Effects of a novel neurodynamic tension technique on muscle extensibility and stretch tolerance: a counterbalanced cross-over study.

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1 Abstract.

2 **Context:** Neurodynamic tension affects hamstring extensibility and stretch tolerance, and is 3 considered important in hamstring injury management. Neurodynamic tension was postulated 4 to affect segmental muscle extensibility and stretch tolerance, and potentially also demonstrate extra-segmental and contralateral effects. Objectives: Assess the effects of a novel sciatic-5 6 tibial neurodynamic tension technique, the modified long sit slump (MLSS), on segmental, 7 extra-segmental and contralateral muscle extensibility and stretch tolerance. Study design: 8 Counterbalanced cross-over study. Setting: University research laboratory. Participants: 9 Thirteen healthy and active subjects (mean \pm SD age 24 \pm 8 y, BMI 23.1 \pm 2.8 kg·m⁻²). 10 **Intervention:** MLSS application (5 seconds, 5 repetitions, 3 sets) on two occasions with a 11 three-week washout period, and either stance or skill leg treated in a counterbalanced manner. 12 Main outcome measures: Segmental and extra-segmental muscle extensibility were measured 13 utilising passive straight leg raise (PSLR) and prone knee bend (PKB) at pre-, immediately 14 post- and one hour post-intervention. Stretch intensity ratings were measured utilising a simple 15 numerical rating scale (SNRS). Results: MLSS significantly increased PSLR and PKB 16 bilaterally (p<0.001). The effect for PSLR was greater in the ipsilateral leg compared to the contralateral leg (baseline to one hour post: $+9\pm6^{\circ}$ and $+5\pm5^{\circ}$ respectively, p<0.001), but not 17 18 for PKB (baseline to one hour post: ipsilateral leg $+5\pm5^\circ$, contralateral leg $+5\pm4^\circ$). For both 19 PSLR and PKB the effect of the first session was retained at the start of the second session 3 20 weeks later. SNRS data were consistent with increased stretch tolerance. Conclusions: 21 Application of a novel sciatic-tibial neurodynamic tension technique, the MLSS, increases 22 muscle extensibility and stretch tolerance segmentally, extra-segmentally and contra-laterally. 23 Level of evidence: 2C Outcomes research.

Key words: flexibility, hamstrings, muscle extensibility, neurodynamics, stretching, neuronal
desensitisation.

26 **INTRODUCTION**

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Hamstring strain injury (HSI) is one of the most common non-contact injuries in athletes.¹⁻³ 28 with high rates of recurrence,⁴ despite considerable research efforts.⁵ The role of hamstring 29 flexibility, also termed extensibility herein, in HSI, ^{4,6-7,11} re-injury and rehabilitation, ^{2,8,12,13} 30 has not been fully elucidated to date. ⁸⁻¹⁰ Neurodynamics is a term describing mobilisation of 31 the nervous system and its surrounding structures.¹⁴⁻¹⁵ Neurodynamic tension techniques 32 elongate the neural tissue and are considered to increase nerve tension and strain, whereas 33 neural sliding techniques aim to maximise nerve excursion.¹⁶ Neurodynamic tension has been 34 demonstrated to significantly influence hamstring extensibility¹⁷⁻¹⁸ and is considered important 35 in HSI, re-injury and rehabilitation.¹⁹⁻²⁰ For example, Turl & George²⁰ demonstrated 57% of 36 elite rugby players with recurring grade one HSI demonstrated positive slump test²¹ after 37 returning to play, suggesting suboptimal neurodynamics may contribute to known high rates 38 of re-injury.^{4,22} Similarly, Kornberg & Lew¹⁹ demonstrated inclusion of a neurodynamic 39 40 tension technique to rehabilitation of Australian Football League players with HSI resulted in significantly faster return to play. 41

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43 Human in-vivo hamstring stretching studies in non-injured subjects strongly supports 44 stretch tolerance as a primary mechanism responsible for lasting increases in hamstring 45 extensibility utilising intervention protocols of up to eight weeks duration, with longer term stretching postulated to potentially induce structural alterations in hamstring muscle length and 46 passive stiffness.²³⁻²⁵ Immediate stretch-induced changes in hamstring passive stiffness are 47 considered to be due to viscoelastic stress relaxation, with effects typically potentiated within 48 five loading cycles and attenuated within an hour.²⁶ Previous research has demonstrated lasting 49 increases in hamstring extensibility are of similar magnitude irrespective of the stretching 50

protocol utilised, citing total weekly stretch time as the most important variable.²⁷⁻²⁹ However, there is some evidence that more intense stretching may effect greater changes in extensibility, or at the very least, saves time and is therefore considered more efficient.^{28,30} As neurodynamic tension is associated with relative increased levels of reported stretch intensity during hamstring stretch for a common ROM,^{17,31} it was postulated that it may have a significant role in afferent modulation of stretch tolerance.^{18,25}

Compared to muscle stretching protocols, there has been relatively little research 57 investigating utilisation of neurodynamic techniques on lasting changes in hamstring 58 extensibility and stretch tolerance.^{18,32-33} For example, Castellote-Caballero and colleagues³² 59 60 demonstrated a significant increase in passive straight leg raise (PSLR) of nine degrees 61 following three sessions of a neurodynamic slider over one week. Although comparatively this 62 is an average PSLR gain for a hamstring extensibility study, it was achieved in a relatively short period of time.³⁴⁻³⁵ More recently, Sharma and co-workers¹⁸ reported significantly greater 63 64 hamstring extensibility gains when neurodynamic techniques and muscle stretching were 65 utilised compared to muscle stretching alone, but the intervention dosing between the groups was inconsistent which lessens the strength of conclusions drawn from this randomised 66 67 controlled trial (RCT).

