshall, 2005). Also, when breathing 40% O₂ was restricted to the period of isometric contraction, postcontraction hyperemia was attenuated, whereas 40% O₂ from contraction cessation had no such effect (Fordy & Marshall, 2012). Thus, we proposed 40% O₂ alleviates the fall in tissue PO₂ decreasing the generation of PO₂-dependent PGs by endothelium and/or skeletal muscle (Fordy & Marshall, 2012; Frisbee, Maier, Falk, Roman, & Lombard, 2002; Marshall & Ray, 2012; Michiels, Arnould, Knott, Dieu, & Remacle, 1993; Win & Marshall, 2005). However, uncertainty remains over this interpretation because the higher PO₂ attained with 40% O₂ may prevent the action, rather than release of PGs. Furthermore, as muscle blood flow is limited persistently during isometric, but intermittently during rhythmic contractions (Kagaya & Homma, 1997; McNeil, Allen, Olympico, Shoemaker, & Rice, 2015; Van Beekvelt, Shoemaker, Tschakovsky, Hopman, & Hughson, 2001), the fall in tissue PO₂ during isometric contraction may have greater effects on PG synthesis.

Separately, there is also uncertainty over the effects of aging on the contribution of PGs to exercise hyperemia. In contrast to young subjects (Schrage et al., 2004), COX inhibition had no effect on hyperemia during 10% MVC rhythmic contractions in older subjects, leading the authors to conclude that the role of PGs is lost with aging (Schrage, Eisenach, & Joyner, 2007). Furthermore, forearm vasodilator responses to infused PGI₂ were smaller in older than young subjects (Nicholson, Vaa, Hesse, Eisenach, & Joyner, 2009). However, the older subjects who took part in those studies were relatively inactive (Nicholson et al., 2009; Schrage et al., 2007). Although muscle VO₂ is maintained during submaximal exercise in both recreationally active and sedentary older men, exercise hyperemia was only blunted in latter (Poole, Lawsonson, Kim, Brown, & Richardson, 2003; Proctor et al., 2003). Thus, the loss of PG involvement in exercise hyperemia with aging (Schrage et al., 2007) may have reflected aging, sedentariness, the light intensity rhythmic exercise and small fall in muscle PO₂ (Van Beekvelt et al., 2001), and/or impaired responsiveness to PGs (Nicholson et al., 2009; Schrage et al., 2007).

With this background, we hypothesized that in recreationally active young and older men, rhythmic and isometric contractions at moderate intensity of 60% MVC would increase venous efflux of both PGI₂ and PGE₂, but their efflux would be greater in isometric contraction and greater in young men. Furthermore, breathing 40% O₂ or COX inhibition would similarly attenuate postcontraction hyperemia and PG efflux following rhythmic and isometric contractions in both young and older men. We focussed on men to avoid the complicating facilitatory influences of estrogen on COX and NOS activity (Orshal & Khalil, 2004). Some of these results have been published in brief (Junejo, Ray, & Marshall, 2014, 2015).

2 | METHODS

This study was approved by the University of Birmingham's Ethical Review Committee (Project ERN_12-1377) and
undertaken in accordance with the revisions of Declaration of Helsinki.

2.1 | Subjects

A total of 12 young and 11 older men (age 21 ± 1 and 66 ± 2 years, respectively) who were students or staff of the University of Birmingham, or members of The Birmingham 1000 Elders Group were recruited for the study. All regularly participated in recreational activities, but none were in training. None took prescribed medication. Prior to the experimental session, participants were requested to refrain from caffeinated drinks and heavy meals for ≥12 hr; alcohol consumption, nonsteroidal anti-inflammatory drugs, or strenuous exercise for ≥24 hr.

2.2 | Experimental procedures

During a familiarization visit, written informed consent was obtained following explanation of the protocol. MVC of the dominant hand was recorded using a handgrip dynamometer (Lafayette 70718, Loughborough, UK) as an average of 3 maximal effort 5 s handgrip contractions separated by at least 30 s. Habituation to experimental conditions was aided by practising the protocol during this session.

