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2	functional electrical stimulation in spinal cord injury
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Femoral artery blood flow and microcirculatory perfusion during acute, low-level

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51 Abstract

52 Objective – Functional electrical stimulation (FES) may help to reduce the risk of developing 53 macro- and microvascular complications in people with SCI. Low-intensity FES has significant 54 clinical potential since this can be applied continuously throughout the day. This study 55 examines the acute effects of low intensity FES using wearable clothing garment on vascular 56 blood flow and oxygen consumption in people with SCI.

57 *Design* – Cross-sectional observation study

Methods – Eight participants with a motor complete SCI received 4x3 minutes of unilateral
FES to the gluteal and hamstring muscles. Skin and deep femoral artery blood flow and oxygen
consumption were measured at baseline and during each bout of stimulation.

61 *Results* – Femoral artery blood flow increased by 18.1% with the application of FES (*P*=0.02).

62 Moreover, femoral artery blood flow increased further during each subsequent block of FES

63 (P=0.004). Skin perfusion did not change during an individual block of stimulation (P=0.66).

64 Skin perfusion progressively increased with each subsequent bout (P<0.001). There was no 65 change in femoral or skin perfusion across time in the non-stimulated leg (all P>0.05).

Conclusion – Low-intensity FES acutely increased blood flow during stimulation, with a
progressive increase across subsequent FES bouts. These observations suggest continuous,
low-intensity FES may represent a practical and effective strategy to improve perfusion and
reduce the risk of vascular complications.

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76	Key words: Spinal cord injury, functional electrical-stimulation, blood flow, oxygen
77	consumption
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79	Abbreviations: FES (Functional electrical stimulation), SCI (spinal cord injury), DFA (deep
80	femoral artery, NO (nitric oxide)
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96 Introduction

97 A spinal cord injury (SCI) leads to significant changes in sub-lesional vascular structure and function. Most characteristic changes involve a decrease in conduit artery diameter¹, 98 increased vascular resistance², increased arterial stiffness³ reduced capillarization⁴ and 99 impaired cutaneous microcirculation^{5, 6} in the paralyzed, inactive limbs. Collectively, such 100 101 vascular changes are associated with endothelial dysfunction and the development of cardiovascular disease, which is a primary cause of death in persons with a SCI⁷. Besides 102 103 the increased risk of cardiovascular disease, below lesion microvascular endothelial 104 dysfunction manifested as impaired cutaneous blood flow also has significant implications for persons with SCI. The incidence and progression of skin breakdown lesions and pressure 105 106 ulcers in persons with SCI have been attributed to factors that are associated with a reduction in cutaneous microcirculation⁸. Interventions that help reverse macro- and microvascular 107 endothelial dysfunction below, and even above, the lesion are therefore of great clinical 108 significance for persons with SCI. 109

110

111 Studies show that elevations in blood flow and shear stress are required for improvement in vascular function and an increase in artery diameter^{9, 10}. Using electrically stimulated leg 112 exercise in individuals with SCI, Thijssen et al. showed evidence of arterial remodeling in 113 areas subject to electrically stimulated muscular contractions, while vascular adaptations 114 were not apparent in the passive, non-stimulated areas of the leg¹¹. In addition to conduit 115 remodeling, studies have shown that functional electrical stimulation (FES) results in 116 increased muscle mass¹², higher muscular oxidative capacity¹³, enhanced capillary supply⁴ 117 and improved blood flow². This highlights the potency of FES to mediate beneficial 118 adaptations. 119

121 Commonly used methods of FES require specialist facilities and trained staff, making regular application difficult, expensive and impractical. A potential alternative is the use of wearable 122 123 clothing garments with embedded surface electrodes that automatically stimulate muscles 124 when the garment is applied. This also allows for the adoption of low-intensity FES that can be applied for prolonged periods (i.e. during awake hours). Using this approach, an acute 125 126 bout of FES to the gluteal and hamstring muscles has shown to reduce pressure over the ischial tuberosity¹⁴ and increase transcutaneous oxygen levels¹⁵. To date, no study has 127 directly examined the acute impact of FES using a wearable clothing garment on both micro-128 129 and macro-vascular perfusion in people with SCI.