68 The specific groups of afferent neurones primarily affected during stretching and modulation of stretch tolerance are yet to be fully elucidated.^{25,36} Small and large diameter 69 proprioceptors are fundamentally implicated in stretch sensation, but a significant role of 70 71 mechanosensitive nociceptors has also been suggested and warrants more detailed consideration.^{24,36-39} As initiation of stretch discomfort has been reported to occur at 85% of 72 muscle passive torque values recorded for maximal stretch tolerance,⁴⁰ direct activation of 73 74 mechanosensitive nociceptors resulting from stretch-induced tensile strain, secondary compression, or a combination of the two, is probable.^{37-38,41} 75

76 Notwithstanding likely short term modulation of stretch tolerance through an inhibitory nociceptive 'gating' mechanism at the spinal dorsal horn through activation of non-nociceptive 77 afferent fibres, ^{36,42-44} proprioceptor and mechanoreceptor discharge in the early stage of muscle 78 79 stretch could sensitise mechanosensitive nociceptor discharge towards activation thresholds,^{38,41,46} particularly as peripheral afferent neuropeptides are largely unspecific to fibre 80 type.^{38,46-47} This is likely accentuated by mechanisms such as the axon reflex and afferent 81 convergence.^{38,45} Furthermore, the same afferent neuropeptides which are utilised distally are 82 produced in dorsal root ganglia,⁴⁶⁻⁴⁷ the neuropeptides having both peripheral and central 83 neuromodulatory effects that may outlast the duration of stretch.^{25,36} Moreover, the parameters 84 85 and context of stretching likely affect spinal and supraspinal processing, which may also alter 86 the diffuse noxious inhibitory system (DNIS), and has also been implicated in modulation of stretch tolerance through conditioned learning.^{36,44} 87

Inter-neuronal activation and recruitment of latent nociceptive circuits is considered a primary mechanism by which pain spreads segmentally, extra-segmentally and contralaterally.⁴⁸⁻⁵² Given such central pain sensitisation has been considered a form of neuronal long term potentiation (LTP) and learning,^{42,44,53-54} it was postulated herein that the increased stretch tolerance from stretching could be a form of neuronal long term depression (LTD),^{43,55} and stretch tolerance may also demonstrate a similar course of segmental, extrasegmental and/or contralateral effect, given the appropriate stimulus.^{51,56}

Therefore the study hypothesis was that application of a novel sciatic/tibial nerve neurodynamic tension technique, the modified long sit slump (MLSS), would increase muscle extensibility and stretch tolerance segmentally, extra-segmentally, and contra-laterally.

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99 METHODOLOGY

101 Study design

102 A counterbalanced crossover experiment over two intervention sessions was utilised, with each 103 intervention session utilising a single limb from each subject (Figure 1). In order to avoid 104 effects of intervention order and/or limb dominance, the treatment order was counterbalanced 105 with 7 subjects having the stance leg treated first and the remaining 6 subjects receiving 106 treatment on the skill leg first, the skill leg defined as that which the subject reported to preferentially use to kick a ball. Previous research has not demonstrated any contralateral 107 effects from unilateral stretching^{24,32,36} and a three week 'wash out' period was deemed 108 sufficient for any treatment effects to wear off.^{28,57} The independent variables were unilateral 109 110 neurodynamic intervention (MLSS) over two sessions, the dependent variables being 111 ipsilateral and contralateral hamstring and rectus-femoris extensibility and stretch tolerance. 112 The dependent variables were measured pre-, immediately post- and one hour post-113 intervention. Subjects were requested not to partake in unfamiliar physical activity for three 114 days prior to testing and strenuous physical activity on the day of testing, and not to stretch the 115 lower limbs between intervention sessions. All testing was performed in a university laboratory. Recruitment and data collection occurred between March and April 2016. 116

117 **Participants**

118 A healthy and active sample of convenience was recruited from a university population. Assuming alpha = 0.05 with 80% power and utilising one degree standard error of measurement 119 120 and four degree minimum detectable difference for a hand held inclinometer, a priori sample calculation was 12.⁵⁸ Subjects were recruited via print poster, electronic university noticeboard. 121 122 and limited e-mail recruitment. One extra subject was recruited in case of drop out, with a final sample size of 13 (9 male, 4 female, mean \pm SD age 24 \pm 8 years, Body Mass Index 23.1 \pm 2.8 123 $kg \cdot m^{-2}$). Healthy and active was defined as no history of significant medical conditions and a 124 minimum Tegner Activity Scale⁵⁹ rating of five, respectively. Further exclusion criteria were 125

126 significant neurological or orthopaedic conditions, past history of HSI, significant low back 127 pain, and participation in a formal hamstring lengthening or strengthening program in the previous six months. Subjects with clinically 'tight' hamstrings were recruited, adopting values 128 equal or lower than 75° for men and 80° for women, with potential participants with PSLR 129 above these values excluded from the study.^{34,60-61} Ethics approval was obtained through the 130 131 University of Bath Research and Ethics Approval Committee for Health (REACH; EP 14/15 201) and suitable subjects were required to provide signed, informed consent. The rights of all 132 133 subjects was protected.

