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(Article begins on next page)

Increased tissue oxygenation explains the attenuation of hyperaemia upon repetitive 1 pneumatic compression of the lower leg 2 3 Alessandro Messere ¹, Gianluca Ceravolo², Walter Franco², Daniela Maffiodo², Carlo Ferraresi², Silvestro 4 5 Roatta¹ 6 **Author contribution** 7 AM: conception and design of the experiment, collection, analysis and interpretation of the data, drafting the 8 9 manuscript 10 GC: collection, analysis and interpretation of the data, drafting of the manuscript WF: design of the experimental set-up, collection, analysis and interpretation of the data 11 DM: design of the experimental set-up, collection, analysis and interpretation of the data 12 13 CF: design of the experiment, critical revision of the manuscript SR: conception and design of the experiment and critical revision of the manuscript 14 All authors approved the final version of the manuscript. 15 16 ¹ Dept. Neuroscience, University of Torino, Torino, Italy 17 ² Dept. of Mechanical and Aerospace Engineering, Politecnico di Torino, Torino, Italy 18 19 Running Head Tissue oxygenation modulates compression-induced hyperaemia 20 21 Corresponding author 22 Silvestro Roatta 23 Dip. Neuroscienze, Università di Torino 24 C.so Raffaello 30, 10125 Torino, Italy 25 Email: silvestro.roatta@uniito.it

ABSTRACT

27 **Aim**

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- 28 The rapid hyperaemia evoked by muscle compression is short-lived and was recently shown to undergo a
- 29 rapid decrease even in spite of continuing mechanical stimulation. The present study aims at investigating
- 30 the mechanisms underlying this attenuation which include local metabolic mechanisms, desensitization of
- 31 mechano-sensitive pathways, and reduced efficacy of the muscle pump.
- 32 Methods
- 33 In 10 healthy subjects short sequences of mechanical compressions (n=3-6; 150 mmHg) of the lower leg
- were delivered at different inter-stimulus intervals (ranging from 20 to 160 s) through a customized
- 35 pneumatic device. Hemodynamic monitoring included near infrared spectroscopy, detecting tissue
- 36 oxygenation and blood volume in calf muscles, as well as simultaneous echo-Doppler measurement of
- 37 arterial (superficial femoral artery) and venous (femoral vein) blood flow.
- 38 **Results**
- 39 The results indicate that: i) a long lasting (>100 s) increase in local tissue oxygenation follows the
- 40 compression-induced hyperaemia; ii) the compression-induced hyperaemia exhibits different patterns of
- 41 attenuation depending on the inter-stimulus interval; iii) the amplitude of the hyperaemia is not correlated
- 42 with the amount of blood volume displaced by the compression; iv) the extent of attenuation negatively
- correlates with tissue oxygenation (r=-0,78, P<0.05).
- 44 Conclusion

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- 45 Increased tissue oxygenation appears to be the key factor for the attenuation of hyperaemia upon repetitive
- 46 compressive stimulation. Tissue oxygenation monitoring is suggested as a useful integration in medical
- 47 treatments aimed at improving local circulation by repetitive tissue compression.

NEW AND NOTEWORTHY

- 50 This study shows that i) the hyperaemia induced by muscle compression produces a long-lasting increase in
- 51 tissue oxygenation; ii) the hyperaemia produced by subsequent muscle compressions exhibits different
- 52 pattern of attenuation at different inter-stimulus intervals; iii) the extent of attenuation of the compression-
- 53 induced hyperaemia is proportional to the level of oxygenation achieved in the tissue. The results support the
- 54 concept that tissue oxygenation is a key variable in blood flow regulation.
- 56 **Keywords**: muscle blood flow, hyperaemia, muscle compression, tissue oxygenation.

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- 59 IPC intermittent pneumatic compression
- 60 ISI inter-stimulus interval
- 61 NIRS near-infrared spectroscopy
- 62 SRS spatially-resolved spectroscopy
- 63 THI total haemoglobin index
- TOI tissue oxygenation index

