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## Experimental knee-related pain enhances attentional interference on postural control

Suda, Eneida Yuri; Hirata, Rogerio Pessoto; Palsson, Thorvaldur; Vuillerme, Nicolas; Sacco, Isabel C N; Graven-Nielsen, Thomas

Published in: European Journal of Applied Physiology

DOI (link to publication from Publisher): 10.1007/s00421-019-04192-9

Publication date: 2019

Document Version Accepted author manuscript, peer reviewed version

Link to publication from Aalborg University

Citation for published version (APA): Suda, E. Y., Hirata, R. P., Palsson, T., Vuillerme, N., Sacco, I. C. N., & Graven-Nielsen, T. (2019). Experimental knee-related pain enhances attentional interference on postural control. European Journal of Applied Physiology, 119(9), 2053-2064. https://doi.org/10.1007/s00421-019-04192-9

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               REXPERIMENTAL KNEE-RELATED PAIN ENHANCES ATTENTIONAL INTERFERENCE ON POSTURAL CONTROL
                       Eneida Yuri Suda<sup>1</sup>, Rogerio Pessoto Hirata<sup>2*</sup>, Thorvaldur Palsson<sup>2</sup>, Nicolas Vuillerme<sup>3</sup>, Isabel C N Sacco<sup>1</sup>,
          11
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                                                                                                                               Thomas Graven-Nielsen<sup>4</sup>
   4 13
   _{5} ^{15} Laboratory of Biomechanics of Human Movement, Dept. Physical Therapy, Speech and Occupational
   16
6 17 Therapy, School of Medicine, University of Sao Paulo, Sao Paulo, Brazil
   7 18 Department of Health Science and Technology, SMI, Faculty of Medicine, Aalborg University, Aalborg,
   8\ \frac{19}{20} Denmark
   9 2 Juniv. Grenoble-Alpes, EA AGEIS, Grenoble, France & Institut Universitaire de France
 10 22 Center for Neuroplasticity and Pain (CNAP), SMI, Department of Health Science and Technology, Aalborg
\frac{23}{24} University, Denmark
12 25
\begin{array}{c} {\bf 26} \\ {\bf 67} \\ {\bf 77} \end{array} \\ {\bf 77} \\ {\bf 67} \\ {\bf 77} \\ {\bf 13} \\ {\bf 67} \\ {\bf 77} \\ {\bf 13} \\ {\bf 14} \\ {\bf 14} \\ {\bf 14} \\ {\bf 15} \\ {\bf 
14 2 Number of Pages: 24
15 2 Number of words: 50<u>0983</u>
3 0
16 3 Number of figures: 4
 17 3 Number of tables: 3
          3 3
Abstract: 249
19 36 Corresponding author:
          3 8 Associate Professor Rogerio Pessoto Hirata, Ph.D. Center for Sensory Motor Interaction, SMI, Department of Health Science and Technology, Aalborg University,
22 4 Aalborg, Denmark
23 <sup>4</sup> Department of Health Science and Technology
 24 <sup>4</sup>Aalborg University, Fredrik BajersVej 7D-3
25 4 9 220 Aalborg E, Denmark
26 4 ft-mail: rirata@hst.aau.dk
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28 4 Acknowledgement
         4 Center for Neuroplasticity and Pain (CNAP) is supported by the Danish National Research Foundation
          5 (DNRF121). The authors thank the State of São Paulo Research Foundation (FAPESP) for the Suda scholarship
31 51
52 (FAPESP 2017/15449-4, 2015/00214-6).
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      Abstract
^9_1 Rurpose: To quantify how postural stability is modified during experimental pain while performing different
34 \frac{11}{12} cognitively demanding tasks.
35 \frac{1}{M} Methods: Sixteen healthy young adults participated in the experiment. Pain was induced by intramuscular 14
^{1} mjection of hypertonic saline solution (1mL, 6%) in both vastus medialis and vastus lateralis muscles (0.9% ^{1}6
37 1 kotonic saline was used as control). The participants stood barefoot in tandem position for one minute on a
38 1 9 orce plate. Center of pressure (CoP) was recorded before and immediately after injections, while performing
39 21wo cognitive tasks: (i) counting forwards by adding one; (ii) counting backwards by subtracting three. CoP
\frac{22}{2} yariables – total area of displacement, velocity in anterior-posterior (AP-velocity) and medial-lateral (ML-
\frac{24}{25} elocity) directions, and CoP sample entropy in anterior-posterior and medial-lateral directions were
42 \frac{26}{27} displayed as the difference between the values obtained after and before each injection and compared
^{28} between tasks and injections.
44 3 Results: CoP total area (-84.5 ± 145.5 vs. 28.9 ± 78.5 cm²) and ML-velocity (-1.71 ± 2.61 vs. 0.98 ± 1.93 cm/s)
45 3 2 decreased after the painful injection vs. Control injection while counting forward (P < 0.05). CoP total area
46 3412.8 \pm 53.9 \text{ vs.} -84.5 \pm 145.5 \text{ cm}^2), ML-velocity (-0.34 \pm 1.92 vs. -1.71 \pm 2.61 cm/s) and AP-velocity (1.07 \pm
\frac{35}{36.35} vs. -0.39 ± 1.82 cm/s) increased while counting backwards vs. forwards after the painful injection (P <
49 \frac{39}{40} onclusion: Pain interfered with postural stability according to the type of cognitive task performed,
\begin{array}{cc} 41 \\ 42 \\ \end{array} \text{uggesting that pain may occupy cognitive resources, potentially resulting in poorer balance performance.} \\ \end{array}
51 43
52 4 Keywords: postural stability, center of pressure, attention, distraction, pain
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1 2 3 30 31 3 2o quantify the extent to which attention is associated with postural control. Decreases in postural sway 83 43 44 46 85 48 50 51 52 53 57

4

Introduction Controlling of upright posture requires a significant amount of attention to constantly gather  $11\atop1$  information from the body and the environment and to generate adapted and accurate muscle activation 67  $\frac{1}{1}$  for postural control (Morasso and Sanguineti 2002). Although the majority of postural control is regulated 68  $^{1}$  $\S$ ia automatic neural processes (Bronstein and Buckwell 1997), higher cortical centers are significantly 69~1 7 hvolved in processing sensory information to plan and execute the best motor strategy for postural control 70 1 Winter 1995). In daily life, postural control is challenging as several tasks simultaneously compete for the 71 2 pognitive resources available (Woollacott and Shumway-Cook 2002), limited by the capacity of higher  $\frac{22}{25}$  enters to process sensory information (Kahneman 1973). Therefore, sharing attentional resources may  $\frac{24}{25}$  ause impairments in the performance of daily living activities (Brauer et al. 2004). Evidence suggests that

 $\frac{26}{\text{a}\text{For example,}}\text{ competition for cognitive resources during tasks involving postural stability results in body }27$  $^{2\,\text{g}}$  tability being prioritized over secondary tasks (Liston et al. 2014). Dual tasks paradigms, where subjects perform an additional task during quiet-standing, are employed

3 4 while performing a secondary task compared with control conditions have been reported (Andersson et al. 36002; Pellecchia 2003) whereby focusing the attention on standing as still as possible increased postural 3 gway compared with conditions without similar instructions (Vuillerme and Nafati 2007). Altogether, these  $\frac{39}{40}$  results suggest that postural control demands attention (Woollacott and Shumway-Cook 2002) and that

82  $\frac{41}{5}$  simultaneous cognitive loading plays an important role in balance stability (Swan et al. 2007). Although detrimental effects of cognitive loading on postural sway during unperturbed standing are

45 more commonly reported for older adults and patients, studies using dual-task approaches in young and

4 7-entrol-subjects show controversial results (Huxhold et al. 2006; Fraizer and Mitra 2008). Young healthy

4 gubjects have probably more ability to allocate the attentional resources during upright standing without