134 **Procedures**

135 Subjects were screened for clinically 'tight' hamstrings by PSLR utilising a hand held inclinometer (Isomed AcuAngle).^{58,62} The subject lay supine with the non-tested thigh secured 136 137 to the plinth with a firm adjustable strap. The base of the inclinometer was marked on the 138 anterior distal tibia of the tested leg, corresponding to the zero value. The inclinometer was 139 secured with Velcro straps and the subject was instructed to fully relax during testing. The 140 examiner raised the leg slowly until the subject expressed maximal stretch tolerance was 141 reached or firm resistance to further elevation was encountered. The subjects were given a standard set of scripted instructions for the PSLR procedure, with only one measure utilised 142 143 for screening, consistent with clinical practice.

144 Assessment

PSLR was utilised as the ipsilateral and contralateral segmental muscle extensibility measure, as described above. A simple numerical rating scale (SNRS), with zero representing 'no muscle stretch' and ten representing 'the worst muscle stretch imaginable' was utilised as a subjective measure of stretch intensity.³⁶ SNRS measures were taken at maximal PSLR ROM for pre and post intervention time points (SNRS Max), and at the pre intervention maximal PSLR ROM for the post intervention time points (SNRS Com). If post intervention PSLR was less than pre

intervention, SNRS Com was not assessed. Ipsilateral and contralateral extra-segmental 151 152 extensibility of the rectus-femoris was measured utilising a prone knee bend (PKB) procedure. 153 Subjects lay prone with a strap stabilising the pelvis applied at the level of the lower half of the 154 sacrum. The subject's tested hip was positioned in approximately 10° extension by placing a high density foam roll between the thigh and the plinth, immediately proximal to the superior 155 156 patella. The examiner slowly flexed the knee until the subject expressed maximal stretch 157 tolerance was reached or further ROM was blocked by the posterior thigh. The examiner then 158 placed the inclinometer on the previously marked points on the tibia to measure ROM. PKB 159 SNRS stretch intensity measurement procedures were as for PSLR. All measurements were 160 repeated 5 times, the fifth of which was recorded. Subjects remained in the laboratory resting 161 room between immediate and one hour post-intervention assessments.

162 Warm-up

A light warm-up of 10 minutes of cycling on a stationary bicycle at a minimal resistance was
adopted immediately prior to intervention, with subjects instructed to maintain an intensity
whereby they were not short of breath.

166 Intervention

167 The MLSS intervention is shown in (Figure 2): In the starting position, subjects were 168 positioned hemi-sitting on a plinth (adjusted to height approximately 15 cm below greater 169 trochanter), with the stretched limb resting on the plinth while the other limb rested parallel on 170 the floor. With the knee on the plinth flexed in the starting position, the subject used their 171 opposite hand to reach forward to hold the lateral border of the opposite foot, placing it in dorsiflexion and eversion. This action maintains trunk flexion and relative internal rotation of 172 173 the tensioned leg. The subject was then instructed to straighten the knee and internally rotate 174 the femur with overpressure on the anterolateral distal thigh with the ipsilateral hand. The 175 therapist assisted to facilitate sciatic/tibial tract tension positions and if full neurodynamic 176 elongation was well tolerated the patient was asked to add further trunk and cervical flexion, 177 but only two subjects tolerated the additional trunk and cervical MLSS component in this 178 sample with clinically tight hamstrings. Stretch duration was 5 seconds, 5 repetitions and 3 179 sets, paced with a mobile metronome set at 1 Hz (Android 1.2.4; 2012). Subjects were given 180 10 seconds rest between repetitions and two to three minutes between sets. Subjects were 181 clearly instructed before and during the intervention sessions that the stretch procedure aimed 182 to achieve maximal stretch tolerance and may involve some discomfort, however, if the stretch 183 became too uncomfortable they should notify the tester immediately to reduce stretch intensity. 184 Similarly, subjects were also instructed to report symptoms such as pins and needles, numbress 185 or discomfort proximal to the ischial tuberosity.

186 Data analysis

187 Data analysis was performed using SPSS for windows. Exploratory data analysis and 188 significance testing utilising the Shapiro-Wilk test suggested the pre-intervention data was 189 normally distributed. Comparison of mean pre- to post-intervention PSLR and PKB ROM and 190 SNRS ratings was carried out utilising 3-way repeated measures analysis of variance 191 (ANOVA) with the factors session (1 / 2), side (ipsilateral / contralateral) and time (pre / post / post 1 hour). Post hoc analysis using Bonferroni correction was performed to determine 192 193 differences between time points for analyses with a significant main effect of time. If 194 assumption of sphericity was violated utilising Mauchley's test, the data was corrected with 195 the Greenhous-Geisser equation. Post hoc correlation analysis was also performed utilising 196 Pearson's correlation coefficient. Significance was set at alpha = 0.05 for all statistical tests.