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INTRODUCTION

- 68 Since the seminal work of Mohrman and Sparks (39) several studies have demonstrated that a rapid and
- transient hyperaemic response can be elicited by a short-lasting muscle compression (10, 30, 38, 56-60).
- Although the underlying mechanisms have not been fully identified, this phenomenon has been well
- documented in different experimental models, such as the isolated muscle (39), awake and anesthetized
- animals (57, 58, 60) and humans (10, 30, 38, 56). In addition a rapid dilatory response to compressive
- stimuli has also been observed in isolated feed arteries (7). More controversial is the hemodynamic response
- 74 to repeated compressive stimuli. Kirby et al (30) observed that the response to 5 consecutive compressions
- 75 was non-significantly attenuated with respect to the response to a single compression. Conversely, Clifford et
- al (7) using the same pattern of 5 consecutive compressive stimuli on an isolated muscle feed arteries
- 77 observed a significant increase of the dilatory response as compared to the single compression.
- 78 In a recent work Turturici and colleagues investigated the blood flow response to a longer lasting sequence
- 79 of mechanical stimulations (20 compressions, 1 s ON /1 s OFF) reporting that the initial hyperaemic
- 80 response progressively fades away in spite of continuing stimulation and hypothesized that the
- 81 mechanosensitive mechanism underlying the response could undergo some kind of transient inactivation
- 82 (60). In fact, the attenuation of the compression-induced hyperaemia was observed to increase at increasing
- 83 stimulation frequencies (60). A similar behavior was recently observed also in humans (38).
- 84 Surprisingly this phenomenon has been poorly described in the several investigations concerning the
- 85 hyperaemic effect of intermittent pneumatic compressions (IPC) (14, 15, 32-34), and in experimental studies
- investigating the mechanisms underlying compression and contraction-induced hyperaemia (9, 24, 31, 40,
- 87 44), with the exception of a short report by Tschakowsky et al (56). In this pioneering investigation the
- 88 authors observed that repetitive compression of the forearm below heart level exhibited a transient
- 89 hyperaemia settling to a lower level after 10-20 s from the beginning of the treatment (56). More recently
- 90 Sheldon et al (47) also reported attenuation of the hyperaemia during IPC treatment, although on a larger

91 time scale (45 vs. 5 min from the beginning) and observed that the effect was dependent on the frequency of 92 stimulation. The issue is relevant because improving limb perfusion is a major aim in the treatment of disorders such as 93 94 the peripheral arterial disease and is pursued in sport medicine for accelerated recovering from fatigue (1, 95 35). Understanding of the underlying mechanisms is essential for implementing optimal treatments (46). 96 Potential mechanisms underlying attenuation of the hyperaemia during repetitive mechanical stimulation 97 include: 1) inactivation of the mechano-sensitive vasodilatory pathways (60), 2) diminished efficacy of the muscle pump (56), and 3) local regulatory mechanisms that may be activated in response to hyper-perfusion 98 99 (30, 56). Unfortunately, none of these possibilities is supported by a solid experimental evidence. In 100 particular, 1) mechano-sensitive channels exhibiting inactivation properties have been identified (17, 26), but 101 their actual involvement in the rapid compression-induced dilatation was not ascertained, 2) at high stimulation frequencies incomplete vascular refilling may reduce the contribution of the pump, however, a 102 role for the muscle pump was excluded in a previous animal study (60), and 3) local vasoconstrictory 103 104 mechanisms are known to act in response to hyper-perfusion but little is known about the actual regulatory variable (O₂, CO₂, pH, etc.) and about the strength and timing of this vascular reaction (6, 45). However, in a 105 recent reformulation of the metabolic control of blood flow, a primary role for tissue pO₂ has been postulated 106 (23). According to their model, an excessive rise in O₂ concentration within the tissue would trigger a 107 108 vasoconstrictory response, mediated by the inhibition of a tonically released vasodilator (23). Along this line, a rise in tissue O2 occurring during a compression-induced hyperaemia could then trigger a constrictor 109 response and limit further hyperaemic events in response to subsequent mechanical stimuli. 110 111 On this basis the present study was aimed to test the following hypotheses: 1) the compression-induced hyperaemia elicits a rise in tissue oxygenation, 2) the attenuation of the hyperaemic response to subsequent 112 113 compressive stimuli is related to the extent of hyper-oxygenation achieved in the tissue, and 3) the other 114 mechanisms, namely, the intrinsic inactivation of mechano-sensitive pathways and the muscle pump would 115 have a minor role in the attenuation of the hyperaemic response upon repetitive compressive stimulation. In order to assess changes in tissue oxygenation, the near infrared spectroscopy (NIRS) was adopted. By 116 117 locating the NIRS probe under the compressive cuff, continuous monitoring of local oxygenation and blood 118 volume changes from the relevant muscles was be achieved. Moreover, in addition to arterial inflow, venous outflow was also monitored as its response to the compression is an indicator of the extent of filling of the 119 120 venous compartments and thus, of the efficacy of the muscle pump exerted by compressive stimuli.

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MATERIALS AND METHODS

- 123 Ethical approval
- Ten healthy subjects (8 men and 2 women; age: 27.1 ± 3.0 years; weight: 67.9 ± 11.7 kg; height: 176.7 ± 9.7
- cm) were recruited for the present study. All subjects were normotensive and non-obese.
- The study conformed to the standards set by the Declaration of Helsinki and was approved by the Local