1 2 3 gacrificing postural stability, showing that a system without impairments prioritizes postural stability when  $^9_{\mbox{\sc d}}$  dealing with dual-cognitive tasks (Siu and Woollacott 2007). 11 12 Evidence suggests that Subjects with pain demonstrate increased postural sway compared with  $\frac{1}{1}$  controls (Hirata et al. 2011). Among several  $\underline{\underline{A}}$  potential possible explanations for this finding, one hypothesis  $^{1}\$$  that the increased postural sway may relate to a disrupting effect of nociceptive stimuli on attention to 92 176ther simultaneous non-nociceptive tasks (Eccleston et al. 1999), underlining that processing of nociceptive 93 1 9 timuli is cognitively demanding (Veldhuijzen et al. 2006). Thus, the execution of cognitive tasks during pain 94 2 might interfere with postural control. Although previous studies have shown that patients with pain present  $\frac{22}{2}$  impaired balance while performing a secondary cognitive task in comparison to health subjects (Van Daele  $\frac{24}{26}$ t al. 2010; Larivière et al. 2013; Mazaheri et al. 2014; Sherafat et al. 2014; Etemadi et al. 2016; Levinger et  $^{26}_{27}$  al. 2016), it is not clear yet the isolate effect of pain in these conditions and comparisons, since in clinical  $^{27}$  $^{2}$   $^{8}$   $^{pain}$  populations, besides pain, other factors like reduced muscle strength, reduced flexibility and 3 degenerative changes at the affected segment also cause both stiffness and instability in patients suffering 3 <u>2rom chronic pain (Knoop et al. 2012)</u>. Therefore, further investigation of the interaction between pain, 101 3 4ognition and postural stability is warranted. This investigation is of particular interest for clinical practice 102 3 gince there are evidences that attention can be directed away from pain using some specific strategies (Van  $\frac{37}{38}$  yckeghem et al. 2018). If selective attention could be directed away from the painful stimulus and modify  $\frac{39}{40}$  the deleterious effect of muscle pain on postural control, these results could have important implications  $\frac{41}{100}$  for clinical settings. Likewise, if the execution of cognitive tasks impairs postural control in the presence of  $\frac{41}{42}$ 106  $^4$  ain, this should also be taken into account in rehabilitation context. 107 45 Considering that posture can be defined as the dynamic stability of a continuous moving body 108 4 Tharbourne and Stergiou 2003; Madeleine et al. 2011), nonlinear analysis of the dynamic structure of the 4 genter of pressure (CoP) time series would contribute to understand the physiological complexity of posture 109  $\frac{1}{5}$  by accessing motor patterns that would be implicit in the CoP variability. Sample entropy (SaEn) measures 52 53

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2 gariations in the system output along time, which is independent of the signal magnitude (Slifkin and Newell  $1^{\frac{1}{999}$ ; Richman and Moorman 2000).\_\_Therefore, measures of physiological complexity of the postural sway  $\frac{11}{2}$  during quiet standing may relate to the system functionality as they are defined as the capacity of generating  $^{13}$  adaptive answers to an ever-changing environment such as controlling posture (Manor et al. 2010). SaEn  $^{14}$ 115  $\frac{1}{9}$  rovides a measure of "orderly structure" within the time series since it tests if there are any repeated 16 116~1  $\beta$  atterns of various lengths, including the ones that are not repeated at regular intervals (Duarte and Sternad 117 1 9008). So, the lower the SaEn values are, the higher the similarity and lesser the complexity in the temporal 11 2 peries is (Richman and Moorman 2000). SaEn has been used to measure the structure of the CoP variability  $\frac{1}{2}$  Roerdink et al. 2006; Donker et al. 2007; Duarte and Sternad 2008; Stins et al. 2009) and thus address the 119 24 25 complexity of the signal. 120 26 121 Most definitions of complexity are driven by operational considerations on the number of system 27 281 122 <del>ients and their functional interactions. Therefore, c</del>Complexity depends on the number of structural 29 3 components of the system, the existing coupling among these components and how this interaction is 123 3 2nfluenced by the intrinsic dynamic properties of the system and the motor task demands (Vaillancourt and 125 3 Newell 2002). Thus, if the presence of pain and the execution of a cognitive task are both concurring with  $_{
m 3}$  the attentional resources used in postural control, then the coupling between the components of the system gesponsible for balance may be affected and, consequently, the complexity of the postural sway is affected.  $\frac{39}{40} \frac{1}{10} \frac{1}{10}$  $^4$  tomplexity of the postural sway, and this increase has been attributed to a more automatized postural sway,  $^42$ 129 43 when less attention is directed to the balance control (Donker et al. 2007; Stins et al. 2009; Kuczyński et al. 130 131 45011). On the other hand, there is some evidence that the complexity of postural control decreases with 132 4 pain. Søndergaard et al. (2010) found a decrease SaEn of CoP displacement during sitting with increased 4 perceived discomfort in healthy young subjects (Søndergaard et al. 2010). The same Similar finding was 138 134 5 Teported in young subjects with transient acute episode of low back pain during two continuous hours of 52 53

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1 2 gtanding, but without history of low back pain (Fewster et al. 2017), showing a relation between the  $\frac{9}{100}$  courrence of pain and the decrease in CoP complexity. Therefore, examining the complexity of postural  $\frac{11}{12}$  way in a dual task context and the effect of experimental pain in this condition may improve the 13 $\beta$  1 understanding of the decrease in postural stability (Levinger et al. 2016) -and complexity (Fewster et al. 2017) 14 139  $^{1}$  hat may exist as a result of pain in an otherwise healthy system. 140 17 The aim of this study was to quantify how postural stability, i. e., CoP sway [{CoP sway velocity and 14 1 grea of displacement) and CoP complexity (CoP SaEn)], is modified during experimental pain while 142 2 performing a cognitive task. It was hypothesized that (i) the kind of cognitive task (more or less demanding)  $\frac{22}{2}$  143  $\frac{2}{3}$  n a non-painful condition will not interfere with CoP sway or CoP complexity, since the system would have  $\begin{array}{c} 24 \\ 144 \\ 2 \end{array}$  enough cognitive resources to overcome it; (ii) experimental pain will increase CoP sway and decrease CoP 145  $\frac{26}{27}$  complexity, regardless the type of cognitive task performed; (iii) the presence of experimental pain while  $^{28}\text{performing a difficult cognitive task will overload the cognitive resources and impair postural stability, <math display="inline">^{29}$ 147 <sup>3</sup> Chcreasing CoP sway and decreasing CoP complexity. 148 3**2.** Methods 149 34.1. Subjects <sup>150</sup> 36 Sixteen young adults, all university students, (to control for the effect of education level on  $\frac{37}{38}$  multitasking performance (Voos et al. 2015)), participated in the experiment – 8 males (mean ± SD: age = 152  $\frac{39}{40}$ 6.9 ± 2.8 years; body mass = 74.9 ± 13.8 kg; height = 1.76 ± 0.08 m) and 8 females (mean ± SD: age = 27.1 ± 153  $\frac{41}{4.0}$  years; body mass = 68.8 ± 5.2 kg; height = 1.68 ± 0.06 m). The exclusion criteria were body mass index  $\frac{41}{4.0}$  $^{4}$ above 25 kg/m², pregnancy, drug addiction, previous neurologic, musculoskeletal or mental illness, lack of 155 45 bility to cooperate, current use of medications (e.g. analgesics, anti-inflammatory medicine), consumption 156 4 of alcohol, caffeine, nicotine or painkillers 8 hours prior to the data collection, recent history of acute pain 15 4 affecting the upper lower limb and/or trunk, past history of chronic pain conditions, participation in other  $\frac{1}{5}$  pain trials throughout the study period. All procedures performed in studies involving human participants

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8 were in accordance with the ethical standards of the institutional and/or national research committee and 160 1 with the 1964 Helsinki declaration and its later amendments or comparable ethical standards. The study was 11 approved by the local Ethics Committee (N-20120077). This sample size was calculated to detect a minimum 162 difference of 40% in the CoP area assuming type error 1 as 5% and power of 80% between the conditions 14 before and after the induction of experimental pain. All participants gave signed informed consents prior to 16 17 inclusion in the study.