197

198 **RESULTS**

200 Figure 3A shows the changes in PSLR following MLSS. MLSS significantly increased PSLR 201 directly after the intervention, with no further increase 1 hr later (main effect of time: p<0.001). 202 The effect of the unilateral MLSS intervention was evident in both legs, but greater in the 203 ipsilateral leg compared to the contralateral leg (baseline to one hour post: $+9\pm6^{\circ}$ and $+5\pm5^{\circ}$ 204 respectively, main effect of side: p<0.001). PSLR increased to a similar extent in both sessions 205 (no significant session x time interaction effect), despite the fact that the effect of the first 206 session was retained at the start of the second session 3 weeks later (main effect of session: 207 p<0.001).

208 The effects of the MLSS intervention on PKB were mostly similar (Figure 3B), with 209 significant main effects of time (p<0.001) and session (p<0.001). PKB increased from baseline 210 to directly post (p<0.001), but there was no further significant increase one hour following the 211 intervention. There was no significant effect of side, with similar effects on the ipsilateral leg and the contralateral leg (baseline to one hour post: $+5\pm5^{\circ}$ and $+5\pm4^{\circ}$ respectively). Post-hoc 212 213 analysis also revealed moderate to strong negative correlation between pre-intervention ROM 214 and the size of the ROM treatment effect for both PSLR (r=-0.32; p<0.05) and PKB 215 immediately (r=-0.56; p<0.001), and one hour post intervention (r=-0.53; p<0.001; r=-0.68, p<0.001). 216

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Subjective stretch intensity ratings were consistent with increased stretch tolerance following the MLSS intervention (**Table 1**). Post-intervention ratings taken at the preintervention maximal joint angle decreased for the PSLR (main effect of time: p<0.001), with a greater decrease in the ipsilateral side (main effect of side: p<0.001; time x side interaction effect: p<0.05). Conversely, ratings at the maximal joint angle achieved at each time point increased (main effect of time: p<0.01), again with a greater change in the ipsilateral side (main effect of side: NS; time x side interaction effect: p<0.001). PSLR stretch intensity ratings were
higher in the second session compared to the first session (main effect of session: p<0.001).

226 PKB stretch intensity ratings at the pre-intervention joint angle followed a pattern 227 similar to the PSLR ratings, with a significant decrease following the intervention (main effect 228 of time: p<0.001), and higher ratings during the second session (main effect of session: 229 p<0.05), but no significant main effect of side or time x side interaction effect (**Table 1**). No 230 significant main effects of time, session, or side, and no interaction effects were observed for 231 PKB stretch intensity ratings at the maximal joint angle achieved at each time point. No 232 differences were observed in the responses for any parameters between participants who 233 received the initial treatment on their skill leg or stance leg.

234

235 **DISCUSSION**

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237 The purpose of the study was to assess potential segmental, extra-segmental and contra-lateral 238 effects of applying a novel sciatic nerve neurodynamic tension technique, the MLSS, in healthy 239 and active adults. We observed significant mean increases in ipsilateral and contralateral PSLR 240 and PKB immediately and one hour post intervention, which is consistent with neurodynamic 241 tension being an important neuro-modulator of muscle extensibility, and is further supported 242 by the finding that these effects were significant after the first intervention session and 243 maintained for three weeks. As to the authors' knowledge lasting extra-segmental and 244 contralateral muscle extensibility gains from unilateral intervention have not previously been reported,^{24,32,36} these results require verification through additional studies. 245

The pooled mean increase in PSLR from pre first intervention to one hour post second intervention of $15\pm6^{\circ}$ represents a relative increase of $19\pm8\%$, utilising a total stretch time of 75 seconds per leg. This may be considered above average for PSLR gain in a hamstring

extensibility study,³⁵ but achieved with considerably less total stretch time than previously 249 reported.^{28,34} For example, Ayala and colleagues³⁴ demonstrated a mean increase of 14° in 250 251 PSLR utilising 540 seconds total weekly stretching over 12 weeks. Therefore the results of the 252 current study provide a novel finding in that neurodynamic tension and stretch intensity appear to have a highly significant role in muscle extensibility,^{18,30} compared to previous research 253 which has purported total weekly stretch time as the most important parameter.²⁷⁻²⁹ Thus MLSS 254 intervention could potentially be utilised to make stretching practices more efficient in 255 256 increasing hamstring extensibility by reducing total stretch time. However, further research is 257 required as the current study utilised a narrow sample of young and healthy adults, whereas 258 less robust populations, such as the elderly or those with irritable musculoskeletal conditions, 259 may not tolerate application of higher levels of stretch intensity and neurodynamic tension, and thus be inappropriate for MLSS intervention.^{26,36} Moreover, given the lack of blinding and 260 cross-over design of the current study, a follow-up investigation to verify and compare the 261 262 effects of MLSS intervention utilising single blinded RCT design is indicated.