Ethical Committee (Prot. # 60195) and all subjects gave their written informed consent after they were 127 128 instructed about purpose and procedures of the experiment. 129 130 Mechanical leg compressions A previously tested prototype of IPC device was employed in the present study to deliver controlled and 131 repeatable compressions to the leg of the subject (19, 20). Briefly the device exerts a compressive action by 132 133 inflating five different bladders wrapped around the foot and the calf of the subject, with programmable 134 pressure levels and timing. In the present study all bladders were inflated simultaneously to a supra-systolic 135 pressure of 150 mmHg, with inflation and deflation times of about 3 s each. Two digital pulses are generated by the device to signal the starting time of both inflation and deflation. 136 137 138 *Near-infrared spectroscopy* Local hemodynamic changes induced by leg compression were measured using a continuous wave NIRS 139 140 device (NIRO-200NX, Hamamatsu Photonics, Hamamatsu City, Japan), which, besides the classical modified-Lambert-Beer method, supports spatially-resolved spectroscopy (SRS) (16, 52). Since mechano-141 sensitive vascular reactivity appears to be more prominently expressed by muscular than cutaneous tissues 142 (57) we focused our attention on SRS parameters which, being less affected by cutaneous circulation, 143 provide a more specific monitoring of muscle tissue (2, 36, 37). Since NIRS cannot discriminate between 144 haemoglobin (Hb) and myoglobin (Mb), all measurements always refer to Hb+Mb in the sample volume 145 146 (51). In particular, TOI (tissue oxygenation index) indicates the ratio (MbO₂+HbO₂)/(Mbtot+Hbtot) expressed in percentage, and THI (tissue haemoglobin index) indicates the concentration of (Hb+Mb) in 147 148 arbitrary units and is therefore an indicator of blood volume changes. Classical Lambert-Beer Parameters 149 (O₂Hb and HHb detecting changes in the concentration of oxygenated and deoxygenated (Hb+Mb), respectively) are only displayed in Fig. 1 and not further considered in the study. 150 151 152 Hemodynamic measurements 153 Measurements of blood velocity in femoral artery and vein were performed simultaneously using two 154 ultrasound systems (Mylab 25 XVision and MyLab 25 Gold, Esaote, Genoa, Italy) equipped with linear 155 arrays (LA 523, Esaote, Genoa, Italy). Superficial femoral artery and femoral vein were insonated distally to 156 the inguinal ligament. Since these instruments could not measure blood velocity and vessel diameter 157 simultaneously, the latter was measured at the beginning and at the end of every stimulation protocol. 158 Doppler measurements were performed by extending the sample volume over the whole vessel size, 159 echographically displayed (transversal approach) in real time. All blood velocity measurements in femoral artery were obtained with insonation angle of about 60° (operating frequency of 6.6 MHz) instead, a higher 160 angle of about 70° (operating frequency of 5 MHz) was used in order to avoid saturation of the recording 161 162 when assessing the high-speed venous outflow propelled by leg compression. The two probes were placed

163164165	few centimeters apart with the ultrasound beam of the proximal probe oriented proximally and the one of the distal probe oriented distally, in order to avoid interference between the measurements.
166	Experimental set-up
167 168 169	A schematic representation of the experimental setup is reported in Fig 1 A. All experiments were performed in a quiet room with a constant ambient temperature of about 22-23 °C. The subject sat upright on an adjustable chair with the back supported by a back rest.
170 171 172 173	The NIRS probe was located on the lateral head of gastrocnemius muscle of the right leg (inter-optode distance = 4 cm). The IPC device was wrapped around the lower leg, over the NIRS probe. The two echographic probes were maintained in place by dedicated holders for the whole duration of the protocol.
174	Experimental protocol
175 176 177 178 179 180 181	After 15 min of rest, an initial series of 3 compressive stimuli with inter-stimulus interval (ISI) of 160 s was delivered to the subject. After other 4 min of rest four series of 6 compressive stimuli were delivered at different frequency (ISI= 20, 40, 60 and 80 s) in randomized order and separated by 4-min resting intervals. Femoral artery and femoral vein diameters were collected at the beginning and at the end of every stimulation protocol. Diameters were measured along a single direction, since both vessels present a circular cross-section in these experimental conditions. Average diameter of the artery was calculated as (Ds+2*Dd)/3, Ds being the systolic and Dd the diastolic diameter.
183	Data acquisition and processing
184 185 186 187	The NIRS signals were digitally acquired along with both Doppler audio signals and the digital synchronism signal from the IPC device by a single acquisition system (CED Micro 1041, Cambridge Electronic Design, Cambridge, UK) and stored on the computer for later analysis with Spike2 software (version 6.10, Cambridge Electronic Design, UK).
188	A specific algorithm was implemented in the Spike2 script language to calculate blood velocity from
189 190	Doppler audio signals (11, 25). Briefly, power spectra of the audio signals were computed by the Fast Fourier Transform over non-overlapping epochs lasting 25.6 ms. From each spectrum the maximum
191	frequency of the signal (corresponding to maximum blood velocity) was estimated according to D'Alessio
192	(11), then the mean frequency was calculated as the average of all frequencies below the maximum,
193	weighted according to spectral amplitude (25). The mean frequency was then time-averaged over each
194	cardiac cycle and converted into blood velocity, $BV = (MF * C)/(2F * \cos\theta)$, where MF is the mean
195	frequency calculated from Doppler shift, C the averaging speed of ultrasound in soft tissue (1540 m/s), F the
196	operating frequency of the Doppler, and ϑ the insonation angle). Blood flow, in ml/min, was then calculated