# 165 $\frac{19}{20}$ .2. Experimental protocol

166 21 22 Since in healthy individuals approximately 70% of the information used for controlling posture  $^{23}$  originates from proprioceptive systems (Peterka 2003), we controlled the effect of different footwear on  $^{24}$ 168 2 postural control by asking the subjects to stand barefoot during the experiment. -The participants stood on 26 169 27 triangular force plate that measures vertical forces (Good Balance System, Metitur, Jyväsklä, Finland; 170 2 glimensions: equilateral triangle - 800-mm; sampling frequency: 50-Hz as suggested by the International 171 3 pociety for Posture and Gait Research Standardization Committee (Scoppa et al. 2013)). This is a valid and  $\frac{32}{3}$  geliable system for postural sway measurements (Era et al. 2006; Ha et al. 2014) with accuracy better than  $\frac{34}{35}$ -mm for the CoP position measurement (Good Balance System User Manual). The CoP position was 174 3 calculated via the Good Balance Software (Metitur, Jyväsklä, Finland) which uses the weighted arithmetic  $^{3}$  nean between the vertical force measured by four sensors and their corresponding position: one in each  $^{176}$   $^{4}$  Qorner of the force-plate and the last one in the centroid of the force-plate (Fig. 1). The rational for using the 177 4 22 andem position for the feet was based in previous studies showing that greater pain effects are presented 178 4 4 when posture is challenged (Hirata et al. 2013). This was important to ensure that postural stability 4 daptations due to pain could be observed.- Therefore, subjects were asked to stand in tandem position, to  $\frac{47}{48}$  increase postural challenge during the tasks, with the right leg behind (Fig. 1), arms hanging relaxed  $\frac{49}{50}$  alongside the body, and were instructed to maintain balance while looking forward. Tape markers were

 $\frac{5}{5}$  placed on the force plate to ensure that the same foot position was maintained through all conditions. During  $\frac{5}{5}$ 

the assessment of postural control, subjects were instructed to look forward at a target positioned at eye- $^{9}$  184  $^{1}$  level approximately 45-cm from the subjects to minimize the influence of the target distance on postural  $\frac{11}{2}$  way (Kapoula and Lê 2006). CoP records were made under eight experimental conditions, depending on the 186  $\frac{13}{14}$  type of injection (control or painful), the dual-task (counting forward or counting backward as the less and 187  $\frac{15}{2}$  more challenging tasks, respectively), before (pre-injection) and immediately after the injection. The  $188\,$   $1\,$ 7ounting forward task consisted of adding one and the counting backward was performed by subtracting 189 1 hree, beginning from a random number. The total number of answers and the number of correct answers

190 2 during each trial were recorded. The order of the injections and the order of the tasks were randomized,

 $\frac{22}{23}$  with the same number of subjects receiving the hypertonic or isotonic injections first.

The experiment always followed the same order for all participants: (i) CoP measurement while 193  $\frac{26}{27}$  performing the first randomly assigned task (cognitive task 1 or 2) over 60-s (pre-injection 1); (ii) 1-min rest; 194  $\frac{2}{2}$  (iii) CoP measurement over 60-s while performing the second randomly assigned task (cognitive task 1 or 2) 195 <sup>3</sup> Ever 60-s (pre-injection 2); (iv) injections of the first saline solution (painful or control) into vastus medialis

196 3 (VM) and vastus lateralis (VL) muscles; (v) assessment of pain intensity by visual analogue scale (VAS); (vi)

197 3 CoP measurement over 60-s while performing task A; (vii) collecting VAS scores of the pain intensity and 1-

198 3 Anin rest; (viii) CoP measurement over 60-s while performing task B; (ix) collecting VAS scores of the pain

 $\frac{37}{38}$  intensity. After the final step, the pain VAS scores were taken each minute until the pain had subsided which

 $\frac{39}{40}$  was followed by a 5-min break. Following the break, all steps of the experiment were performed again with

 $^{20}$ l  $^{4}$ the injection of the other saline solution, including new pre-injection CoP recordings. Before each CoP  $^{4}$ 2

202 4 measurement, all subjects confirmed that no tiredness or other problems were presented. The duration of

203 45 he CoP measurements were performed according to guidelines proposed by the International Society for

204 4 Posture and Gait Research (Scoppa et al. 2013). Fig. 2 summarizes the study procedures along time.

## Experimental muscle pain

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Before the experiment all subjects were instructed about the nature and effects of the injections,

1 2 and that one type of injection would be painful while the other would be a non-painful stimulus, although  $\frac{9}{1}$  they would not know which kind of injection they would be receiving. Pain was induced through  $\frac{11}{12}$  intramuscular injection of 1-mL of 6% sterile hypertonic saline solution or as a control condition 1-mL of 21) 1 sotonic (0.9%) saline solution (Graven-Nielsen et al. 1997; Farina 2003; Schulte et al. 2004; Falla et al. 2006)  $^{211}$   $^{15}$ he injections were performed with a 2-mL syringe with a disposable needle (27G, 40-mm) into right VM 212  $\,^1$ 7nuscle and right VL muscle. Both injections locations were marked to ensure that they were applied 213 1 Approximately in the same location. The VM muscle injection was performed 5-cm proximal and 5-cm medial 214 2 to the medial corner of the patella (Shiozawa et al. 2013), and in the VL muscle, injections were performed  $\frac{22}{23}$ t two thirds of the distance from the anterior spina iliaca to the lateral side of the patella (Fig. 3). The depth  $^{24}_{26}$  of the injection was determined by an ultrasound scanner (LOGIQ $^{\text{m}}$  S7, General Electric, USA). This pain  $\frac{26}{27}$  model has been successfully used previously to mimic knee-related pain during quiet standing tasks  $^{28}$  providing moderate pain intensities for approximately five minutes (Hirata et al. 2011). Hypertonic saline  $^{29}$ 219 3 hjections have been shown to activate nociceptors around the injected site (Mense 1993) whereas the 0.9% 220 3 Eotonic saline injections have induced little or no pain during postural control tasks similar to the one used 221 3 4n the present study (Hirata et al. 2010, 2011, 2013). 222 38.4. Assessment of pain intensity 223 38 The subjects were asked to rate the pain intensity using a 10-cm VAS from 0-cm to 10-cm (0-cm 224  $\frac{39}{40}$  means "no pain" and 10-cm means "maximum pain") immediately after the injections and after each balance  $^{225}$   $^{41}_{\text{measurement}}$ . Therefore, three VAS scores were obtained for each set of experiments (balance  $^{42}$  $^{226}$  measurements after isotonic injection and balance measurements after hypertonic injection, respectively; 227 4 \$\frac{1}{3}\$ig. 2), and the mean values of the three VAS scores were considered as the pain intensity after each injection 228 4 paradigm. Additionally, following each set of experiments subjects were asked to indicate the overall pain 229 4 greas during the trials on a body chart and to respond the McGill Pain Questionnaire (Melzack 1975). The  $\frac{1}{5}$  area of pain was extracted from the body charts with VistaMetrix 1.38 software. The pain rating index based 52 53

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233  $\frac{1}{1}$  the total pain rating index were determined as the sum of the ranked values of the words (Melzack 1975). 234 13.5. 14

Data analysis

All variables for postural sway were calculated based on 50-s of the standing tasks, with the first and

on the rank values of the words chosen within each category (sensory, affective, evaluative and

 $\frac{9}{1}$  Miscellaneous) from McGill Pain Questionnaire were obtained and the score for each category, as well as

237 19oftware (Mathworks, Massachusetts, USA). The area fitted to 95% confidence interval of the CoP

238 2 plisplacement was calculated as representative of the total CoP area displacement (95% confidence interval

 $\frac{22}{2}$  gllipse), along with the CoP velocity in both directions (anterior-posterior and medial-lateral). The structural

 $\begin{array}{c} 24 \\ 24 \\ 25 \end{array}$  variability of the CoP was calculated by means of SaEn with the embedding dimension (m) and the tolerance

241  $\frac{26}{27}$  distance (r) set to m=2 and r=0.2xSD (Vaillancourt and Newell 2000). All CoP variables are displayed as the

 $^{28}$  difference between the values obtained immediately after the injection and the correspondent pre-injection

243 3 Condition. Negative values show that the CoP variable decreased after the injection of the saline solution

244 32ompared to its respective pre-injection condition. Likewise, positive values show that the CoP variable

245 3 increased after the injection compared to its respective pre-injection condition.

#### 246 3*8.6.* Statistical analysis

Pain outcomes were compared between injection types (isotonic or hypertonic injections) with

248  $\frac{39}{40}$  paired T-tests when normal distribution was present (VAS scores and pain area data) and with the Wilcoxon

 $^{41}$  Signed Rank Test when the data distribution was non-normal (McGill scores). The task measures (number of  $^{42}$ 

 $^{4}$  answers, number of correct answers) were evaluated with a 3-way RM-ANOVA with *injection* (isotonic vs

251 4 Sypertonic), time (pre-injection vs after injection) and task (counting forward vs backwards) as main factors.

252 4 The CoP parameters were compared with a 2-Way RM-ANOVA with task and injection as main factors, and

253 4 the p-values are shown in the table 3. Bonferroni post-hoc correction for multiple comparisons was applied

254  $\frac{1}{5}$  and p-values are shown in the results texts. The alfa-value ( $\alpha$ ) for statistical significance was set to 0.05.