For each experimental session, the subject rested supine on a couch with the backrest at ~65° and both arms supported at heart level. An intravenous cannula (22–24 G, BD Venflon, BD and Co.) was inserted in the antecubital vein of the dominant (exercising) forearm to allow blood sampling. Beat by beat arterial blood pressure (ABP) was monitored from a finger on the nondominant hand using photoplethysmography (Finapres, Ohmeda 2300, Englewood, USA); Mean ABP (MABP), and heart rate (HR) were computed over the same cardiac-cycle/s and expressed as conductance units (CU).

2.3 | Experimental protocol

The protocol was carried out on four different days in a randomized, single-blind, cross-over design under control conditions (placebo/air breathing) and with three treatments: aspirin, 40% O2, and the combination; aspirin + 40% O2. On arrival, the subject consumed an orange-flavoured drink either without (placebo), or with aspirin (600 mg), which produces its maximal inhibition of COX at ~30 min (Heavey, Barrow, Hickling, & Ritter, 1985). The recording equipment and venous cannula were put in place and an equilibration period of ~20 min was allowed; the cannula was kept patent by infusion of 3 ml sterile saline bolus immediately after insertion and 15–20 s prior to each blood sampling (PosiFlush SP Syringe 0.9% NaCl, BD and Co.). Subject then breathed either medical grade air or 40% O2 via the facemask (100% O2 titrated using a Venturi valve); an O2 sensor (ProOx 110, Biospherix) ensured appropriate delivery of the required O2 concentrations. After 5 min, resting venous samples were taken and baseline measurements of FBF were recorded and at 30 min after placebo/aspirin drink, the subject performed rhythmic handgrip contractions at 60% MVC (1 s contraction: 1 s relaxation) for 3 min. FBF was recorded immediately the final contraction ceased (0 s), at 30 s, 1 min and at 1 min intervals until 7 min. Values of MABP and HR were extracted for analysis over the same time periods as the FBF measurements and at the mid-point of the 1st, 2nd, and 3rd min of contractions. In all subjects, venous blood samples were taken for blood gas and metabolite analysis (see below) at rest, immediately the contractions ceased (0 s), 3-, 5-, and 7-min postexercise. Furthermore, in six randomly selected subjects from each age group, additional blood samples were taken at rest and immediately the contractions ceased for assay of PG metabolites (see below). The subject then rested whilst breathing normal room air for 25 min.

This protocol was repeated except the subject performed 60% MVC isometric handgrip contraction for 3 min, or as long as possible, with vigorous verbal encouragement. Care was taken to ensure the subject did not engage in a Valsalva manoeuvre. Blood samples were taken as described for rhythmic contractions.

2.4 | Blood analysis

Venous blood samples were collected into a 1 ml syringe for immediate analysis of pO2, pCO2, pH, K+, and lactate (GEM Premier 4000, Instrumentation Laboratory). Samples
for PG assay were collected into ice-cold heparinized vacu-
tainers with 1 µl/ml of 10 µM indomethacin (Sigma Aldrich).
Vacutainers were centrifuged at 700 g at 4°C for 20 min
(Mistral 3000i, MSE Ltd). Supernatant plasma was collected
in 1 ml Eppendorf tubes, snap frozen in liquid nitrogen and
stored in −80°C. Enzyme-linked immune-sorbent assays
(ELISAs) were performed for PGI₂ and PGE₂ derivatives
[6-Keto PGF₁α and PGE₂ metabolite (PGEM)] using com-
mercially available kits (Cayman Chemical Co.). The efflux
of each PG was calculated for each subject at baseline and on
cessation of exercise as the product of venous PG concentra-
tion and the associated FBF value.