130

The purpose of this study, therefore, was to examine the acute effects of low-intensity FES (involving the gluteal and hamstring muscles) on deep femoral artery blood flow (i.e. supplying the active muscles) and skin microcirculatory perfusion (i.e. covering the active areas). It was hypothesized that an increase in conduit artery and skin blood flow would occur with muscle stimulation, whilst also having a cumulative effect leading to a gradual increase in baseline perfusion with repeated application of FES.

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138 Materials and Methods

Participants. Eight male individuals with ASIA A or B classified SCI participated in this study. All participants were outpatients and frequently visited Reade rehabilitation center for checkups with their physician and to participate in sporting activities. All injuries were traumatic in origin and existed for at least 1 year prior to undergoing the study. None of the participants had any known cardiovascular diseases or took any medication known to interfere with the cardiovascular system. Exclusion criteria included individuals with flaccid paralysis (i.e. inability to activate the muscles through nerve stimulation), a previous history
of autonomic dysreflexia during FES (i.e. for safety purposes) and intolerance or
contraindication for the use of FES. The local institutional medical ethical board of Reade
Rehabilitation center approved the study and all participants provided written informed
consent after receiving and understanding full details of the research study. This study is
reported in accordance with the STROBE guidelines and conforms to all items on the
checklist accordingly (see supplementary checklist).

152

153 Electrical stimulation. FES was applied using a specially developed garment with embedded surface electrodes (Axiobionics, Ann Arbor, MI, USA), connected to a portable 154 155 battery-operated stimulator (Neuropro, Berkelbikes, Nijmegen, The Netherlands). All wires 156 and leads were embedded within the seam of the garment to prevent them becoming 157 entangled with the patient. The FES garment was made from elastic lycra and secured to the 158 body using foldable Velcro straps (Fig 1). One surface electrode was positioned at the upper 159 (proximal) part of the gluteal muscle and a second about halfway down the hamstring area, preventing the participants from lying directly on the electrodes with their buttocks. 160 161 Ultrasound gel was placed in small Velcro pouches to be used as a conductor between the electrodes and the participants' skin. FES was applied to the right leg only at a standard 162 163 constant voltage of 150V using 50Hz biphasic impulse frequency to induce a visible tetanic 164 contraction. The amplitude needed to induce a strong muscle contraction depends on muscle denervation and the amount of muscle nerve fibers that can be recruited and activated. Due to 165 the variability between individuals, the current amplitude was subjectively determined by the 166 167 researcher and individualized for each participant with increments of 5 to 10mA to a level that did not cause discomfort or excessive movement. To minimize muscle fatigue and ensure 168

169 continuous muscle contractions, a 1:4 duty cycle, consisting of 1-second stimulation followed
170 by 4 seconds without stimulation for a period of 3 minutes was used¹⁴.

Protocol and testing procedure. Participants attended the laboratory at Reade rehabilitation 171 172 center once to undergo testing. Due to sympathetic nervous system activation and the effects on hemodynamics and blood pressure, all participants were asked to refrain from alcohol and 173 174 caffeine consumption 24 hours prior to testing. On arrival, the protocol and testing procedures were explained in full to each participant. Participants were transferred from their 175 176 wheelchair to a bed and positioned comfortably in the supine position. Subsequently, the 177 shorts were applied to ensure correct placement of the electrodes. After a 10-minute rest 178 period and before the start of stimulation, baseline measurements were made for oxygen 179 consumption (VO₂), skin blood flow, and deep femoral artery (DFA) blood flow in the 180 control and intervention leg. After baseline measurements, the protocol included four blocks 181 of stimulation lasting 3 minutes interspersed with 17 minutes of no stimulation (Fig 2). We 182 chose four blocks of stimulation so we could determine the response and potential benefits of 183 repeated exposure to FES (i.e. a pattern that would be applied in practice). Recordings for all 184 measures were collected 1 minute before and 3 minutes throughout stimulation. 185 Measurements of DFA diameter and blood flow velocity during stimulation were performed in the intervention leg only. Since it was unlikely that FES would alter blood flow in the 186 187 contra-lateral, non-stimulated leg (i.e. a systemic effect), we did not measure blood flow in 188 the non-stimulated leg.