1 2 3 12 255 Results -Experimental muscle pain and cognitive task performance Formatted: Justified, Indent: First line: 0.49' <u>Area and amplitude of perceived pain'</u> Formatted: Font: Italic Formatted: Indent: First line: 0" 13 Fig. 4 shows the reported pain areas following both isotonic and hypertonic injections. Pain was 258 Formatted: Font: 12 pt, Italic 259  $^{1}$ 5 present in the anterior and lateral portions of the thigh after both isotonic and hypertonic injections, being 260 1 More concentrated in the lower half of the thigh after the isotonic injections. The hypertonic saline injections 261 1 9nduced higher pain area (mean area ± SD: isotonic = 518.6 ± 690.6 au; hypertonic = 1659.3 ± 1574.0 au; 262 2P=0.003) and higher VAS scores (mean score  $\pm$  SD: isotonic = 0.9  $\pm$  1.1 cm; hypertonic = 4.7  $\pm$  1.7 cm; P<0.001)  $\frac{22}{2}$  than isotonic saline injections. Table 1 shows the scores for each class of words from McGill Pain  $\begin{array}{c} 24\\ 26 \end{array}$  Questionnaire and the pain rating index. Subjects presented a higher total pain rating index and scored  $\frac{26}{27}$  higher in all the categories, with the exception of the affective class, after the hypertonic injections (*P*<0.05). 266 Cognitive task performance Formatted: Indent: First line: 0" Formatted: Font: 12 pt. Italic Only for the analysis of the cognitive task performance, one subject was not included due to problems 267 Formatted: Font: Italic Formatted: Normal, Justified, No bullets or numbering 3 2n the answers recording. The total number of answers and the number of correct answers decreased during 268 3 packwards counting conditions compared with forwards counting despite the injection effect (significant 269 3 main effect for task factor; Table 2). 37 271 38 272 3.3 Center of pressure Formatted: Normal, Justified, No bullets or numbering Formatted: Font: 12 pt, Italic 41 273 Effect of experimental pain in CoP variables <sub>274</sub> 43 There were no statistical differences between the different conditions for the factor injection on any 275 45 of the CoP variables (Table 3). 46 Effect of cognitive task in CoP variables 276 47 48 49 50 53 57 58 59 60 61

<sub>280</sub> 13 <sub>281</sub> 15

283 1 9 oth variables decreased after the hypertonic injection in comparison to the condition with isotonic injection

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 $^{41}_{\rm D}$  performance of a difficult cognitive task increased CoP sway but did not change CoP complexity.

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 $^{9}$  278  $_{1}$  increased AP-velocity during the counting backwards task compared to the counting forwards task, 279  $\begin{array}{c} 11 \\ 12 \end{array}$  regardless the type of injection (Table 3).

Effect of the interaction between experimental pain and cognitive task in CoP variables An interaction effect was found between injection and task factors for CoP total area and CoP ML-

282  $1\sqrt[n]{e}$  locity (CoP total F=7.78, P=0.049; CoP ML F=4.69, P=0.021) (Table 3). Post-hoc comparisons showed that

284 2 when subjects where counting forward (Bonferroni: P = 0.010 for total area; P = 0.015 for ML-velocity). After  $\frac{22}{2}$  the hypertonic injection, CoP total area increased when subjects were counting backwards in comparison to

240 25 when they were counting forwards (Bonferroni: P = 0.019). ML-velocity showed differences between the 287  $\frac{26}{0}$  different cognitive tasks also after the injection of hypertonic solution, with a smaller decrease of ML-velocity

288  $^{2}$ 8 while counting backwards (Bonferroni: P = 0.049).

## Discussion

The present study aimed at quantifying how postural stability, represented by CoP sway (velocity and

A main effect of task was found for the CoP AP-velocity (F=5.82; P=0.028), showing that there was an

291 3 4 rea of displacement) and CoP complexity (CoP SaEn), is modified during experimental pain while performing

292 38 cognitive task. The main results showed that the kind of cognitive task did not interfere with postural

 $\frac{37}{38}$ tability in the absence of pain. Experimental pain around the knee joint reduced CoP sway but did not affect

 $\frac{39}{40}$  coP complexity during the performance of an easier cognitive task. During experimentally induced pain, the

# 296 <sup>4</sup> <u>Pain intensity and counting performance</u>

The subjects showed higher pain intensity for the hypertonic saline injection and a larger pain area

298 4 Tompared with the isotonic saline injection, as expected, indicating that experimental pain occurred (Hirata

299 4 oft al. 2011). The McGill pain questionnaire indicated that hypertonic saline was perceived more impairing

4 shan the isotonic injection in all subscales except for the affective one. It is important to note that during
 301 1 sotonic injections subjects rated pain around 1/10, which cannot be classified as a totally pain free condition.

Counting performance requires the use of cognitive process which relies on the working memory of

 $^{13}$  the subject (Lemaire 1996), impairing motor output performance when executed simultaneously with a  $^{14}$ 

 $^{15}$ motor task (Vuillerme and Nafati 2007). Seminowicz and Davis (2007) showed that subjects are able to

17 maintain performance of difficult cognitive task while experiencing different levels of pain. In this study, the

306 1 painful condition did not affect the counting performance while performing a motor task (standing still)

 $\frac{1}{2}$  indicating that healthy subjects are able to engage multiple tasks (motor and cognitive) during pain without

 $\frac{22}{25}$  ompromising performance. This suggests that sufficient cognitive resources were available to manage the

 $\frac{24}{25}$  cognitive process of counting forwards or backwards despite the interpretation of painful stimuli and the

 $\frac{26}{27}$  postural control task (Eccleston et al. 1999). Finally, education level is associate with both motor and

 $^{28}$  perceptual performance, where higher education level is associated with better performance (Voos et al.  $^{29}$ 

30015). -Since our subjects were all university students, we believe that bias due to education level did not 31

313 32 affect the present results.

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#### 314 3 Leffect of cognitive tasks on postural stability

Our first initial hypothesis, that (i) the kind of cognitive task (more or less demanding) in a non-painful

 $\frac{37}{38}$  ondition would not interfere with CoP sway or CoP complexity, was confirmed. The factor task affected the

 $\frac{39}{40}$  coP anterior-posterior velocity, indicating an increased velocity during the execution of the more difficult

 $\frac{41}{42}$  task (counting backwards) in comparison to the easier task (counting backwards forward). Nevertheless, the  $\frac{41}{42}$ 

319 4 cop SaEn was not affected by the kind of the performed cognitive task. These results indicate that enough

320 45 ognitive resources were available to overcome the demands of both cognitive and postural tasks, which 46

321 4 Was expected since they were young individuals without any sensory-motor alterations.

# 322 4 fffect of experimental knee-related pain on postural stability

2  $\frac{1}{5}$  Reducing postural sway might reflect a motor strategy available for healthy subjects to avoid excessive 52 53

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Our second initial hypothesis, that (ii) experimental pain would increase CoP sway and decrease CoP  $\frac{9}{1.6}$  omplexity was not confirmed since the type of saline solution injected did not affect the CoP variables.  $\frac{1}{12}$  However, even though the factor *injection* did not show statistical differences between the different  $\frac{1}{2}$  conditions for any of the studied CoP variables, there was a difference between total area and ML-velocity 327  $^{1}8$ etween the control and the painful condition when the subjects were counting forwards, i.e., in conditions 328  $\,1\,$ Where the kind of cognitive task performed was the same. Interestingly, during the counting forward, the 329 1 \$ype of injection resulted significant changes in postural sway (total area and ML-velocity) in opposite 330 2 directions: positive values of the difference between pre-injection and after injection of the isotonic solution,  $\frac{22}{2}$  whereas after the injection of the hypertonic solution both variables showed negative values. Additionally,  $\begin{array}{c} 24\\ 25 \\ \end{array}$  significant changes were observed in the structural variability of the CoP signal. This is contrary to the 333  $\frac{26}{27}$  initial hypothesis, where an increase in postural sway and a decrease in structural variability during painful  $\frac{28}{2}$  conditions were expected. It is also in contrast with previous findings (Mazaheri et al. 2013) but may relate 338 othe different position of the feet used in this study, which affects the postural sway (Day et al. 1993). The 336 322andem feet position adopted allows less displacement of the CoP due to the limited base of support 337 3 4 ompared to side-by-side feet position, since if the subjects increase the CoP amplitude they may fall (Day 338 3 et al. 1993). This also may reflect a voluntary strategy, requiring a greater amount of cognitive resources and  $\frac{37}{38}$  ttention (Morasso and Sanguineti 2002), attempting to avoid large excursions of the body and consequent  $\frac{39}{40}$  loss of balance. For the current study, this might indicate that the subjects prioritized the balance task over  $^{4}$  the other tasks, also known as *posture first strategy* (Vuillerme and Nafati 2007). The subjects were able to  $^{4}$  $^{4}$  educe the postural sway without compromising the counting performance during the easy cognitive task, 343 45 uggesting that the available cognitive resource was sufficient to perform the less challenging cognitive task 344 4 Without compromising postural stability. Therefore, these results indicate that healthy subjects have the 345 4 gapacity to perform easy cognitive tasks while ensuring postural stability (Siu and Woollacott 2007).