2.5 Data Analysis

Hemodynamic and handgrip contraction data were digitally
collected at 400 Hz using Power-Lab and Lab Chart data
acquisition software (Version 7.3.3 AD Instruments) and
stored on a desktop computer (Dell Inc.). Data were ana-
ysed on JMP for Windows (Version 13.0.0, SAS Institute
Inc.). Tension time integral (TTI) from the dynamometer
output, relative percentage change in PG efflux and in post-
contraction hyperemia (FVC values from 0 s until 7 min
after contraction) were computed offline. Participant char-
acteristics were compared between the age groups using
Student's t-test. Factorial mixed-model analysis of variance
(ANOVA) was used to identify time, treatment, age, and
interaction effects. Additionally, time effect within each
treatment were analyzed using one-way repeated-measures
ANOVA. Once a significant effect was detected, Tukey's
HSD was used as post hoc test. Statistical significance was
set at p < .05.

3 RESULTS

3.1 Subject characteristics

Both groups were closely matched for height, weight, body
mass index, and forearm circumference; cardiovascular base-
lines were also similar. However, as expected, age and MVC
were different between the two groups. These variables along
with all nutritional supplementation use are shown Table 1.

3.2 Magnitude of forearm contractions

The older men achieved a MVC that was ~25% less than that
of the young men (p < .05; Table 1). Within each age group,
the TTI for rhythmic handgrip was lower than for isometric
handgrip (Table 2). The TTIs for rhythmic and isometric con-
tractions were not significantly different between older and
young men, but fewer young men completed the full 3 min of
isometric handgrip. None of the treatments (aspirin, 40% O₂,
or aspirin + 40% O₂) affected TTI for rhythmic or isometric
contraction in young, or older men.

3.3 Hemodynamic responses

Under control conditions, rhythmic and isometric contrac-
tions evoked smaller increases in HR in older men (p < .05;
Figures 1 and 2), as described previously (Proctor et al., 2003)
for submaximal exercise in recreationally active older men.
None of the three treatments affected the increases in HR or
MABP evoked by rhythmic or isometric handgrip contrac-
tions in young, or older men (Figures 1 and 2). However,
all three treatments caused similar attenuation in absolute
terms, of the postcontraction increases in FBF and FVC for
rhythmic and isometric contraction in both young and older men (Figures 1 and 2).

3.4 | Venous prostaglandins (PGs)

Under control conditions, baseline venous concentrations of 6-Keto PGF$_{1\alpha}$ and PGEM did not differ between rhythmic and isometric handgrip contraction in either age group. There were also no differences between young and older men for baseline concentrations of 6-Keto PGF$_{1\alpha}$ or PGEM (see online Figure S1, which shows the effects of the three treatments on venous PG concentrations before and following handgrip contractions). Considering the venous efflux data, under control conditions, both rhythmic and isometric contractions led to significant increases in the venous efflux of 6-Keto PGF$_{1\alpha}$ and PGEM in both young and older men (Figure 3). There were no differences between the age groups for venous efflux of either PG, but there was a strong trend for PGEM efflux to be greater in older men (Figure 3; $p = .05$ for isometric contractions). The effluxes of both PG metabolites caused by rhythmic and isometric contractions were substantially attenuated after aspirin in both young and older men (Figure 3). They were also reduced in both groups by 40% O$_2$ and by combined aspirin + 40% O$_2$ (Figure 3).

3.5 | Venous PO$_2$ (PvO$_2$) and other metabolites

Overall, in young men, PvO$_2$ values following both rhythmic and isometric handgrip contractions were significantly higher during 40% O$_2$ and combined aspirin + 40% O$_2$ than during control, or aspirin conditions (Treatment effects: $p < .05$ in each case, see Figure 4, Tables S1 and S2). Considering the time points in more detail, immediately following rhythmic and isometric contractions (i.e., at time 0), PvO$_2$ values were lower than their respective baselines, except during 40% O$_2$ and combined aspirin + 40% O$_2$ for rhythmic contraction and during 40% O$_2$ for isometric contraction (Figure 4). However, from 3 to 7 min, PvO$_2$ values were higher than their respective baselines during 40% O$_2$ and combined aspirin + 40% O$_2$ for rhythmic contractions and following isometric contractions (Figure 4).