197	as mean blood velocity times cross-sectional area of the vessel (BF = BV * πr^2 *60, where BV is the blood
198	velocity expressed in cm/s, and πr^2 the cross sectional area of the vessel in cm ²).
199	The response to each compressive stimulus was characterized by: pre-compression arterial blood flow,
200	calculated as the average over the 4 s preceding the compression; pre-compression TOI; pre-compression
201	THI; peak arterial blood flow, as the hyperaemic peak reached after the compression; Δ TOI, calculated as
202	the difference between the peak TOI reached after the compression and pre-compression TOI; displaced
203	blood volume, calculated as the product of the area under the curve of the venous blood velocity response
204	and the cross-sectional area of vein.
205	In addition, the amplitude of the hyperaemic response was also calculated as the difference between peak
206	arterial flow and pre-compression flow.
207	In order to assess the extent of attenuation of the response throughout the experimental protocol, changes in
208	blood flow and blood volume were normalized to the changes observed in response to the first delivered
209	compressive stimulus.
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211	Statistics
212	To examine the effect of repetitive leg compression performed at different ISI on peak blood flow, displaced
213	blood volume, pre-compression THI and pre-compression TOI, a two-way repeated-measures ANOVA was
214	used with factors ISI and repetition (GraphPad Prism v 6.0, GraphPad Software, La Jolla, CA). When
215	significance was found, a Dunnett's post hoc test was performed to assess significant changes within each
216	series with respect to the response to the first stimulus. Pearson's coefficient was used to assess the
217	correlation between different variables. All data are expressed as means \pm standard deviation in the text and
218	means \pm standard error in diagrams. The level of statistical significance was set at P< 0.05.
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221	RESULTS
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223	Single leg compression
224	A typical response to a single compressive stimulus is reported in figure 1B. Venous blood velocity exhibits
225	a prompt and short-lasting increase, peaking $1.7 \pm 0.2~\text{s}$ after the beginning and terminating before the end of
226	the compression. The blood volume displaced by compression was on average 28.3 ± 14.8 ml. The increase
227	in arterial blood flow starts immediately after deflation and peaks in $4.9 \pm 1.4~s$ passing from a basal value of
228	74.5 ± 22.7 ml min-1 to 260.2 ± 83.3 ml min-1 during the peak (peak flow is 3.6 ± 1.0 of baseline). Blood
229	flow generally returns within 15-25 s The response in tissue oxygenation is further delayed. TOI slowly
230	increases (from 66.4 \pm 5.1 to 78.0 \pm 4.0 %) and peaks after 20.6 \pm 5.1 s from deflation. Local changes in
231	blood volume are detected by THI exhibiting a rapid decrease during compression followed by a slower
232	return to the basal level, in agreement with the changes in venous and arterial blood flow, respectively.

234 Repeated leg compressions 235 The hemodynamic response to repetitive leg compression at different ISI is summarized in Fig 2, each 236 column representing the response to a single stimulus. The upper two rows show the response in terms of peak arterial blood flow and displaced venous blood volume, both variables exhibiting a significant 237 dependence on ISI (p<0.01) and repetition (p<0.01). It can be observed that when ISI = 160 s the response to 238 subsequent stimuli is unchanged. Unchanged response in terms of peak arterial flow and displaced blood 239 volume is also observed in response to the first compression in each series. Instead, both parameters exhibit a 240 progressive attenuation although with different time course at ISI ranging from 20 to 80 s. In particular, the 241 hyperaemia is consistently reduced starting from the second stimulus in the series, at ISI ranging from 20 to 242 60 s, while displaced blood volume is consistently reduced at ISI = 20 and 40 s, starting from the third 243 stimulus. A peculiar pattern is observed at ISI = 80 s where hyperaemia is only attenuated in response to 244 245 even and not to odd stimuli, while, at the same time displaced blood volume remains unaffected. NIRS parameters, shown in the lower rows of fig 2, exhibited a significant dependence on repetition 246 247 (p<0.01) but not on ISI, along with a significant interaction between the two factors. It can be observed that pre-compression THI, which can be considered an indicator of vascular filling, qualitatively parallels the 248 249 changes in displaced blood volume, remaining unchanged at large ISI and exhibiting the most marked reduction at ISI = 20 s. Pre-compression TOI exhibits instead marked increases at all ISIs lower than 160 s 250 starting from the second stimulus in the sequence. It is interesting to observe that its pattern of change is 251 252 opposite to peak blood flow: i.e., hyperaemic peak is higher if the pre-compression TOI is lower. Note also 253 that the oscillating pattern previously observed in peak blood flow at ISI = 80 s is also exhibited by precompression TOI in an opposite way. 254 In order to provide a better understanding of the interplay between the different parameters in the peculiar 255 response to repetitive compression at ISI = 80 s, original tracings are reported from a representative subject 256 257 in Fig. 3. As described in Fig. 1, the first stimulus elicits a marked hyperaemia which results in a marked increase in oxygenation. The following compression, which occurs when the tissue oxygenation is still high, 258 259 now elicits a much smaller hyperaemia, resulting in a proportionally smaller increase in TOI and attenuated vascular refilling in THI. The third compression occurs when the TOI is almost returned to basal levels and 260 the elicited hyperaemia resumes its original size. Although it cannot be fully appreciated with this time scale, 261 262 the venous blood flow response is comparable in all instances as well as the pre-compression level reached 263 by THI. Another representative recording illustrating the pattern at ISI = 20 s is reported in fig 4. Note the 264 265 disappearance of the hyperaemic response to the second and subsequent stimuli in spite of the fact that arterial blood flow is returned to basal level. A weak hyperaemia reappears only in response to the last 266 267 stimulus, when also TOI is almost returned to basal level. Note that THI indicates that blood volume is