189

190 Experimental Measures.

Femoral artery blood flow. Velocity and diameter in the right DFA was measured using a 2dimensional echo Doppler ultrasound. Using a 10-MHz-multi-frequency linear array probe
attached to a high resolution ultrasound machine (T3000, Terason, Aloka, UK), optimal

longitudinal B-mode images capturing the lumen/arterial wall interface, along with Doppler
velocity measures of the DFA, approximately 2cm from the bifurcation were obtained.
Following image acquisition, 1 min baseline imaging was performed in the control and
intervention leg. The same examiner performed all measurements and images were recorded
for later offline analysis.

199

200 Skin microcirculatory perfusion. We used laser Doppler flowmetry (Periflux system 5000, 201 Perimed AB, Järfälla, Stockholm, Sweden) to obtain an index of microcirculatory perfusion. 202 This is a non-invasive technique that enables evaluation of skin microvascular blood flow over a period of time and is sensitive at detecting changes in response to a stimulus. The 203 204 technique uses a beam of laser light that undergoes a change in wave lengths when it detects 205 moving red blood cells. The specific changes in wavelength are characterized by red blood 206 cell concentration and velocity to give a measurement of skin blood flow expressed as 207 arbitrary perfusion units (PU). After the FES shorts had been applied and the participant was 208 comfortably lying in a supine position, the laser Doppler flowmetry probes were placed at the measurement site. Blood flow was continuously measured at the skin covering the gluteal 209 210 muscle on the stimulated leg. A small incision was made in the shorts to allow placement of 211 the laser Doppler probe in close proximity to the stimulated muscle and to ensure fixation 212 throughout the protocol.

213

Oxygen consumption. Oxygen consumption was collected throughout using a facemask
connected to an online gas analyser (Oxycon Pro, Jaeger, The Netherlands). Volume and gas
concentration calibrations were performed prior to each test. The participants were instructed
not to talk during the measurements.

218 Data Analysis

219 DFA diameter and blood flow. Post-test analysis of the DFA was performed using custom-220 designed edge-detection and wall tracking software which is largely independent of 221 researcher bias. Thorough details of the analysis technique have been described elsewhere ¹⁶. 222 Briefly, data collected on the ultrasound machine were stored as a digital avi file. Subsequent software analysis of the data was performed at 30 Hz using an icon-based graphical 223 224 programming language and toolkit. The initial phase of analysis required selecting an optimal region of interest (ROI) on the B-mode image, which allowed for automated calibration of 225 226 artery diameter. Within the ROI, a pixel density algorithm automatically identified the angle 227 corrected near and far wall e-lines. Finally a ROI was drawn around the Doppler waveform and automatically detected the peak of the envelope for this waveform. The mean diameter 228 229 measure was calculated from within the B-mode ROI and synchronized with the velocity 230 measure which was calculated from the Doppler ROI at 30 Hz. The product of this (artery 231 cross-sectional area and Doppler velocity) gives a measure of average blood flow (mL/s). We 232 have shown that analysis using this semi-automated method produces reproducible diameter 233 calculations that are significantly better than manual methods and producing an intraobserver coefficient of variation of 6.7%¹⁷. 234

235

Skin microcirculatory perfusion. Dedicated software (Perisoft for Windows) was used to
collect, store and analyze the skin blood flow data. Unwanted artefact in the data due to
participant/wire movement was identified and removed from the data prior to analysis.
Resting values were calculated by averaging the last 3 minutes of rest before the start of the
next stimulation block, whilst perfusion during stimulation was presented as averages every
30-s.

Oxygen consumption. Five-second bins of gas analysis data were exported to Excel. Steady
state average values were calculated from the last minute of rest prior to stimulation and
during the entire 3 minutes of stimulation.