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1 during the control injection while counting backwards, probably indicating that a high cognitive load seems  $\frac{1}{1}$  to be interpreted as a treat to postural stability. An alternative explanation for the contrast between the  $\overset{1}{\overset{\circ}{0}}$  present study and the previous studies with pain patients showing larger postural sway (Schulte et al. 2004;  $\overset{1}{\overset{\circ}{0}}$ 351  $^{1}$   $^{1}$  Eevinger et al. 2016) might be the pain model used that is not a complete proxy to the impaired pain patients' 352 17 ensory-motor system. 353 1 Interactions between pain and cognitive load on postural stability 354 21 Our initial third hypothesis, that (iii) the presence of experimental pain would increase CoP sway and

aranslation of the body, which could lead to balance loss (Winter 1995). This strategy was also observed

 $\frac{22}{2}$  decrease CoP complexity only when performing a difficult cognitive task was partially confirmed since CoP  $\frac{24}{25}$  way increased during pain under a difficult cognitive task, but the CoP complexity did not change. ANOVA  $\frac{26}{27}$  results showed an interaction between the task and injection factors for total area and ML-velocity. After  $^{2}$  the hypertonic injection CoP total area increased and CoP ML-velocity decreased less while counting 359 3 backwards in comparison to counting forwards condition, corroborating our hypothesis. ANOVA results also 360 32 showed an effect of the task factor on AP-velocity with post-hoc comparisons showing a difference only 361 3 4 during the hypertonic injection condition: while counting backwards AP-velocity also increased. Altogether 362 3 these results show that CoP sway increases when performing a more demanding cognitive task in the  $\frac{3}{3}$  presence of experimental pain. This might reflect an interference with the information-processing capacity  $\frac{39}{40}$  and an attention disruption from both postural control and cognitive task (Eccleston et al. 1999). Previous  $^{41}$  Studies suggest that disruptions of sensory information lead to worsening of proprioception in the affected  $^{42}$  $^{4}$  area (Matre et al. 2002), further impairing postural sway (Hirata et al. 2010, 2011). The results indicate that 367 45he posture first strategy (Vuillerme and Nafati 2007) found during the easy cognitive task during pain is no

368 4 Tronger feasible when a difficult cognitive task is performed during painful conditions. The increased cognitive 4 goad in painful conditions seems to impair the motor performance maybe due to insufficient cognitive  $\frac{1}{5}$  resource to simultaneously maintain postural stability (which requires significant amount of attention

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17 Morasso and Sanguineti 2002)) and execute a difficult cognitive task. These results might have important  $\frac{9}{10}$  flow implications in understanding the mechanisms related to fall accidents. Postural stability in daily life  $\frac{11}{12}$  activities is usually performed in combination with additional tasks, for example, walking in a busy slippery  $\frac{13}{3}$  idewalk. These daily life activities involves simultaneously competition for the cognitive resources available  $\frac{13}{4}$ 375  $^{1}$  Woollacott and Shumway-Cook 2002) to evaluate the environment constrains in order to promote the best  $_{16}$ 376  $\,1\,$ 7motor strategy (Winter 1995). Our present results indicate that, if the subject performs a challenging 377 1 postural task in pain, his/her capacity for maintain balance while exposed to a difficult cognitive task is 378 2 \*\*puboptimal, which could increase the likelihood of losing balance. The complexity of postural sway did not show any differences between the experimental conditions.  $\frac{24}{25}$  his result is contrary to the literature finding that young healthy subjects present a more regular and less

 $\frac{26}{27}$  automatic postural sway (decreased CoP SaEn) when the motor task is more difficult (e. g. standing with eyes  $^{28}$  closed) and more irregular postural sway and more automatic postural sway (increased CoP SaEn) when a 383 3 Qognitive task is added (Donker et al. 2007; Stins et al. 2009). The fact that the cognitive task did not interfere 384 3 2with CoP complexity may be due to the nature of both motor (standing in tandem position) and cognitive 385 3 4subtraction calculus) tasks used in the experimental setup that did not interfere with the automaticity of 386 3 Rostural control. Besides that, pain also did not affect CoP complexity, showing that experimental knee- $\frac{37}{38}$  gelated pain did not compromise the coupling between the components of the system responsible for  $\frac{39}{40}$  balance in the current experimental setup. Future studies should investigate the interaction between pain,  $^{41}$ cognition and on CoP complexity with different motor and cognitive demands, in addition to different  $^{42}$ 390 <sup>4</sup> populations.

392 4 Televance of the findings for clinical populations should be interpreted with care. The experimental pain 393 4 model used here is convenient to assess the effect of pain without the interference of potential structural  $\frac{1}{5}$  pr pathologies. However, extrapolating the current findings to an older population can only be done to some

Despite interesting results regarding the effects of cognitive tasks in postural control during pain, the

degree. Additionally, chronic pain patients may also suffer from depressive symptoms (Bair et al. 2003) or 9  $_1$  anxiety (McWilliams et al. 2003), which might increase cognitive load (Nebes et al. 2001). Furthermore,

 $\frac{11}{12}$  cognitive impairments are often found in chronic pain patients, decreasing the possibility to maintain

 $^{1398}$  performance of two or more concurrent tasks (Brauer et al. 2004), as opposed to what was observed in this

 $\frac{1}{2}$  study where young healthy subjects were recruited. Also, there was no recording of postural sway without

400 17 any cognitive task. This would have allowed comparisons with a condition where neither pain nor cognitive

401 1 \$asks were influencing postural sway, and could have reduced type 2 errors given that multiple CoP variables

402 2 were analyzed in the study. Thus, it can be considered a limitation to our interpretations.

#### **Conclusions**

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Pain and cognitive task interfered on postural stability, changing its patterns. During the performance

40  $\frac{26}{0}$  of a simple cognitive task, pain, reduced postural sway, while during the performance of a more demanding

406 28 cognitive task, postural sway was increased in young healthy subjects. Since our subjects were young healthy

407 3 Qubjects, the direct translation of the present results to patients suffering from pain should be done with

3 <u>2aution</u>. However, <u>Tt</u>hese results may suggest that rehabilitation approaches should take into account that

3 ⊉ain not only affects directly the motor system, but may occupy cognitive resources, potentially resulting in

410 3 poorer performance when performing rehabilitation exercises. Additionally, rehabilitation strategies using

 $\frac{37}{38}$  both motor and cognitive resources need further investigation to outline the effect of interaction between

412  $\begin{array}{c} 39 \\ 40 \end{array}$  pain and cognition on the performance during activities of daily life in patients.

## 414 4 Compliance with ethical standards

415 4 **5** unding: Center for Neuroplasticity and Pain (CNAP) is supported by the Danish National Research

416 4 Foundation (DNRF121). The authors thank the State of São Paulo Research Foundation (FAPESP) for the Suda

417 4 gcholarship (FAPESP 2013/06123-7, 2015/00214-6).

418  $\frac{1}{5}$  **Conflict of Interest:** The authors declare that they have no conflict of interest.