almost fully returned to basal level after the first stimulus (thanks to the marked hyperaemia) but not 268 afterwards. Accordingly, the venous response is markedly reduced after the third and subsequent stimuli. 269 In general a good correlation was found between the peak blood flow during hyperaemia and the ensuing 270 increase in oxygenation as shown in fig 5 A, in which all subjects have been pooled and each dot represents 271 the response to a single compression. The overall r is 0.76 (p< 0.05). When individually computed for the 272 273 different subjects r ranged between 0.72 and 0.95 (p< 0.05) (average 0.78 ± 0.1). 274 On the contrary the hyperaemic response was not correlated with the amount of displaced blood volume as shown in Fig. 5B (r = 0.34, individual r ranging between -0.4 and + 0.3). 275 276 Fig 5C shows the correlation between pre-compression TOI and the peak of the hyperaemic response which is exhibiting an overall r = -0.434 (p<0.05), however a much higher within- subject correlation is observed: -277 0.78 ± 0.06 , individual r ranging between 0.7 and 0.9 (p<0.05). 278 In Fig. 5D the amplitude of the hyperaemic response (= peak flow-basal flow) instead of peak flow is plot vs. 279 pre-compression TOI. While the general pictures resembles that of Fig. 5C, it is here better evidenced that 280 281 the hyperaemia can be almost abolished at high TOI levels. Moreover, the slope of the regression lines, m, 282 allows to quantify the dependence of the hyperaemic response on tissue oxygenation. On average, m = -0.082 ± 0026 meaning that the compression-induced hyperaemia is attenuated by 8% per unitary increase of 283 284 TOI, with respect to its full amplitude (the one that is evoked in resting conditions). 285 Changes in vessel size 286 287 A slight increase in vessel diameter was detected from the comparison of measurements performed at the beginning and at completion of the experimental protocol in both femoral artery (from 6.0 ± 0.8 to 6.2 ± 0.8 288 mm, p<0.05) and vein (from 8.3 ± 0.9 to 8.6 ± 1.3 mm, p<0.05) 289 290 291 **DISCUSSION** For the first time a comprehensive approach has been employed for the investigation of the rapid 292 293 compression-induced hyperaemia and its adaptation upon repetitive stimulation, which includes continuous assessment of NIRS indicators of changes in local tissue oxygenation and blood volume as well as 294 295 simultaneous monitoring of arterial inflow and venous outflow. This allowed us to describe the early 296 hyperaemic changes taking place at the beginning of IPC treatments at different frequencies, and to confirm 297 our initial hypotheses: i) the compression-induced hyperaemia elicits proportional increases in local tissue oxygenation; ii) the extent of attenuation of the hyperaemic response to subsequent stimuli is related to the 298

current level of tissue oxygenation; iii) the extent of attenuation is not strictly dependent on the extent of 300 vascular filling and on the ISI, therefore the attenuation cannot be attributed to the reduced efficacy of the 301 muscle pump or to a simple, time-dependent, inactivation mechanism of mechano-sensitive pathways.

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Compression-induced hyperaemia increases tissue oxygenation

A novel observation of the present study is that muscle compression elicits a prominent increase in local tissue oxygenation. This increase is consequent to the induced hyperaemia but is much longer lasting. This aspect is important because it reveals that the return to "control conditions" is not achieved at the end of the hyperaemia, which normally occurs within 15-25 s and may instead require up to 100 - 200 s. This pattern has never been reported for compression-induced hyperaemia but it is in agreement with what occurs in the rapid-onset hyperaemia induced by short contractions (53). It is generally accepted that an increase in perfusion, with unchanged metabolism, increases tissue oxygenation (3, 12). In the present condition, different factors could contribute to the observed TOI increase in response to compression-induced hyperaemia: 1) depletion of the venous-compartment, which alters the proportion of arterial/venous blood in the sample volume; 2) increased Hb saturation in venous blood due to decreased oxygen extraction, given that the hyperaemia occurs in a condition of constant metabolism; 3) increased saturation of myoglobin. The voiding of venous compartment does not seem to affect the TOI signal considerably, as no relevant changes are observed immediately after the compression, including those associated with large blood volume changes (see original tracings in Figs. 1, 3 and 4). Unfortunately, NIRS cannot discriminate between Mb and Hb saturation nor between arterial and venous compartments, thus no univocal explanation can be provided. Irrespective of the underlying reason, the increase in tissue oxygenation was a very consistent feature of the hemodynamic response to the compression of the resting

Is compression-induced hyperaemia attenuated by increased tissue oxygenation?

muscle and exhibited a good correlation with the amplitude of hyperaemia (Fig. 5A).