263 Increased stretch tolerance from stretching is considered to occur through decreases in perception of stretch intensity for a common joint angle (SNRS Com) and potentially through 264 increased tolerance to higher intensity stretch sensation (SNRS Max).^{25,36} Consonant with the 265 266 post intervention ROM changes, significant mean decreases in SNRS Com for ipsilateral and contralateral PSLR and PKB are consistent with modulation of stretch tolerance through 267 268 neuronal desensitisation. Interestingly, PSLR but not PKB outcome measures demonstrated 269 small but significant concomitant increase in SNRS Max, suggesting modulation of muscle 270 extensibility by both neuronal desensitisation and increased tolerance of higher stretch intensity 271 segmentally, but not extra-segmentally. This may also be a novel finding, as previous research 272 has largely demonstrated constant maximal stretch intensity ratings pre-post stretching intervention.^{31,36,57} The contrasting result of the present study may be due to the MLSS being 273

a therapist-assisted technique eliciting greater amounts of neurodynamic elongation and stretch
 intensity.^{16,17,31,63}

Previous investigations of neurodynamics and muscle extensibility have reported 276 varying results. For example, Sullivan and colleagues⁶⁴ demonstrated focused hamstring 277 278 muscle stretches were more effective than hamstring stretches in a stooped position that was consistent with elongation of the neuraxis.^{16,63} However, the study by Sullivan and colleagues⁶⁴ 279 reported maintenance of ankle plantar flexion and adoption of a low to moderate stretch 280 281 intensity protocol, which may have elicited only neural unfolding, rather than nerve excursion, tension or strain,^{16,63} with the stooped stretch, and subsequently provided relatively less 282 stimulus to modulate stretch tolerance.^{18,32} Nevertheless, the current study adds to more recent 283 284 reports demonstrating efficacy of neurodynamic interventions in producing lasting increases of hamstring extensibility and stretch tolerance.^{18,32-33} 285

286 The MLSS produces elongation of the sciatic/tibial nerve tract through a combination of ankle dorsiflexion and eversion, knee extension, hip internal rotation and trunk flexion, with 287 likely resultant increases in nerve tension and strain.^{16-17,63,65} Its potential advantage over other 288 sciatic/tibial neurodynamic tension techniques, such as the slump²¹ and long sit slump,^{14,19} is 289 290 that it is postulated to produce maximal tolerated sciatic/tibial nerve tract elongation, with relatively less flexion stress on lower lumbar spinal segments⁶⁶ through antagonistic rotation 291 of the ilia around the sacrum in the hemi-sitting position.⁶⁷ Given unilateral sciatic-tibial sliding 292 has previously demonstrated not to produce contralateral hamstring extensibility effects,³² 293 294 while comparison between a bilateral glider and unilateral tensioner was statistically nonsignificant,¹⁸ further comparative studies of neurodynamic techniques, including the MLSS, on 295 muscle extensibility and stretch tolerance is indicated.³³ 296

297 An interesting *post-hoc* finding of the current study was the significant moderate to 298 strong inverse correlation between pre-intervention PSLR ROM and the magnitude of the

299 ROM increase immediately (r = -0.318; p < 0.05) and one hour (r = -0.526; p < 0.001) post 300 intervention, suggesting a potential 'diminishing returns' effect of the MLSS with respect to muscle extensibility. This is in contrast to the findings by Ayala and colleagues³⁴ who 301 302 demonstrated no significant difference between subjects with and without tight hamstring 303 tightness in response to 12 weeks of muscle stretching. Notwithstanding the large difference in 304 total stretch time, a possible explanation of these seemingly differing results, is that the 305 stretching protocol utilised by Ayala and colleagues,³⁴ through adoption of ankle dorsiflexion 306 in two out of the four techniques, appear a combination of stretches which preferentially target 307 muscle and neural tissue at moderate levels of stretch intensity whereas the MLSS preferentially targets the neural tissue at high stretch intensity.^{16,28,30,63} Although the PKB 308 309 measures in the current study were also significantly inversely correlated to pre-intervention 310 ROM, tight rectus-femoris was not an inclusion criterion so this effect may have been due some subjects achieving full PKB ROM. 311

312 The specific neuronal mechanisms responsible for modulating stretch tolerance are yet 313 to be fully elucidated. Large diameter proprioceptors have been implicated in modulating stretch tolerance through spinal gating,^{24,36} but this mechanism may not have a significant 314 lasting effect.⁴²⁻⁴³ Furthermore, as muscle spindle and golgi organ receptors are considered 315 316 absent outside the musculotendinous tissues,³⁸ and muscle stretching protocols have previously not demonstrated lasting extra-segmental nor contralateral effects, ^{24,32,36} this 317 suggests the effects of the MLSS were probably not modulated primarily by 318 proprioceptors.^{25,68,69} However, this postulation is not inconsistent with the possibility that 319 320 during stretching, low threshold proprioceptors and mechanoreceptors may sensitise high threshold receptors, such as mechanosensitive nociceptors, towards activation thresholds^{38,41,46} 321 322 through mechanisms such as the axon reflex and afferent convergence, as well as nonspecificity of peripheral afferent neuropeptides to fibre type.^{45,47} Conditioned learning and 323

324 increased activation of the DNIS have also previously been implicated in increases of muscle stretch tolerance,³⁶ and is not inconsistent with the results the current study. Compared to 325 previous muscle stretching research, the relatively higher levels of neurodynamic tension and 326 327 stretch intensity with MLSS intervention may have acted as a stronger neural stimulus for 328 subjects' learning to tolerate muscle stretch, which could explain the novel extra-segmental 329 and contralateral effects. A future study utilising the MLSS which includes a muscle 330 extensibility and stretch tolerance outcome measure proximal to the lumbar and lumbosacral 331 plexus may provide further insights into the role of conditioned learning and DNIS activation, 332 versus more local neuronal signalling at the spinal level, but fully elucidating these mechanisms 333 may require corroboration with direct neurophysiological measures.