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**Figure captions** <sup>570</sup> 10  $\frac{1}{12}$  ig 1 Schematic drawing representing the force platform size, sensor locations, and the tandem position of  $\frac{13}{14}$  the subjects during the experiment  $\frac{13}{14}$ <sub>573</sub> 15 574 1 Fig 2 Study design overview: pain assessments were performed immediately after each injection and each 575 1 \( \text{\text{9}}\) alance measurement; the order of the saline injections was randomized in a balanced way 576 21  $\frac{22}{250}$  3 Injections sites for vastus lateralis muscle, performed at two thirds of the distance from the anterior  $\frac{24}{25}$  pina iliaca (a) to the lateral side of the patella (b); and for the vastus medialis muscle, performed 5 cm  $\frac{26}{27}$  proximal and 5 cm medial to the medial corner of the patella (c), <sub>580</sub> 28 581 3 Fig 4 Representation of the experimental pain distribution reported areas after isotonic (top, blue in the 582 32nline version) and hypertonic (bottom, red in the online version saline injections (A); the individual 34 distributions are superimposed in the anatomical drawings <sup>584</sup> 36 

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       REXPERIMENTAL KNEE-RELATED PAIN ENHANCES ATTENTIONAL INTERFERENCE ON POSTURAL CONTROL
          Eneida Yuri Suda<sup>1</sup>, Rogerio Pessoto Hirata<sup>2*</sup>, Thorvaldur Palsson<sup>2</sup>, Nicolas Vuillerme<sup>3</sup>, Isabel C N Sacco<sup>1</sup>,
                                                      Thomas Graven-Nielsen<sup>4</sup>
<sub>588</sub> 13
589 ^{15} Laboratory of Biomechanics of Human Movement, Dept. Physical Therapy, Speech and Occupational
\frac{16}{17} Therapy, School of Medicine, University of Sao Paulo, Sao Paulo, Brazil
591 18 Department of Health Science and Technology, SMI, Faculty of Medicine, Aalborg University, Aalborg,
\begin{array}{c} 19 \\ 20 \end{array} \text{Denmark}
593 2 Juniv. Grenoble-Alpes, EA AGEIS, Grenoble, France & Institut Universitaire de France
594 22 Center for Neuroplasticity and Pain (CNAP), SMI, Department of Health Science and Technology, Aalborg
\begin{array}{c} 23 \\ 24 \end{array} \text{University, Denmark} \\ \end{array}
596 25
597 \frac{26}{27} original article for: European Journal of Applied Physiology
598 2 Number of Pages: 24
599 <sup>2</sup> Number of words: 5009
\begin{array}{c}
30 \\
600 \\
31
\end{array} Number of figures: 4
601 32Number of tables: 3
602 3 Abstract: 249
35
603 3 *Corresponding author:

    38 Associate Professor Rogerio Pessoto Hirata, Ph.D.
    605 Center for Sensory Motor Interaction, SMI, Department of Health Science and Technology, Aalborg University,

606 4 Aalborg, Denmark
607 4 Department of Health Science and Technology
608 <sup>4</sup> Aalborg University, Fredrik BajersVej 7D-3
609 4 § 220 Aalborg E, Denmark
610 4 \ -mail: rirata@hst.aau.dk
611 45
612 4 Acknowledgement
613 4 Center for Neuroplasticity and Pain (CNAP) is supported by the Danish National Research Foundation
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614 5 (DNRF121). The authors thank the State of São Paulo Research Foundation (FAPESP) for the Suda scholarship
615 5 FAPESP 2017/15449-4, 2015/00214-6).
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       Abstract
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617 <sub>1 Purpose</sub>: To quantify how postural stability is modified during experimental pain while performing different
618 \frac{11}{12} cognitively demanding tasks.
619 \frac{1}{M} Methods: Sixteen healthy young adults participated in the experiment. Pain was induced by intramuscular \frac{1}{1}
620 ^{1} hjection of hypertonic saline solution (1mL, 6%) in both vastus medialis and vastus lateralis muscles (0.9%)
621 \, ^{\circ} Tsotonic saline was used as control). The participants stood barefoot in tandem position for one minute on a
622 1 force plate. Center of pressure (CoP) was recorded before and immediately after injections, while performing
623 2 two cognitive tasks: (i) counting forwards by adding one; (ii) counting backwards by subtracting three. CoP
\frac{22}{2} gariables – total area of displacement, velocity in anterior-posterior (AP-velocity) and medial-lateral (ML-
\begin{array}{c} 24 \\ 25 \end{array} velocity) directions, and CoP sample entropy in anterior-posterior and medial-lateral directions were
\frac{26}{27} displayed as the difference between the values obtained after and before each injection and compared
627 ^2 between tasks and injections.
628 3 Results: CoP total area (-84.5 ± 145.5 vs. 28.9 ± 78.5 cm²) and ML-velocity (-1.71 ± 2.61 vs. 0.98 ± 1.93 cm/s)
629 3 Decreased after the painful injection vs. Control injection while counting forward (P < 0.05). CoP total area
630 3412.8 \pm 53.9 vs. -84.5 \pm 145.5 cm<sup>2</sup>), ML-velocity (-0.34 \pm 1.92 vs. -1.71 \pm 2.61 cm/s) and AP-velocity (1.07 \pm 1.00 cm/s)
35 36.35 vs. -0.39 ± 1.82 cm/s) increased while counting backwards vs. forwards after the painful injection (P < 100
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38.05).
\frac{39}{40} conclusion: Pain interfered with postural stability according to the type of cognitive task performed,
^{634} suggesting that pain may occupy cognitive resources, potentially resulting in poorer balance performance.
<sub>635</sub> 43
4 Keywords: postural stability, center of pressure, attention, distraction, pain
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&ist of abbreviations
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            ANOVA
                          Analysis of variance
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             au
                          Arbitrary units
             CoP
                          Center of pressure
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             SaEn
                          Sample entropy
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             SD
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             VAS
                          Visual analogue scale
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 $\frac{1}{1}$  the body and the environment and to generate adapted and accurate muscle activation for postural control 651  $\frac{13}{14}$  Morasso and Sanguineti 2002). Although the majority of postural control is regulated via automatic neural  $^{652}$   $^{15}$  processes (Bronstein and Buckwell 1997), higher cortical centers are significantly involved in processing  $^{16}$  $653~1\,\mathrm{Fensory}$  information to plan and execute the best motor strategy for postural control (Winter 1995). In daily 654 1 Ife, postural control is challenging as several tasks simultaneously compete for the cognitive resources 655 2 available (Woollacott and Shumway-Cook 2002), limited by the capacity of higher centers to process sensory  $\frac{22}{2}$  information (Kahneman 1973). Therefore, sharing attentional resources may cause impairments in the  $\frac{24}{25}$  performance of daily living activities (Brauer et al. 2004). For example, competition for cognitive resources 658 during tasks involving postural stability results in body stability being prioritized over secondary tasks (Liston 2.7 659 <sup>2</sup> et al. 2014).

Controlling of upright posture requires a significant amount of attention to gather information from

661 3 Quantify the extent to which attention is associated with postural control. Decreases in postural sway while 662 3 performing a secondary task compared with control conditions have been reported (Andersson et al. 2002; 663 3 Rellecchia 2003) whereby focusing the attention on standing as still as possible increased postural sway  $\frac{37}{38}$  compared with conditions without similar instructions (Vuillerme and Nafati 2007). Altogether, these results  $\frac{39}{40}$  suggest that postural control demands attention (Woollacott and Shumway-Cook 2002) and that 

Dual tasks paradigms, where subjects perform an additional task during standing, are employed to

Although detrimental effects of cognitive loading on postural sway during unperturbed standing are 668-45 more commonly reported for older adults and patients, studies using dual-task approaches in young subjects 669 43 how controversial results (Huxhold et al. 2006; Fraizer and Mitra 2008). Young healthy subjects have 670-4  $oldsymbol{\wp}$ robably more ability to allocate the attentional resources without sacrificing postural stability, showing that

1 2 R system without impairments prioritizes postural stability when dealing with dual-cognitive tasks (Siu and 672 1 Woollacott 2007). Subjects with pain demonstrate increased postural sway compared with controls (Hirata et al. 2011). 674  $\stackrel{1}{\underset{1}{\overset{A}{A}}}$  possible explanation for this finding is that the increased postural sway may relate to a disrupting effect of 675  $^1$  hociceptive stimuli on attention to other simultaneous non-nociceptive tasks (Eccleston et al. 1999),  $676~1\,\mathrm{Jm}$ derlining that processing of nociceptive stimuli is cognitively demanding (Veldhuijzen et al. 2006). Thus, 677 19he execution of cognitive tasks during pain might interfere with postural control. Although previous studies 678 2 have shown that patients with pain present impaired balance while performing a secondary cognitive task  $\frac{22}{2}$  (or comparison to health subjects (Van Daele et al. 2010; Larivière et al. 2013; Mazaheri et al. 2014; Sherafat  $\begin{array}{c} 24 \\ 261 \\ 261 \end{array}$  et al. 2014; Etemadi et al. 2016; Levinger et al. 2016), it is not clear yet the isolate effect of pain since reduced  $\frac{26}{27}$  muscle strength, reduced flexibility and degenerative changes at the affected segment also cause both  $^{28}$ tiffness and instability in patients suffering from chronic pain (Knoop et al. 2012). Therefore, further 683 3 Provestigation of the interaction between pain, cognition and postural stability is warranted. This investigation 684 32s of particular interest for clinical practice since there are evidences that attention can be directed away 685 3 4 from pain using some specific strategies (Van Ryckeghem et al. 2018). If selective attention could be directed 686 ു Away from the painful stimulus and modify the deleterious effect of muscle pain on postural control, these 687 3 gesults could have important implications for clinical settings. Likewise, if the execution of cognitive tasks  $\frac{39}{40}$  impairs postural control in the presence of pain, this should also be taken into account in rehabilitation 689 <sup>41</sup>context. <sub>690</sub> 43 Considering that posture can be defined as the dynamic stability of a continuous moving body 691 4 \$Harbourne and Stergiou 2003; Madeleine et al. 2011), nonlinear analysis of the dynamic structure of the 692 4 Tenter of pressure (CoP) time series would contribute to understand the physiological complexity of posture 693 4 by accessing motor patterns that would be implicit in the CoP variability. Sample entropy (SaEn) measures  $\frac{1}{5}$  variations in the system output along time. Therefore, measures of physiological complexity of the postural 52 53

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700 1 The temporal series is (Richman and Moorman 2000).