246

247 Statistical Analysis

248 Statistical analysis was conducted using the Statistical Package for the Social Sciences. All data were expressed as means \pm SD and statistical significance was set at P<0.05. Linear 249 250 mixed models were used to examine the impact of FES on femoral artery and skin 251 microcirculatory blood flow (main effect of "stimulation": baseline vs stimulation), but also whether the stimulation-induced changes differed across the 4 blocks of stimulation (main 252 253 effect for "blocks"). The repeated covariance type was compound symmetry and stimulation, 254 blocks and stimulation*blocks were specified as fixed effects and as estimated marginal 255 means. The test of fixed effects stimulation*blocks interaction was interpreted. Significant 256 main effects of stimulation, blocks and stimulation*blocks interaction were followed up with 257 a simple main effects analysis and the least significant difference (LSD) approach to multiple 258 comparisons.

259

260 **Results**

261 *Conduit artery*. There was a significant main effect of stimulation on DFA blood flow

262 (*P*=0.02). On average, arterial blood flow increased by 18.1% from 4.69 mL/s at first baseline

263 (pre-intervention) to 5.52 mL/s during 3 minutes of FES (Fig 3). There was also a significant

264 main effect for "blocks" (P=0.004), indicating that perfusion at each subsequent baseline and

- 265 perfusion during stimulation was different across repeated blocks. More specifically, blood
- flow in block 2 (P=0.02), 3 (P=0.01) and 4 (P<0.001) were all significantly higher than
- during block 1. There was no stimulation*block interaction (P=0.74).

268 To assess changes in arterial blood flow in the control leg we used a paired samples t-test.

Femoral blood flow in the control leg did not change from pre (3.86±1.66 mL/s) to post-

270 stimulation (3.64 \pm 1.52 mL/s; $t_3 = 0.97$, P = 0.41).

271

272 Skin blood flow. There was no significant main effect for stimulation (Fig 4) (P=0.66),

273 indicating that there was no immediate change in perfusion with stimulation when compared

to baseline. However, perfusion did increase over time with repeated stimulation resulting in

a significant main effect for "blocks" (P < 0.001). Skin blood flow, expressed as perfusion

units (PU) significantly increased from block 1 (12 ± 6 PU) to block 2 (17 ± 9 PU; P=0.01) and

block 3 (22±13 PU; P<0.001) and was ~80% higher during block 4 compared to block 1

278 (22 \pm 13 PU; P<0.001). Blocks 3 and 4 were also greater than block 2 (P=0.004), but

279 plateaued between blocks 3 and 4. There was no stimulation*blocks interaction (P=0.99).

280

281 *Oxygen consumption.* Oxygen consumption did not change throughout the stimulation 282 protocol. There was no significant main effect for stimulation (P=0.98), the number of 283 stimulation blocks (P=0.94) or stimulation* block interaction (P=0.87).

284

285

286 Discussion

287 The main finding of this study was that unilateral FES acutely increased femoral blood flow

in the stimulated leg, most likely a direct result of the increased oxygen demand of the

activated gluteal muscles. Skin microcirculatory perfusion also increased from pre-

intervention baseline, although the response was more gradual and was not evident during the

291 3-minute stimulation blocks. Additionally, resting femoral artery blood flow and skin

292 perfusion both progressively increased with repeated bouts of stimulation. Collectively, these

results indicate that low-intensity FES was effective at inducing hemodynamic changes in the
superficial and deep layers of the gluteal region. Since frequent increases in blood flow
represent a key stimulus for improvement in micro- and macrovascular function and
structure¹⁰, these observations warrant further research to examine the potential effects of
repeated exposure to low-intensity FES on the vasculature in individuals with SCI.