In contrast, in older men, PvO$_2$ values were not different between treatment conditions following rhythmic, or isometric contractions (Treatment effects: $p = .91$ and $p = .99$, respectively, see Figure 4). In fact, immediately following rhythmic and isometric contractions (at time 0), PvO$_2$ values were lower than their respective baselines under all treatment conditions irrespective of whether 40% O$_2$ was breathed ($p < .01$ vs. respective baselines). Furthermore, at 3–7 min following rhythmic...
and isometric contractions, PvO2 values were generally not significantly different from their respective baselines. It was only at 3 min following isometric contractions, that PvO2 reached values significantly higher than baselines during 40% O2 and combined aspirin + 40% O2 conditions (Figure 4). Tables S1 and S2 provide the numerical data for these changes.

As expected, venous PCO2 (PvCO2), K+, and lactate were increased, whereas venous pH was decreased immediately following both rhythmic and isometric contractions in both young and older men (Tables S1 and S2). Thereafter, PvCO2 and K+ returned to baseline levels by 3-min post exercise, lactate was still raised at 7 min following both types of exercise, whereas pH tended to return to baseline more quickly in older, than young subjects. There were no treatment effects on the changes in PvCO2, K+, lactate, or pH. The only age-dependent effects were on lactate and pH, which showed greater changes from baseline in young men (Tables S1 and S2).
3.6 Relative effects of aspirin, 40% O₂ and aspirin/40% O₂ in young and older subjects

Considering just the six young men in whom PG efflux was assayed, aspirin, 40% O₂, and combined aspirin + 40% O₂ attenuated the postcontraction vasodilatation (increase in FVC) of rhythmic and isometric handgrip contractions by ~24 and ~30%, respectively, relative to control responses, whereas in the six older men, the attenuations were significantly smaller: ~17 and ~21% for rhythmic and isometric contractions, respectively (see Figure 5; p < .05 vs. young). Table S3 provides the numerical data for these percentage changes.

In contrast, as shown in Figure 5 and Table S3, aspirin caused similar, substantial reductions in young and older men of contraction-induced venous effluxes of 6-Keto PGF₁α, by 70%–85% and of PGEM, by 52%–75%. Furthermore, 40%
O₂ and combined aspirin + 40% O₂ caused similar reductions as aspirin for both types of contraction in both young and older men. The relative reductions in effluxes of PG metabolites were not significantly different between young and older men.

**FIGURE 3** Increases in venous efflux of 6-Keto PGF₁α and PGEM evoked in young men and older men by rhythmic (a) and isometric (b) contractions under control conditions (placebo), after Aspirin, during 40% O₂, and during combined Aspirin + 40% O₂. Absolute values are shown as mean ± SEM (n = 6 for each set of values) at baseline and immediately contractions ceased (time 0). *p < .05 versus respective baselines for all conditions included in bracket, †p < .05 for all treatments versus placebo.

**4 | DISCUSSION**

This is the first study to make direct comparisons between recreationally active young and older men of the contributions made by newly generated PGs to postexercise...
hyperemia for isometric, or rhythmic contractions. Taken together, our findings indicate that in both young and older men, PGs make substantial O₂-dependent contributions to postexercise hyperemia evoked by contractions of moderate intensity, whether they are isometric, or rhythmic.

4.1 Control responses

Maximum voluntary contraction was smaller in older than young men, consistent with a loss of muscle mass and relative increase in the proportion of oxidative fibers with aging (McGregor, Cameron-Smith, & Poppitt, 2014). Nevertheless, in both young and older men, the TTI was greater for isometric, than rhythmic contractions, but postcontraction hyperemia was similar for the two types of contraction. This may be attributed to a greater metabolic cost of rhythmic contractions (Newham, Jones, Turner, & McIntyre, 1995), and greater release of vasodilator metabolites. Importantly, compared at the same relative workload (60% MVC), postcontraction vasodilator responses were similar in young and older men. Others made similar observations in recreationally active young and older subjects (Jasperse, Seals, & Callister, 1994; Proctor et al., 2003), whereas in sedentary older subjects, exercise hyperemia was blunted (Poole et al., 2003).