Several lines of evidence from the present study support the finding that elevated tissue oxygenation is the factor responsible for the attenuation of the hyperaemia and for the reduced responsiveness to the mechanical stimulus. By looking at the original tracings of Fig. 3 it can be observed that the response to the second compression is smaller as compared to the first and the third responses, while TOI is higher than baseline. The same is visible in Fig.4: the hyperaemic response almost disappears during the initial high oxygenation phase and only later exhibits a tendency to recover, concomitantly with a decrease in TOI. This dependence of peak hyperaemia on pre-compression TOI is also supported by the histograms of Fig. 2 (see opposite patterns of peak blood flow and pre-compression TOI) and is quantitatively assessed by the correlations in Fig. 5 C and D. Moreover, it appears to be rather linear and rather similar between different subjects. According to these indications, the amplitude of the hyperaemic response is attenuated by 8 ± 2 % per unitary increase of TOI meaning that an increase in TOI by 12.5 points virtually abolishes the response.

Notably, the dependence of the active vessel dilatation on tissue oxygenation may explain why the same 335 short sequence of compressive stimuli elicited opposite effects in vitro (7), where tissue hyperoxia does not 336 take place, and in vivo (30). 337 338 In the several studies investigating hemodynamic effects during IPC treatments this pattern of adaptation of 339 the hyperaemia has not been described, possibly because the attention was focused on medium-long term rather than on early effects. Although different devices and patterns of stimulation have been used in 340 previous investigations, an increase in limb perfusion is generally reported, ranging between 20 and 240 %, 341 and being assessed at 5-60 min from the beginning of the treatment (9, 15, 24, 33, 40, 44, 47), which also 342

appear to be little dependent on the stimulation frequency (47). These results are not readily comparable with

the present ones because no steady state was reached in our study. It is reasonable to expected that a certain

stable increase in perfusion is obtained with prolonged stimulation, once steady tissue oxygenation is

achieved.

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348 *Underlying mechanisms and implications*

As discussed above, the attenuation of the mechano-sensitive dilatory response to multiple compressions could result as a reaction of the tissue to the hyper-perfusion (generated in response to the first stimulus), which entails the washout of metabolites and alteration of the local milieu in which PO2 is a most relevant variable (4, 23, 27). It is well known that low oxygenation stimulates vasodilatation and, conversely, that increased oxygenation leads to vasoconstriction, although the effects generally observed in humans exposed to increased levels of inspired PO₂ are rather small (5, 62). In the latter study, increasing arterial PO₂ from 100 to 2100 mmHg increased resting vascular conductance only by 20-25% and reduced functional hyperaemia by 20% (5). However it must be observed that tissue PO₂ is differently affected by increased arterial PO₂ and hyper-perfusion. In fact while the hyperbaric hypoxia at 2100 mmHg increases the amount of oxygen carried to the tissue by about 30% (5) a 2-fold increase in perfusion results in a 200% increase in oxygen flow. In early studies reactions to hyper-perfusion were investigated on isolated preparations with externally-controlled blood supply (21). However these studies could not provide a clear indication of the time course of the local tissue response, nor could they discriminate between "metabolic" and myogenic response, given that hyper-perfusion was produced by increased perfusion pressure which also resulted in increased transmural pressure (48). In this respect, the compression-induced hyperaemia offers a peculiar model of (transient) tissue hyper-perfusion, characterized by unchanged tissue metabolism, unchanged arterial PO₂ and most likely unchanged neuro-hormonal drive.

The prompt counter-reaction to the compression-induced hyperaemia and the concomitant inactivation of the mechano-sensitive dilatation upon increased tissue oxygenation fits with the "bang-bang" model of blood flow control, recently proposed by Golub & Pittman (23) according to which the feedback signal (O_2 , whose concentration increases in response to increased O_2 availability) carries the information of excessive

perfusion and operates a vasoconstriction by inactivating the tonically released vasodilators (namely, nitric oxide), aim of this regulation being to protect the tissue from hyperoxia and prevent excessive perfusion. Accordingly, the vascular mechano-sensitivity, which is considered to mediate the rapid dilatation and the anticipatory (feed-forward) hyperaemia at the beginning of exercise (8, 30, 43, 60) is promptly abolished if the exercise does not take place, due to the hyper-oxygenation produced by the hyper-perfusion. Instead, in the case of exercise the hyper-oxygenation is quickly reduced even below control levels (18) by increased metabolism and no limitation to vasodilation takes place, which results in the "functional hyperaemia". The same mechanism is likely to explain why both passive movement hyperaemia is attenuated upon repeated stimulation (54, 55) and contraction-induced hyperaemia is attenuated after a sequence of muscle compressions (38). Surprisingly, with one exception (34) no study has ever included NIRS in the characterization of the hyperaemic response to compression and IPC. Although tissue oxygenation can be considered a major outcome of perfusion, in the short term it does not strictly follow arterial blood flow, e.g., in Fig. 2 TOI is maintained at high levels for some time, after the end of hyperaemia. On this basis, it might be more appropriate to monitor TOI rather than blood flow in order to better appreciate the actual effects of the treatment. In addition, adopting NIRS as the monitoring technique gives the possibility to assess the effects specifically on the tissue of interest, as compared to the more global information provided by blood flow in

Alternative hypothesis 1: Vascular refilling and the muscle pump

 an large supplying artery.