298

299 Blood flow in Stimulated Leg

300 This study is the first to examine conduit artery blood flow and skin microcirculatory 301 perfusion in SCI following acute application of FES using a wearable clothing garment. As anticipated, the results show an immediate increase in deep femoral blood flow, even when 302 303 performed using our low-intensity FES protocol. These findings are consistent with previous data from studies in able bodied¹⁸ and individuals with SCI¹⁹. These previous studies 304 observed a 95% increase in blood flow in the femoral artery during FES. Although we 305 306 observed a modest increase of 20%, this difference between studies is most likely attributable 307 to distinct stimulation parameters. Whilst in the current study, only two muscle groups were stimulated using a stimulation level that allowed for muscle contractions without overt limb 308 309 movement (m=75mA), previous work used whole leg muscle stimulation inducing significant muscle movement and therefore marked oxygen demand of the activated muscles. The co-310 311 contractions used in the aforementioned studies are also likely to further increase oxygen 312 demand and contribute to greater arterial inflow and blood distribution throughout the entire limb. Nonetheless, it must be emphasized that the large muscle stimulation with marked 313 314 movement can only be applied for ~20 minutes. Muscle fatigue and energy source depletion 315 prevents longer duration stimulation, whereas low-intensity FES can be applied throughout the day and night and on a day-to-day basis. Although our protocol only increased blood flow 316 by $\sim 20\%$, the ability for prolonged exposure to low-intensity FES in individuals with SCI 317

318 make the FES-protocol applied in the present study a physiologically significant and319 potentially clinically relevant stimulus.

320

321 An important question relates to the mechanisms responsible for the increase in perfusion. 322 Since the current study found no changes in DFA blood flow in the non-stimulated leg, the 323 possibility of systemic stimuli affecting perfusion (e.g. blood pressure) can be excluded. During muscular contractions, a number of mechanisms are known to regulate arterial blood 324 325 flow supplying the active muscles. Firstly, an increase in cell metabolism initiates the 326 localized release of vasodilator metabolites such as nitric oxide (NO), prostacyclin, ATP, adenosine and potassium from contracting skeletal muscle and the vascular endothelium^{20, 21}. 327 328 The release of such compounds initiates vascular smooth muscle relaxation, vasodilation of 329 the artery and a subsequent increase in blood flow to the stimulated region. During exercise, skeletal muscle blood flow increases in proportion with metabolic activity to meet the oxygen 330 demands of the contracting muscle²². Considering the direct relationship between skeletal 331 332 muscle blood flow and metabolic load, it seems sensible to assume that the small, albeit significant, increase in arterial blood flow is due to the low-intensity stimulation protocol we 333 334 used.

335

Another physiological impact of low-intensity FES must be considered. The dynamic and mechanical effect of muscle contractions and relaxations, or the 'muscle pump' mechanism, importantly influences blood flow in the vasculature. During muscle contraction, a decrease in venous pressure occurs as venous blood empties from peripheral areas (i.e. the legs) and is propelled to the central circulation^{23, 24}. The emptying of venous segments leads to an increase in arteriovenous pressure gradient facilitating an increase in arterial inflow as the muscle relaxes^{25 24}. Although this study did not differentiate changes in blood flow during the 343 contraction and relaxation phases of muscle stimulation, the increase in arterial blood flow
344 may, at least partially, be explained through increased muscle pump activity and increases in
345 the arteriovenous pressure gradient.

346

347 Microcirculatory Perfusion

348 Changes in skin microcirculation occur as a reflex thermoregulatory control mechanism during whole body and/or localized changes in temperature²⁶. In the current study, we 349 observed little change in skin perfusion during an individual block of stimulation. However, 350 351 the combined effect of consecutive and repeated exposure to stimulation did result in a successive rise in skin perfusion over the duration of the protocol. Considering there was no 352 353 change in whole body VO₂, it is unlikely that an increase in core body temperature could 354 explain the progressive rise in skin perfusion. A more likely explanation relates to localized 355 heat production and a subsequent gradual warming of the skin covering the activated 356 muscles. This would result in a sustained rise in skin blood flow during localized heating which, is mediated through the release of NO from the vascular endothelium²⁷. Regardless of 357 any change in skin temperature, previous work confirms a NO mediated increase in skin 358 perfusion in response to FES^{28} . Petrofsky and colleagues observed an increase in skin blood 359 360 flow during FES that was prevented with the infusion of L-NAME, a NO inhibitor. Although 361 the current study nor Petrofsky et al. controlled for potential changes in skin temperature, its contribution to the gradual rise in skin perfusion should not be excluded. Future research 362 should consider the exact mechanisms involved in the increase in skin perfusion during FES 363 364