Under control conditions, the concentrations of PGI₂ and PGE₂ metabolites in plasma were, as expected, in the low pg/ml range in young and older men (Heavey et al., 1985; Trappe & Liu, 2013). In young men, the increases in venous efflux of both PGs following rhythmic contractions were consistent with previous assays of venous blood and muscle interstitial fluid (Boushel et al., 2002; Karamouzis et al., 2001; Wilson & Kapoor, 1993). We now show that venous effluxes of PGI₂ and PGE₂ are also increased following isometric contraction. Moreover, we report the novel finding that venous effluxes of PGI₂ and PGE₂ are increased in older men following both rhythmic and isometric contraction at 60% MVC.

**FIGURE 4** Changes in \( P_O_2 \) evoked in young men and older men by rhythmic (a) and isometric (b) contractions under control conditions (placebo), after Aspirin, during 40% O₂ and during combined Aspirin + 40% O₂. Absolute values are shown as mean ± SEM at baseline, immediately contractions ceased (time 0) and at 3-, 5-, and 7-min postcontractions. *p < 0.05 versus respective baselines.
evidence that the same dose of aspirin inhibited bradykinin-induced generation of PGI₂ by 85% and ~70% at 30 and 90 min, respectively (Heavey et al., 1985). That PGE₂ efflux was also inhibited by 50%–75% accords with evidence that oral aspirin attenuates the increase in interstitial PGE₂ evoked by rhythmic contractions (Trappe & Liu, 2013).

By infusing COX inhibitor Wilson and Kapoor (Wilson & Kapoor, 1993) achieved almost complete inhibition of PGI₂ and PGE₂ efflux evoked by rhythmic forearm contractions at low and moderate intensity: they concluded PGs were responsible for 10%–20% of the hyperemia that occurred in the relaxation phases of rhythmic contractions. Even the incomplete COX inhibition we achieved indicated that in young men, PGs are responsible for 24%–32% of postcontraction hyperemia following rhythmic and isometric contractions at moderate intensity. This agrees with estimates made from the effects of COX inhibitors on postcontraction hyperemia of moderate-high intensity exercise (Cowley et al., 1985; Duffy et al., 1999; Kilbom & Wennmalm, 1976; Win & Marshall, 2005). But, our results also suggest PGs are responsible for at least 17%–21% of postcontraction hyperemia in recreationally active older men following rhythmic or isometric contractions at 60% MVC. Clearly, this contrasts with the proposal that the contribution of PGs to exercise hyperemia is lost with aging (Schrage et al., 2007).

However, our finding that the percentage reduction in postcontraction hyperemia caused by COX inhibition was smaller in older, than young men agrees with evidence that the action of PGs is blunted with aging (Nicholson et al., 2009).

### 4.3 Effects of 40% O₂

In both young and older men, 40% O₂ caused similar attenuation as aspirin of both PGI₂ and PGE₂ efflux, and the
postcontraction hyperemia associated with rhythmic and isometric contractions. Moreover, 40% O₂ and aspirin applied together had no greater effect on PG efflux, or the vasodilator responses. We previously showed in young men, that when breathing 40% O₂ was restricted to the period of isometric contraction, it attenuated postcontraction hyperemia, but when given immediately after contraction, it had no such effect (Fordy & Marshall, 2012). Thus, we can now argue it is the release, rather than the action of PGI₂ and PGE₂ that allows PGs to make an O₂-dependent contribution to postcontraction hyperemia during both types of contraction, even though FBF and O₂ delivery are more restricted during isometric contractions (Kagaya & Homma, 1997; Van Beekvelt et al., 2001). Importantly, our results indicate that PG release is just as O₂-dependent in recreationally active older men, as in young men.