The parallelism observed between changes in pre-compression THI and in displaced blood volume (Fig.2), suggests that pre-compression THI is a good indicator of current vascular filling and that its changes mostly reflect volume changes of the venous compartment. By observing its time course after the compressive stimulus we can detect a fast refilling phase, associated to the possible concomitant hyperaemia, and a subsequent slow phase, associated to "resting" blood flow. At high ISI, i.e., 80 and 160 s, a complete vascular refill is granted by both a consistent hyperaemia and a large time interval. Accordingly, the compressive action of the device displaces comparable amount of blood volume at every stimulus. At lower ISI, the lack of hyperaemia and/or insufficient time for the slow phase to yield a significant contribution may result in incomplete vascular refilling and in a reduction of the blood volume displaced by the subsequent compression. This observation is in agreement with the study by Delis and colleagues who reported 3 to 4 compressions per minute (i.e., ISI = 20 or 15 s) as the optimum stimulation frequency to maintain low venous pressure in the treated limb (13). Valic et al (61), in the anesthetized dog estimated a refilling time of less than 1 s due to the large contraction-induced hyperaemia. Based on direct foot venous pressure estimation, two human studies reported refilling times of 16 - 40 s after 10 tip—toe movements (42) and pneumatic compression (22). In the present conditions the refill could take place in 10-15 s through the rapid

phase in the presence of large hyperaemia but could otherwise require more than one minute when hyperaemia was blunted (Fig. 3).

According to the "muscle pump" effect, an increase in intramuscular pressure empties the venous

compartments producing a decrease in venous pressure, which in turn increases the artero-venous pressure gradient thus contributing to the ensuing hyperaemia. This mechanism is activated both with active muscle contraction as well as with the compression of the passive muscle and has been often considered to explain the larger hyperaemic responses observed when compressing (10, 56) or contracting (41, 50) limbs muscles below as compared to above heart level. However the issue is still debated (7, 29, 49) due to the conflicting evidence provided by other studies (24, 28, 61). In particular, Jasperse et al. investigated the effect of positional differences on reactive hyperaemia, as a model of hyperaemia dissociated from the muscle pump. They showed that also reactive hyperaemia is larger when evoked below, with respect to above heart level, suggesting that positional effects may be secondary to differences in driving pressure rather than to the muscle pump. The present results support this view through a complementary model, i.e., the muscle pump action dissociated from the hyperaemia. This particular condition was observed in several instances such as the responses to the second compression at ISI ranging from 20 to 80 s (in Fig 2 and in Fig 4), in which maintained vascular filling and compression-displaced blood volume, i.e., an effective muscle pump, was associated with a considerably reduced hyperaemia, as compared to the first compression in the series. This proves that the muscle pump mechanism is not involved in the attenuation of the hyperaemia in response to multiple compressions. Whether the muscle pump plays a role in the hyperaemic response to the first compressive stimulus cannot be ruled out based on the present data. In fact, from scatter plot in Fig 5B we can observe that the largest hyperaemic responses were never associated with low displaced blood volume, which suggests that adequate vascular filling may be a necessary condition to express the full response. Investigating the mechanisms behind compression-induced hyperaemia was not an aim of this study; further investigations will be necessary to elucidate this issue.

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Alternative hypothesis 2: Desensitization of mechano-sensitive pathways

It was previously observed that the hyperaemic response to the compressive stimulus progressively reduced to 26% of its original amplitude, with decreasing ISI from 4 min to 2 s (60). On this basis the hypothesis was put forward that the attenuation could be due to some transient inactivation (desensitization) of mechanosensitive dilatory mechanisms. This hypothesis was supported by the observation that desensitization upon repeated activation is a characteristic of certain vascular mechano-sensitive channels (17, 26). A subsequent human study in which similar stimulation protocols were applied to the forearm, qualitatively confirmed the attenuation pattern, although with a less gradual dependence on the ISI (38)