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Complexity depends on the number of structural components of the system, the existing coupling 20
2 mmong these components and how this interaction is influenced by the intrinsic dynamic properties of the 22
3 system and the motor task demands (Vaillancourt and Newell 2002). Thus, if the presence of pain and the 24
2 execution of a cognitive task are both concurring with the attentional resources used in postural control, 26 then the coupling between the components of the system responsible for balance may be affected and, 27

3 Quring standing increases the complexity of the postural sway, and this increase has been attributed to a

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708 3 2 nore automatized postural sway, when less attention is directed to the balance control (Donker et al. 2007;

35 3 4 tins et al. 2009; Kuczyński et al. 2011). On the other hand, there is some evidence that the complexity of

33 gostural control decreases with pain during sitting with increased perceived discomfort in healthy young

 $\frac{37}{38}$  ubjects (Søndergaard et al. 2010). Similar finding was reported in young subjects with transient acute

712  $\frac{39}{40}$  episode of low back pain during two continuous hours of standing, but without history of low back pain

 $^{41}$ Fewster et al. 2017), showing a relation between the occurrence of pain and the decrease in CoP complexity.  $^{42}$ 

 $^{4}$  herefore, examining the complexity of postural sway in a dual task context and the effect of experimental  $^{44}$ 

715 45pain in this condition may improve the understanding of the decrease in postural stability (Levinger et al. 46

716 4 **7**016) and complexity (Fewster et al. 2017) that may exist as a result of pain in an otherwise healthy system.

The aim of this study was to quantify how postural stability [CoP sway velocity and area of

718  $5\dot{d}$  isplacement and complexity (CoP SaEn)], is modified during experimental pain while performing a cognitive

2 gask. It was hypothesized that (i) the kind of cognitive task (more or less demanding) in a non-painful 9 720 <sub>1</sub> Condition will not interfere with CoP sway or CoP complexity, since the system would have enough cognitive  $\frac{11}{1}$  resources to overcome it; (ii) experimental pain will increase CoP sway and decrease CoP complexity, 722  $\frac{13}{12}$  regardless the type of cognitive task performed; (iii) the presence of experimental pain while performing a  $\frac{13}{14}$ 723  $^{1}$  difficult cognitive task will overload the cognitive resources and impair postural stability, increasing CoP sway 724 17 and decreasing CoP complexity. 725 1**2**. Methods 20 726 2**½**.1. Subjects 22 727 23 Sixteen young adults, all university students, (to control for the effect of education level on 728  $\frac{24}{25}$  multitasking performance (Voos et al. 2015)), participated in the experiment – 8 males (mean ± SD: age = 729  $\frac{26}{27}$  26.9 ± 2.8 years; body mass = 74.9 ± 13.8 kg; height = 1.76 ± 0.08 m) and 8 females (mean ± SD: age = 27.1 ± 730  $\frac{28}{9}$  .0 years; body mass = 68.8 ± 5.2 kg; height = 1.68 ± 0.06 m). The exclusion criteria were body mass index 731 3 bove 25 kg/m², pregnancy, drug addiction, previous neurologic, musculoskeletal or mental illness, lack of 732 3 ability to cooperate, current use of medications (e.g. analgesics, anti-inflammatory medicine), consumption 733 3 of alcohol, caffeine, nicotine or painkillers 8 hours prior to the data collection, recent history of acute pain 734 ج $oldsymbol{\mathsf{g}}$ ffecting the lower limb and/or trunk, past history of chronic pain conditions, participation in other pain  $\frac{37}{38}$  rials throughout the study period. All procedures performed in studies involving human participants were 736  $\frac{39}{40}$  n accordance with the ethical standards of the institutional and/or national research committee and with  $^{41}$ the 1964 Helsinki declaration and its later amendments or comparable ethical standards. The study was  $^{42}$ 738  $^4$   $^4$  approved by the local Ethics Committee (N-20120077). This sample size was calculated to detect a minimum 739  $4\,\Xi$  ifference of 40% in the CoP area assuming type error 1 as 5% and power of 80% between the conditions 740 4 before and after the induction of experimental pain. All participants gave signed informed consents prior to 741 4 inclusion in the study. 742 51.2. 52 Experimental protocol 53

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 $\frac{9}{744}$   $\frac{9}{1}$  Originates from proprioceptive systems (Peterka 2003), we controlled the effect of different footwear on 745  $\frac{11}{12}$  postural control by asking the subjects to stand barefoot during the experiment. The participants stood on 746  $\frac{13}{4}$  triangular force plate that measures vertical forces (Good Balance System, Metitur, Jyväsklä, Finland; 747  $^{1}\sqrt[3]{}$  dimensions: equilateral triangle – 800-mm; sampling frequency: 50-Hz as suggested by the International 749 1 Peliable system for postural sway measurements (Era et al. 2006; Ha et al. 2014) with accuracy better than 750  $\,2\,1$ -mm for the CoP position measurement (Good Balance System User Manual). The CoP position was  $\frac{22}{2}$  Falculated via the Good Balance Software (Metitur, Jyväsklä, Finland) which uses the weighted arithmetic  $\begin{array}{c} 24 \\ 25 \\ \end{array}$  mean between the vertical force measured by four sensors and their corresponding position: one in each 753  $\frac{26}{27}$  corner of the force-plate and the last one in the centroid of the force-plate (Fig. 1). The rational for using the 754  $^{28}$ andem position for the feet was based in previous studies showing that greater pain effects are presented 755 3 when posture is challenged (Hirata et al. 2013). This was important to ensure that postural stability 756 32daptations due to pain could be observed. Therefore, subjects were asked to stand in tandem position, to 757 34ncrease postural challenge during the tasks, with the right leg behind (Fig. 1), arms hanging relaxed 758 ു $_{f A}$ longside the body, and were instructed to maintain balance while looking forward. Tape markers were 759  $\frac{3}{3}$  placed on the force plate to ensure that the same foot position was maintained through all conditions. During  $\frac{39}{40}$  the assessment of postural control, subjects were instructed to look forward at a target positioned at eye- $^{41}$ evel approximately 45-cm from the subjects to minimize the influence of the target distance on postural  $^{42}$ 762  $^{4}$   $^{3}$ way (Kapoula and Lê 2006). CoP records were made under eight experimental conditions, depending on the 763  $\,4\,$  Sype of injection (control or painful), the dual-task (counting forward or counting backward as the less and 764 47 more challenging tasks, respectively), before (pre-injection) and immediately after the injection. The 765 4 gounting forward task consisted of adding one and the counting backward was performed by subtracting

 $\frac{1}{5}$  three, beginning from a random number. The total number of answers and the number of correct answers

Since in healthy individuals approximately 70% of the information used for controlling posture

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during each trial were recorded. The order of the injections and the order of the tasks were randomized, with the same number of subjects receiving the hypertonic or isotonic injections first.

The experiment always followed the same order for all participants: (i) CoP measurement while

The experiment always followed the same order for all participants: (i) CoP measurement while

performing the first randomly assigned task (cognitive task 1 or 2) over 60-s (pre-injection 1); (ii) 1-min rest;

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The experiment always followed task (cognitive task 1 or 2) over 60-s (pre-injection 2); (iv) injections of the first saline solution (painful or control) into vastus mediants.

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773 1 **9**VM) and vastus lateralis (VL) muscles; (v) assessment of pain intensity by visual analogue scale (VAS); (vi)

774 2 CoP measurement over 60-s while performing task A; (vii) collecting VAS scores of the pain intensity and 1-

22 775 2 min rest; (viii) CoP measurement over 60-s while performing task B; (ix) collecting VAS scores of the pain

776  $2\frac{4}{2}$  intensity. After the final step, the pain VAS scores were taken each minute until the pain had subsided which

777  $\frac{26}{27}$  was followed by a 5-min break. Following the break, all steps of the experiment were performed again with

778 <sup>2</sup> the injection of the other saline solution, including new pre-injection CoP recordings. Before each CoP 29
 779 <sup>3</sup> neasurement, all subjects confirmed that no tiredness or other problems were presented. The duration of

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The Cop measurements were performed asserting to guidelines proposed by the International Society for

780 3 2he CoP measurements were performed according to guidelines proposed by the International Society for

781 3 Posture and Gait Research (Scoppa et al. 2013). Fig. 2 summarizes the study procedures along time.