365 Oxygen Consumption

There was no change in VO₂ during the stimulation protocol, which is in contrast to other
studies using FES whilst sitting or lying^{18, 29}. In the current study, only two muscle groups

368 were stimulated using low level FES for 3 minutes. Given the increase in blood flow, it 369 seems logical that energy expenditure in these muscles increased. However, energy 370 expenditure has previously been shown to increase in a dose response relationship with 371 stimulation intensity and the number of muscles stimulated. The small dose of stimulation 372 adopted in the current protocol may be insufficient to detect a significant increase in whole 373 body VO₂. Indeed, previous work that reported higher oxygen consumption upon FES adopted higher stimulation (100 mA and 93 mA), but also stimulated a larger muscle mass¹⁸, 374 ²⁹. These previous studies confirm that FES has the potential to increase VO_2 and energy 375 376 expenditure which, is indirectly supported by our observation of increased perfusion, and 377 therefore oxygen delivery to the large muscle mass in the legs and gluteal region. One should 378 also consider that changes in oxygen consumption in the current study (involving unilateral 379 FES) may increase exponentially more when FES is applied in a clinical situation using 380 bilateral stimulation.

381

382 Study Limitations

The small sample size we used may overestimate the true effect of stimulation on vascular 383 perfusion. That said, our data clearly show a distinctive increase in perfusion with FES and 384 we are therefore confident that the results of this study are representative of the wider SCI 385 386 population. Secondly, due to equipment failure, we were unable to obtain skin blood flow 387 measures in the contralateral, unstimulated leg. However, the low intensity stimulation 388 protocol we used is unlikely to induce any systemic effects on cutaneous perfusion. This is 389 supported by the absent changes in the deep femoral artery in the contra-lateral, non-390 stimulated leg. Finally, FES induced autonomic dysreflexia is a potential side effect that may 391 limit its usage in some individuals. Although this was an exclusion criterion in the current 392 study, it has previously been reported to occur at higher current amplitudes (160mA) during FES-assisted hydraulic resistance training exercise.³⁰ Blood pressure monitoring is therefore 393

394 recommended for novice users.

396	In conclusion, this study clearly shows an increase in superficial and deep vascular perfusion
397	during low level FES. A ~20% increase in blood flow occurred through the deep femoral
398	artery supplying the gluteal muscles, most likely through local increased oxygen demand and
399	muscle pump activation. The results also show a gradual and consistent increase in skin
400	perfusion over the duration of the protocol. This may represent a potent stimulus when this
401	type of low-intensity FES is applied for several hours. Future work is required whether such
402	physiological changes translate to a clinically relevant effect, especially given its simplicity
403	and ability for home-based, day-to-day use.
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Does arterial shear explain the magnitude of flow-mediated dilation?: a comparison

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506	Figure Legends
507	
508	Figure 1: Example of electrical stimulation shorts and how they are worn
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510	Figure 2: Schematic of stimulation protocol
511	
512	Figure 3: Deep femoral artery blood flow at baseline and during stimulation using low-
513	intensity ES in the stimulated leg. Data are presented for each block of stimulation. Error bars
514	represent standard deviations. *P<0.05 vs. Block 1 #P<0.05 vs. Baseline
515	
516	Figure 4: Skin blood flow at baseline and during stimulation using low-intensity ES in the
517	stimulated leg. Data are presented for each block of stimulation. Error bars represent standard
518	deviations. *P<0.05 vs. Block 1 †P<0.05 vs. Block 2
519	
520 521	
522	

Subject	Age (yr.)	Level of	ASIA	Time Since Injury	Stimulation
		injury	score	(yr.)	level
1	40	T9	А	10	75
2	30	C6	А	16	70
3	57	C8/T1	В	15	60
4	54	C6	А	28	85
5	34	T2	А	10	75
6	29	T8	А	9	85
7	60	T8	А	8	70
8	43	C6	А	16	80
Mean	43	-	-	14	75mA

TABLE 1. Characteristics of SCI individuals



Figure 1.





tago

End



P=0.02

P=0.004

P=0.74