The up-and-down pattern exhibited by compression-induced hyperaemia at ISI = 80 s (Fig. 2 and Fig. 3)

seems to exclude a simple, frequency-dependent, desensitization mechanism of mechano-sensitive pathways,

as previously hypothesized (38, 60). More complex desensitization patterns possibly affecting multiple

mechanosensitive pathways cannot be excluded based on the present data. However, in order to explain the 441 peculiar hyperaemic responses observed at ISI=80 s, such desensitization pattern should exhibit an up-and-442 down time course, as exhibited by TOI, which would appear a quite unlikely coincidence. 443 444 445 Limitations Manual assessment of insonation angles, as required with the transversal approach, is not very accurate and 446 447 may introduce systematic errors in the calculation of absolute flow values. This is particularly true for 448 assessment of venous blood flow since a wide angle between the vessel axis and the ultrasound beam had to 449 be adopted in order to avoid saturation of the velocity signal (aliasing). However, the analysis was here focused on relative changes, thereby eliminating errors associated with measurement of the insonation angle. 450 Diameter of the femoral vein was not continuously monitored. Possible enlargement of the vessel during the 451 passage of the blood volume displaced by the compression may have resulted in underestimation of venous 452 453 flow. 454 Diameter of both femoral artery and vein exhibited a small increase throughout the experimental protocol, which was not accounted for. This may also have led to increasing underestimation of blood flow with time. 455 Since the sequence of the series was randomized this aspect should not have affected the results. 456 457 458 Conclusions This study allowed to prove that the attenuation of hyperaemia upon repetitive limb compression is not 459 dependent on vascular filling and the muscle pump nor on a simple ISI-dependent desensitization of 460 mechano-sensitive pathways. In addition, strong evidence is provided, supporting the concept that tissue 461 hyper-oxygenation is the key signal underlying the inactivation of the rapid dilatory response to muscle 462 compression. This evidence is however indirect and other studies are necessary to conclusively prove this 463 464 assertion. Irrespective of the underlying mechanisms, inactivation of mechano-sensitive pathways may almost 465 466 completely abolish the compression-induced hyperemia, suggesting a role for this phenomenon in protecting the tissue form hyperperfusion and oxidative stress. 467 The hyperaemic response to muscle compression is proposed as a peculiar model for the investigation of the 468 response to hyper-perfusion characterized by constant arterial pO₂, constant tissue metabolism as well as 469 470 modest or absent systemic reactions. Finally, tissue oxygenation monitoring is recommended to assess the efficacy of IPC treatments, oriented to 471 improve blood perfusion in limbs. 472

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- No competing interest to declare.

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LEGENDS TO FIGURES

626

- 627 **Fig 1**
- Experimental setup and typical hemodynamic response to a compressive stimulus.
- A) The experimental setup includes: the IPC system for the compression of the lower limb, eco-Doppler
- monitoring of blood flow from femoral vein and femoral artery, and NIRS monitoring at lateral head of
- gastrocnemius muscle. B) Typical response to leg compression in a representative subject. From top to
- bottom: blood velocity in femoral vein (BVFV), blood velocity in femoral artery (BVFA), tissue
- oxygenation index (TOI), total hemoglobin index (THI), changes in oxygenated hemoglobin (O₂Hb) and in
- deoxygenated hemoglobin (HHb) and the synchronism signal (Sync.), the thick and thin bars indicating start
- of inflation and deflation of the cuff, respectively.
- 636 **Fig 2**
- 637 Hemodynamic responses to repetitive compression at different inter-stimulus intervals (ISI).
- The ISI is indicated at the bottom of each column of bar-diagrams; each bar refers to the response to a single
- compressive stimulus. From top to bottom: Peak (arterial) blood flow, displaced (venous) blood volume, Pre-
- compression THI (indicating local vascular filling reached before the delivery of the compressive stimulus);
- Pre-compression TOI (indicating local tissue oxygenation before the stimulus). For the first three variables

642	and for each subject, responses have been normalized to the response to the first stimulus in the 160-s series
643	(white bar). * significantly different from the first response in the series (p<0.05)
644	Fig 3
645	Original recordings of the response to repetitive leg compression at inter-stimulus interval = 80 s, from
646	a representative subject.
647	Notations as in Fig.1. Note the pattern of response of arterial blood velocity in relation to tissue oxygenation.
648	The dotted line represents the initial TOI baseline.
649	Fig 4
650	Original recordings of the response to repetitive leg compression at inter-stimulus interval = 20 s, from
651	a representative subject.
652	Notations as in Fig.1. Note the complete disappearance of the hyperaemic response (BVFA) after the first
653	compressive stimulation, as long as tissue oxygenation (TOI) remains elevated, and the agreement between
654	the displaced blood volume (area under BVFV) and the current vascular filling (THI).
655	Fig 5
656	Scatter plots for assessing the correlation between different variables. Each dot indicates the response to
657	a single compressive stimulus in a single subject. Notations as in Fig. 2. (n=10). Straight lines indicate linear
658	regressions for individual subjects. Note that: the increase in tissue oxygenation is related to the peak blood
659	fow (A); Peak blood flow is not related to the displaced blood volume (B) but is inversely related to pre-
660	compression oxygenation level. In D the amplitude of the hyperaemic response (peak-baseline) is plot vs
661	pre-compression TOI to indicate that at high oxygenation levels the hyperaemic response may be almost
662	completely abolished.