334 Desensitisation of mechanosensitive nociceptors has previously been implicated in 335 modulation of muscle stretch tolerance and is also consistent with the results of the current study.^{24,36} The extra-segmental and contralateral effects induced by the MLSS are also 336 337 consonant with the proposition that increased stretch tolerance may be a form of nociceptive LTD,^{43,55} akin to sensitisation as a form of LTP,^{42,44,53} through recruitment of latent neuronal 338 circuits.^{48,51,54} Interestingly, A-delta but not A-beta afferent stimulation has been demonstrated 339 to induce C-fibre LTD and de-potentiate LTP in the rat spinal dorsal horn, which provides a 340 plausible mechanism for future investigations of stretch tolerance modulation in humans.⁴³ 341

Additionally, the sympathetic nervous system (SNS) and autonomic balance may also have a significant role in modulating stretch tolerance as sympathetic efferent and afferent fibres are considered to constitute a substantial proportion of lower limb peripheral nerve⁷⁰⁻⁷² and co-utilise noradrenaline and substance P, which are strongly implicated in nociceptor sensitivity and neuronal recruitment.^{38,42,48,53,73} Moreover, SNS tracts possess complex anatomical and physiological configurations including multiple segments and bilateral midline crossing spinally. multi-segmental serial and parallel processing supra-spinally, and likely

rapid autocrine and paracrine autonomic signalling.⁷⁴⁻⁷⁷ Notwithstanding the aforementioned 349 350 potential role of the SNS modulating stretch tolerance through neuronal desensitisation, significantly higher SNRS ratings in session two compared to session one for most of the 351 352 outcome measures could be due to autonomic modulation of stretch tolerance through attenuation of 'threat' perception during stretch.⁷⁸ However, some contrasting findings, 353 354 predominantly for the PKB data, further supports a difference between segmental and extrasegmental stretch tolerance modulation, but the potential of type 2 error, due to small sample 355 356 sizes, should also be considered. Moreover, given modulation of autonomic balance is a primary mechanism proposed to underlie yoga efficacy⁷⁹ and the likely overlap between yoga 357 postures and neurodynamic tension positions,⁸⁰ further investigation of the role of the 358 359 autonomic nervous system and its role in muscle extensibility, neurodynamics and HSI, is warranted.81 360

There were several limitations to the current study. Although there is in-vivo evidence 361 demonstrating validity in administering targeted nerve excursion and strain through 362 neurodynamics,^{16,82} there is an absence of studies which demonstrate differentiation between 363 364 muscle and nerve biomechanics with neurodynamic intervention, obviating a need for further research to improve content and construct validity.⁸³ Another major limitation of the current 365 366 study, due to resource limitations at MSc study level, was that all measurements and 367 intervention were performed by the same experienced musculoskeletal physiotherapist, raising the internal bias of the study.⁸⁴ Therefore verification of the study's results in a single blinded 368 369 RCT is indicated. Another limitation was that the PKB procedure utilised has not been 370 validated for rectus-femoris muscle extensibility, despite common clinical utilisation. 371 Nevertheless, the high consonance between mean PKB ROM and SNRS changes suggests high 372 measurement error was probably not a significant factor. Given the PKB procedure is simple 373 and efficient for a single examiner, future investigation of its validity is warranted. An

374 additional potential source of bias was not testing SNRS Com measures when post intervention 375 ROM was less than pre-intervention, which avoided moving the limb beyond the maximally 376 tolerated point. However, this only occurred with PSLR measures in one subject in the first 377 intervention session, and with several PKB measures in subjects who had full PKB ROM, and 378 is not considered to have significantly affected the results. Lastly, the study was limited to 379 healthy and active adults with clinically tight hamstrings recruited from a university population, 380 resulting in a relatively young and robust sample. Notwithstanding due care required in 381 applying neurodynamic tension techniques in less robust populations, investigation of the effects of the MLSS in a slightly older sample, or those with past HSI, is indicated.¹⁶ 382

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384 CONCLUSIONS

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Application of a novel sciatic-tibial neurodynamic tension technique, the MLSS, produced significant and lasting segmental, extra-segmental and contralateral increases of muscle extensibility and stretch tolerance in a healthy, active sample with clinically tight hamstrings. Additional studies are indicated to verify the findings and further investigate potential MLSS effects in different samples.

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TABLE 1. Mean stretch intensity ratings on a simple numerical rating scale (SNRS) from 0 ('no muscle stretch') to 10 ('the worst muscle stretch imaginable'). 'Com' represents the score taken at the pre-intervention joint angle for that session, whereas 'Max' represents the score at maximal stretch tolerance for each time-point. Effect of time: * p<0.05, ** p<0.01, *** p<0.001 compared to pre within the session; effect of side: †† p<0.01 compared to ipsilateral side; effect of session: # p<0.05, ### p<0.001 compared to session 1. Values shown are mean±SD.