Breathing 40% O₂ raises arterial PO₂ from ~90 to ~240 mmHg (Fordy & Marshall, 2012), and must have considerably steepened the O₂ gradients within muscle. To be specific, with air breathing, PvO₂ falls from ~40 mmHg at rest to 30–35 mmHg during submaximal exercise (Dufour et al., 2010), whereas intracellular PO₂ in muscle fibers falls from ~20–35 mmHg to ~3 mmHg during contractions at all intensities ≥50% maximum workload (Richardson, Newcomer, & Noyziewski, 2001). In this study, 40% O₂ had no effect on PvO₂ at rest, but alleviated the fall in PvO₂ at the end of both rhythmic and isometric contractions, as reported by others (Dufour et al., 2010). Hyperoxia also raises muscle intracellular PO₂ (Richardson, Noyziewski, Leigh, Wagner, 1998). Thus, at rest, additional O₂ must have diffused to the muscle fibers before reaching the veins, whereas during both types of contraction, O₂ delivery must have exceeded O₂ extraction such that the immediate postcontraction fall in PvO₂ was prevented. Accordingly, 40% O₂ must have raised the PO₂ gradient along the vascular pathway and from vasculature to muscle fibers. During recovery, PvO₂ increased above resting values as reported by others (Van Beekvelt et al., 2001) and 40% O₂ exaggerated this. Thus, O₂ delivery was greater than O₂ extraction and 40% O₂ exaggerated the disparity even though postcontraction hyperemia was attenuated by 40% O₂.

Presumably 40% O₂ increased arterial PO₂ to a similar extent in older, as in young men and affected the PO₂ gradients in a similar way. However, 40% O₂ did not affect the fall in PvO₂ in older men at the end of rhythmic, or isometric contractions. Moreover, PvO₂ did not rise above resting values during recovery from either type of contraction, whereas 40% O₂ raised PvO₂ only at the 3rd min following isometric contraction. Thus, even though 40% O₂ must have raised PO₂ at least part way along the vascular pathway toward muscle fibers, additional O₂ was extracted during both types of contractions and in recovery. This agrees with evidence that in recreationally active older men, O₂ extraction normally reaches maximum in submaximal exercise (Proctor et al., 2003), and suggests 40% O₂ can improve O₂ extraction not only during contraction but also in recovery, even though 40% O₂ attenuates postcontraction hyperemia.

### 4.4 Locations of PG release

PGI₂ is the major PG released by the endothelium, the expression of PGI₂ synthase being ~100-fold greater than PGE₂ synthase (Féletou, Huang, & Vanhoucke, 2011). On the other hand, PGE₂ is the dominant PG released by skeletal muscle fibers (McLennan & Macdonald, 1991; Trappe & Liu, 2013). Endothelial PGI₂ synthesis is maintained with aging (Féletou et al., 2011), whereas PGE₂ synthesis in muscle during exercise increases with age (Trappe & Liu, 2013), consistent with our finding of a trend for PGE₂ efflux following isometric contraction to be greater in older, than in young men. Thus, it is reasonable to conclude that in recreationally active young and older men, the PGI₂ and PGE₂ in venous efflux following isometric and rhythmic contractions originated mainly from endothelium and muscle fibers, respectively.