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		Session 1			Session 2		
		Pre	Post	Post 1 hour	Pre	Post	Post 1 hour
	Com	7.4±0.8	5.1±1.4***	5.4±1.5***	8.1±0.9###	6.2±1.0***###	6.9±1.3***###
Ipsilateral PSLK	Max		7.9±1.0**	8.0±1.2**		8.7±0.6**###	9.0±0.8** ^{###}
	Com	7.8±0.8 [†]	6.3±0.9**††	5.4±1.4**††	8.4±1.1 ^{†###}	7.1±0.9**††	7.3±1.1**††
Contralateral PSLR	Max		7.5±0.7	8.0±0.9		8.6±0.7###	8.7±0.9###
	Com	70.11	5.8±1.8***	5.6±1.7***	7.6±1.2	5.6±1.8***#	6.4±1.6***#
Ipsilateral PKB	Max	/.2±1.1	7.2±1.4	7.4±1.4		7.2±1.5	7.6±1.3
	Com	7.1±1.6	6.0±1.7***	5.4±1.6***	7.8±1.0	6.6±1.4***#	6.5±1.7***#
Contralateral PKB	Max		7.3±1.4	7.2±1.6		7.7±1.4	7.6±1.7

Figure 1. During session 1, half the subjects received the MLSS intervention on the stance leg and the other half of the subjects received the intervention on the skill leg. Measurements were taken pre-, directly post, and one hour post-intervention. Following a 3-week washout period the intervention was repeated on the other leg.

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Figure 2. Modified long sit slump (MLSS). Start position (top row) and end position (bottom row). The subject starts hemi-sitting with the stretched limb on the plinth and the knee flexed. The subject uses their opposite hand to reach forward and hold the lateral border of the foot, placing it in dorsiflexion and eversion. They are then instructed to extend the knee and internally rotate the femur. The therapist assists to facilitate neurodynamic tension positions, and if the position is well tolerated, the subject is facilitated to add further trunk and cervical flexion.

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Figure 3: Effect of the MLSS intervention on: A) passive straight leg raise (PSLR), and B) prone knee bend (PKB). The intervention was performed on either the stance leg (n=6) or skill leg (n=7) in session 1, and on the other leg 3 weeks later in a counterbalanced manner. Main effects for PSLR: time p<0.001, side p<0.001, session p<0.001. Main effects for PKB: time p<0.001, side NS, session p<0.001.





CONSORT 2010 checklist of information to include when reporting a randomised trial*

	Item		Reported
Section/Topic	No	Checklist item	on page No
Title and abstract			
	1a	Identification as a randomised trial in the title	1
	1b	Structured summary of trial design, methods, results, and conclusions (for specific guidance see CONSORT for abstracts)	3-4
Introduction			
Background and	2a	Scientific background and explanation of rationale	5=7,
objectives	2b	Specific objectives or hypotheses	
			8
Methods			
Trial design	3a	Description of trial design (such as parallel, factorial) including allocation ratio	8
	3b	Important changes to methods after trial commencement (such as eligibility criteria), with reasons	N/A
Participants	4a	Eligibility criteria for participants	8-9
	4b	Settings and locations where the data were collected	8-9
Interventions	5	The interventions for each group with sufficient details to allow replication, including how and when they were	
		actually administered	11, Figure 2
Outcomes	6a	Completely defined pre-specified primary and secondary outcome measures, including how and when they	
		were assessed	9-10
	6b	Any changes to trial outcomes after the trial commenced, with reasons	N/A
Sample size	7a	How sample size was determined	8
	7b	When applicable, explanation of any interim analyses and stopping guidelines	N/A
Randomisation:			
Sequence	8a	Method used to generate the random allocation sequence	N/A 8
generation			(counterbalan
			ced)

	8b	Type of randomisation; details of any restriction (such as blocking and block size)	N/A 8 (counterbalan ced)
Allocation concealment mechanism	9	Mechanism used to implement the random allocation sequence (such as sequentially numbered containers), describing any steps taken to conceal the sequence until interventions were assigned	N/A 8 counterbalanc ed N/A
Implementation	10	Who generated the random allocation sequence, who enrolled participants, and who assigned participants to interventions	N/A
Blinding	11a	If done, who was blinded after assignment to interventions (for example, participants, care providers, those assessing outcomes) and how	N/A
	11b	If relevant, description of the similarity of interventions	N/A
Statistical methods	12a	Statistical methods used to compare groups for primary and secondary outcomes	11=12
	12b	Methods for additional analyses, such as subgroup analyses and adjusted analyses	11=12
Results			
Participant flow (a diagram is strongly	13a	For each group, the numbers of participants who were randomly assigned, received intended treatment, and were analysed for the primary outcome	Figure 1, 35
recommended)	13b	For each group, losses and exclusions after randomisation, together with reasons	35
Recruitment	14a	Dates defining the periods of recruitment and follow-up	8
	14b	Why the trial ended or was stopped	N/A
Baseline data	15	A table showing baseline demographic and clinical characteristics for each group	8-9 (Participants section in text)
Numbers analysed	16	For each group, number of participants (denominator) included in each analysis and whether the analysis was	/
-		by original assigned groups	Figure 1, 35
Outcomes and estimation	17a	For each primary and secondary outcome, results for each group, and the estimated effect size and its precision (such as 95% confidence interval)	8,12-13, Figure3
	17b	For binary outcomes, presentation of both absolute and relative effect sizes is recommended	N/A

Ancillary analyses	18	Results of any other analyses performed, including subgroup analyses and adjusted analyses, distinguishing pre-specified from exploratory	12-13
Harms	19	All important harms or unintended effects in each group (for specific guidance see CONSORT for harms)	N/A 35 (see flowchart)
Discussion			
Limitations	20	Trial limitations, addressing sources of potential bias, imprecision, and, if relevant, multiplicity of analyses	18
Generalisability	21	Generalisability (external validity, applicability) of the trial findings	18
Interpretation	22	Interpretation consistent with results, balancing benefits and harms, and considering other relevant evidence	13-18
Other information			
Registration	23	Registration number and name of trial registry	N/A Not a
			clinical trial
Protocol	24	Where the full trial protocol can be accessed, if available	N/A
Funding	25	Sources of funding and other support (such as supply of drugs), role of funders	N/A page
			1disclosure

*We strongly recommend reading this statement in conjunction with the CONSORT 2010 Explanation and Elaboration for important clarifications on all the items. If relevant, we also recommend reading CONSORT extensions for cluster randomised trials, non-inferiority and equivalence trials, non-pharmacological treatments, herbal interventions, and pragmatic trials. Additional extensions are forthcoming: for those and for up to date references relevant to this checklist, see <u>www.consort-statement.org</u>.

TDieR

The TIDieR (Template for Intervention Description and Replication) Checklist*:

Template for Intervention Description and Replication

Information to include when describing an intervention and the location of the information

Item	Item	Where lo	Where located **		
number		Primary paper	Other [†] (details)		
		(page or appendix			
		number)			
	RRIFF NAME				
1.	Provide the name or a phrase that describes the intervention.	_1,3			

	WHY		
2.	Describe any rationale, theory, or goal of the elements essential to the intervention.	_3-7	
	WHAT		
3.	Materials: Describe any physical or informational materials used in the intervention, including those provided	N/A	
	to participants or used in intervention delivery or in training of intervention providers. Provide information on		
	where the materials can be accessed (e.g. online appendix, URL).		
4.	Procedures: Describe each of the procedures, activities, and/or processes used in the intervention, including any	9-11	
	enabling or support activities.		
	WHO PROVIDED		
5.	For each category of intervention provider (e.g. psychologist, nursing assistant), describe their expertise,	_18	
	background and any specific training given.		
	HOW		
6.	Describe the modes of delivery (e.g. face-to-face or by some other mechanism, such as internet or telephone) of	8-11	
	the intervention and whether it was provided individually or in a group.		
	WHERE		
7.	Describe the type(s) of location(s) where the intervention occurred, including any necessary infrastructure or	8-11	
	relevant features.		
	WHEN and HOW MUCH		
8.	Describe the number of times the intervention was delivered and over what period of time including the	_8-11	
	number of sessions, their schedule, and their duration, intensity or dose.		
	TAILORING		
9.	If the intervention was planned to be personalised, titrated or adapted, then describe what, why, when, and how.	_8-11	
	MODIFICATIONS		
		-	1

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10. [‡]	If the intervention was modified during the course of the study, describe the changes (what, why, when, and	N/A 8-11	
	how).		
	HOW WELL		
11.	Planned: If intervention adherence or fidelity was assessed, describe how and by whom, and if any strategies	35	
	were used to maintain or improve fidelity, describe them.		
12. [‡]	Actual: If intervention adherence or fidelity was assessed, describe the extent to which the intervention was	35	
	delivered as planned.		

** Authors - use N/A if an item is not applicable for the intervention being described. Reviewers – use '?' if information about the element is not reported/not sufficiently reported.

† If the information is not provided in the primary paper, give details of where this information is available. This may include locations such as a published protocol or other published papers (provide citation details) or a website (provide the URL).

[‡] If completing the TIDieR checklist for a protocol, these items are not relevant to the protocol and cannot be described until the study is complete.

* We strongly recommend using this checklist in conjunction with the TIDieR guide (see BMJ 2014;348:g1687) which contains an explanation and elaboration for each item.

* The focus of TIDieR is on reporting details of the intervention elements (and where relevant, comparison elements) of a study. Other elements and methodological features of studies are covered by other reporting statements and checklists and have not been duplicated as part of the TIDieR checklist. When a **randomised trial** is being reported, the TIDieR checklist should be used in conjunction with the CONSORT statement (see <u>www.consort-statement.org</u>) as an extension of **Item 5 of the CONSORT 2010 Statement**. When a **clinical trial protocol** is being reported, the TIDieR checklist should be used in conjunction with the SPIRIT statement as an extension of **Item 11 of the SPIRIT 2013 Statement** (see <u>www.spirit-statement.org</u>). For alternate study designs, TIDieR can be used in conjunction with the appropriate checklist for that study design (see <u>www.equator-network.org</u>).



CONSORT 2010 Flow Diagram –adapted for a within subjects experiment over two intervention sessions with a 3 week washout period