In resting muscle, perivascular PO₂ falls from ~50 mmHg around larger arterioles, to 28–30 mmHg around capillaries and postcapillary venules and ~33 mmHg around larger venules (Lash & Bohlen, 1987). During submaximal muscle contractions, periarteriolar PO₂ falls only transiently by ~10–20 mmHg, returning to resting values as the arterioles dilate, whereas pericapillary and perivenular PO₂ falls by ~50% with little recovery until contraction ceases (Lash & Bohlen, 1987). Thus, capillaries and postcapillary venules are the most likely sites for a fall in endothelial PO₂ to act as a stimulus for synthesis and release of PGI₂ into venous efflux and from their extraluminal surfaces into the interstitium (Hester & Hammer, 2002; Lash & Bohlen, 1987). This proposal is consistent with evidence that endothelial cells release ~10-fold more PGI₂ than PGE₂ at PO₂ levels comparable to those reached during contraction (Michiels et al., 1993) and with venules releasing PGs during muscle contraction which dilate arterioles (Hester & Hammer, 2002; McKay, Gardner, Boyd, & Hester, 1998). On the other hand, arterial occlusion of forearm, which profoundly reduces muscle intracellular PO₂ (Richardson et al., 2001) led to a threefold increase in efflux of PGE₂-like substance (Kilbom & Wennmalm, 1976). Furthermore, contractions at ≥50% MVC reduce muscle intracellular PO₂ (Richardson et al., 2001) and cause PGE₂ release (Trappe & Liu, 2013), whereas 70% MVC contraction during arterial occlusion exacerbated PGE₂ efflux (Symons et al., 1991). Thus, it seems likely the fall in muscle intracellular PO₂ that occurs during contractions modulates, or triggers PGE₂ release from muscle fibers.

Set against this background, it seems reasonable to propose that in both young and older men 40% O₂ limits the fall in local PO₂ sufficiently to attenuate release of PGI₂.
from capillary and venular endothelium and PGE₂ from skeletal muscle fibers such that interstitial PG levels are reduced and postcontraction dilatation and hyperemia are attenuated.

4.5 Interdependent influences

ATP release from contracting muscle fibers is dependent on acidosis and lactic acid efflux (Tu, Lu, Cai, & Ballard, 2012), whereas ATP metabolism to adenosine is facilitated when PO₂ falls (Marshall, 2007). Since 40% O₂ did not affect lactate or H⁺ efflux in young or older men, it seems unlikely this ATP release was affected by contractions at 60% MVC. But, adenosine and ATP are also released from endothelial cells by a fall in PO₂ (Edmunds, Moncada, & Marshall, 2003; To, Kumar, & Marshall, 2015), whereas erythrocytes release ATP when hemoglobin off-loads O₂ (Ellsworth & Sprague, 2012) and both adenosine and ATP release PGI₂ from endothelial cells (Nyberg et al., 2013; Nyberg, Mortensen, Thaning, Saltin, & Hellsten, 2010; Ray, Abbas, Coney, & Marshall, 2002). Thus, 40% O₂ may have attenuated exercise hyperemia by reducing the release and actions of ATP and adenosine whose contributions are partly mediated by PGI₂ (Marshall & Ray, 2012; Nyberg et al., 2010, 2013).

4.6 Experimental considerations

We always tested rhythmic and isometric contractions in that order rather than randomizing, so as to avoid variability caused by fatigue from one type of contraction affecting the response to the other. An “order effect” seems unlikely to have influenced our results for young and older subjects were able to maintain rhythmic contractions at 60% MVC for 3 min and FBF had fully recovered before isometric handgrip. Assays of arterial PGs would have allowed more accurate assessment of PG efflux. However, forearm exercise at moderate intensity did not previously increase arterial PGI₂, or PGE₂ (Wilson & Kapoor, 1993). Thus, it is unlikely arterial assays would have changed our conclusions.

Intra-arterial infusion of COX inhibitor might have achieved more complete COX blockade than one oral dose (Wilson & Kapoor, 1993) and allowed better assessment of the PG contribution to postcontraction hyperemia. However, we avoided using O₂ at concentrations >40% to more rigorously test the O₂-dependency of PG release, or hyperemia because high O₂ concentrations generate reactive oxygen species, which attenuate endothelium-dependent dilatation (Rousseau, Tesslera, Henricson, & Sjöberg, 2010) and may blunt exercise hyperemia. Our evidence indicates 40% O₂ does not have this effect (Caruana & Marshall, 2015; Marshall, Junejo, D'Souza, & Ray, 2015).

Although we recorded postcontraction hyperemia following isometric contraction, we probably captured the majority of the hyperemia, for continuous Doppler recordings showed FBF increased less during isometric contraction at 60% MVC than 30% MVC, with a much larger postcontraction hyperemia (McNeil et al., 2015). Thus, the attenuating effects of 40% O₂ and COX inhibition on peak (postcontraction) hyperemia reported in this study probably reflected majority of the PG contribution. During rhythmic contractions, FBF averaged over contraction and relaxation phases increased more at 75% than 25% MVC; with a dramatic increase in postexercise FBF only at 75 MVC (Van Beekvelt et al., 2001). But, the peak postexercise FBF equaled FBF recorded in the relaxation phases between rhythmic contractions (Van Beekvelt et al., 2001). Thus, it is likely the FBF we recorded immediately postexercise for rhythmic contractions at 60% MVC was comparable to FBF during the relaxation phases. Since PG efflux increased on cessation of rhythmic contractions at 60% MVC, it seems reasonable to propose PGs contributed to the increases in FBF during the contraction period as reported by others for moderate rhythmic contractions (Wilson & Kapoor, 1993). Continuous recordings of FBF during rhythmic contractions at 60% MVC will be required to assess the time course and magnitude of this PG contribution.

4.7 Perspectives

Postcontraction hyperemia not only allows wash out of vasodilator mediators generated during contraction, but it restores creatine phosphate (PCr) at a rate determined by O₂ delivery, O₂ diffusion, and capacity for oxidative phosphorylation (Haseler, Lin, & Richardson, 2004; Kemps Harel, Prompers Jeanine, & Wessels, 2009; Layec, Haseler, & Richardson, 2013). Thus, blunted postcontraction hyperemia in aging and cardiovascular disease contribute to slower PCr recovery rates and may limit the ability to perform repetitive daily activities (Haseler et al., 2004; Kemps Harel et al., 2009; Layec et al., 2013). The present findings indicate that use of proprietary COX inhibitors might similarly slow PCr recovery kinetics and hasten fatigue, particularly in older people. On the other hand, breathing 40% O₂ during exercise may avoid the deleterious effects of COX inhibition while facilitating the benefits. For, raised PO₂ in muscle during exercise would inhibit PGI₂ and PGE₂ synthesis, but allow increased O₂ extraction during recovery at least in older men (Figure 3) despite the attenuated postcontraction hyperemia. Certainly, even in young men, 40% O₂ given selectively during recovery from a maximal fatiguing forearm contraction improved performance in a second contraction undertaken a few minutes later (Fordy & Marshall, 2012).
4.8 Conclusions

Against a background of controversy over whether PGs are necessary for full expression of exercise hyperemia, we have shown that isometric and rhythmic contractions at moderate intensity (60% MVC) caused substantial release of PGI2 and PGE2 from forearm of young and older men and that COX inhibition attenuated postcontraction hyperemia for both types of contraction by at least 20% in both young and older men. Furthermore, in young and older men, the release of PGI2 and PGE2 and contribution to postcontraction hyperemia were similarly attenuated by breathing 40% O2, whereas combined 40% O2 and COX inhibition had no greater effect. Thus, we propose the release of PGs during moderate intensity (60% MVC) isometric and rhythmic contractions is O2-dependent in both young and older men.

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CONFLICT OF INTERESTS

The authors have no competing interests to declare.

AUTHOR CONTRIBUTIONS

JMM conceptualized the study; RTJ, CJR, and JMM designed the work; RTJ performed acquisition and data analysis; RTJ, CJR, and JMM interpreted the data; RTJ drafted the manuscript, which was critically revised by CJR and JMM. All authors approved the final manuscript.

ORCID

Rehan T. Junejo https://orcid.org/0000-0002-0670-8339
Clare J. Ray https://orcid.org/0000-0001-7410-1013
Janice M. Marshall https://orcid.org/0000-0003-3609-5884

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Additional supporting information may be found online in the Supporting Information section.

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