# 782 <sub>3</sub> **2**.3. Experimental muscle pain

Before the experiment all subjects were instructed about the nature and effects of the injections,

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leading the subject of the injection would be a non-painful stimulus, although

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4 Intramuscular injection of 1-mL of 6% sterile hypertonic saline solution or as a control condition 1-mL of 44
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788 4 The injections were performed with a 2-mL syringe with a disposable needle (27G, 40-mm) into right VM 48

4 gnuscle and right VL muscle. Both injections locations were marked to ensure that they were applied
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 5 approximately in the same location. The VM muscle injection was performed 5-cm proximal and 5-cm medial

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791 \$\footnote{\text{two the medial corner of the patella (Shiozawa et al. 2013), and in the VL muscle, injections were performed

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792 \$\frac{9}{1}\$ at two thirds of the distance from the anterior spina iliaca to the lateral side of the patella (Fig. 3). The depth

793  $\frac{11}{12}$  of the injection was determined by an ultrasound scanner (LOGIQ<sup>™</sup> S7, General Electric, USA). This pain

794  $\frac{13}{14}$  model has been successfully used previously to mimic knee-related pain during quiet standing tasks

 $^{1}5$  providing moderate pain intensities for approximately five minutes (Hirata et al. 2011). Hypertonic saline  $^{1}6$ 

 $^{17}$ njections have been shown to activate nociceptors around the injected site (Mense 1993) whereas the 0.9%

797 1 % sotonic saline injections have induced little or no pain during postural control tasks similar to the one used

798 2 in the present study (Hirata et al. 2010, 2011, 2013).

## 22 799 23.4. Assessment of pain intensity

The subjects were asked to rate the pain intensity using a 10-cm VAS from 0-cm to 10-cm (0-cm 26 means "no pain" and 10-cm means "maximum pain") immediately after the injections and after each balance 28 measurement. Therefore, three VAS scores were obtained for each set of experiments (balance 29 a measurements after isotonic injection and balance measurements after hypertonic injection, respectively; 31 a paradigm. Additionally, following each set of experiments subjects were asked to indicate the overall pain 3 paradigm. Additionally, following each set of experiments subjects were asked to indicate the overall pain

35  $_{3}$  ereas during the trials on a body chart and to respond the McGill Pain Questionnaire (Melzack 1975). The  $_{3}$  erea of pain was extracted from the body charts with VistaMetrix 1.38 software. The pain rating index based

 $\frac{39}{40}$  rea of pain was extracted from the body charts with vistametrix 1.38 software. The pain rating index based the values of the words chosen within each category (sensory, affective, evaluative and

 $\frac{41}{10}$  miscellaneous) from McGill Pain Questionnaire were obtained and the score for each category, as well as  $\frac{41}{10}$ 

810 4 he total pain rating index were determined as the sum of the ranked values of the words (Melzack 1975).

# 811 45.5. Data analysis

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All variables for postural sway were calculated based on 50-s of the standing tasks, with the first and

813 4 gast 5-s from the original 60-s time series being excluded. The analyses were performed with Matlab R2016a

814 5 oftware (Mathworks, Massachusetts, USA). The area fitted to 95% confidence interval of the CoP

1 2 displacement was calculated as representative of the total CoP area displacement (95% confidence interval  $\frac{9}{1}$  (Blipse), along with the CoP velocity in both directions (anterior-posterior and medial-lateral). The structural 817  $\frac{1}{1}$  xariability of the CoP was calculated by means of SaEn with the embedding dimension (m) and the tolerance 818  $\frac{1}{4}$  distance (r) set to m=2 and r=0.2xSD (Vaillancourt and Newell 2000). All CoP variables are displayed as the  $\frac{1}{4}$  $^{1}$  difference between the values obtained immediately after the injection and the correspondent pre-injection 820  $\,^17$  ondition. Negative values show that the CoP variable decreased after the injection of the saline solution 821 1@ompared to its respective pre-injection condition. Likewise, positive values show that the CoP variable 822 2 increased after the injection compared to its respective pre-injection condition. Statistical analysis Pain outcomes were compared between injection types (isotonic or hypertonic injections) with 825  $\frac{26}{27}$  paired T-tests when normal distribution was present (VAS scores and pain area data) and with the Wilcoxon 826  $\frac{28}{3}$  igned Rank Test when the data distribution was non-normal (McGill scores). The task measures (number of 827 3 nswers, number of correct answers) were evaluated with a 3-way RM-ANOVA with injection (isotonic vs 828 3 Dypertonic), time (pre-injection vs after injection) and task (counting forward vs backwards) as main factors. 829 3 The CoP parameters were compared with a 2-Way RM-ANOVA with task and injection as main factors, and 830 3 the p-values are shown in the table 3. Bonferroni post-hoc correction for multiple comparisons was applied  $\frac{37}{38}$  and p-values are shown in the results texts. The alfa-value ( $\alpha$ ) for statistical significance was set to 0.05. Area and amplitude of perceived pain' <sub>834</sub> 43 Fig. 4 shows the reported pain areas following both isotonic and hypertonic injections. Pain was 835  $^4$  present in the anterior and lateral portions of the thigh after both isotonic and hypertonic injections, being

836 4 Thore concentrated in the lower half of the thigh after the isotonic injections. The hypertonic saline injections

837 4  $\frac{1}{2}$   $\frac{1}{2$ 

838  $\frac{1}{5}$  f = 0.003) and higher VAS scores (mean score ± SD: isotonic = 0.9 ± 1.1 cm; hypertonic = 4.7 ± 1.7 cm; P<0.001)

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1 2 than isotonic saline injections. Table 1 shows the scores for each class of words from McGill Pain  $\frac{9}{10}$  Ruestionnaire and the pain rating index. Subjects presented a higher total pain rating index and scored 841  $\frac{1}{1}$  higher in all the categories, with the exception of the affective class, after the hypertonic injections (P<0.05). 842 13.*2* Cognitive task performance <sub>843</sub> 15 Only for the analysis of the cognitive task performance, one subject was not included due to problems  $844~1\,\mathrm{fh}$  the answers recording. The total number of answers and the number of correct answers decreased during 845 1 hackwards counting conditions compared with forwards counting despite the injection effect (significant 20 846 2 main effect for task factor; Table 2). 22 847 2**3**.3 Center of pressure 848 25 Effect of experimental pain in CoP variables <sub>849</sub> 26 There were no statistical differences between the different conditions for the factor *injection* on any 850  $^{2}$ 8f the CoP variables (Table 3). 851 30 Effect of cognitive task in CoP variables 31 852 32 A main effect of task was found for the CoP AP-velocity (F=5.82; P=0.028), showing that there was an 853 3 Amcreased AP-velocity during the counting backwards task compared to the counting forwards task, 3% egardless the type of injection (Table 3). 855 38 Effect of the interaction between experimental pain and cognitive task in CoP variables 856 39 An interaction effect was found between injection and task factors for CoP total area and CoP ML-857 4 velocity (CoP total F=7.78, P=0.049; CoP ML F=4.69, P=0.021) (Table 3). Post-hoc comparisons showed that 858  $^4$  both variables decreased after the hypertonic injection in comparison to the condition with isotonic injection 859 45when subjects where counting forward (Bonferroni: P = 0.010 for total area; P = 0.015 for ML-velocity). After 860 4 The hypertonic injection, CoP total area increased when subjects were counting backwards in comparison to 861 4 gwhen they were counting forwards (Bonferroni: P = 0.019). ML-velocity showed differences between the 50 51 52 53 57 58 59 60 61

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#### Discussion

The present study aimed at quantifying how postural stability, represented by CoP sway (velocity and

866 1 area of displacement) and CoP complexity (CoP SaEn), is modified during experimental pain while performing

different cognitive tasks also after the injection of hypertonic solution, with a smaller decrease of ML-velocity

867 17 cognitive task. The main results showed that the kind of cognitive task did not interfere with postural

868 1 9 tability in the absence of pain. Experimental pain around the knee joint reduced CoP sway but did not affect

869 2 CoP complexity during the performance of an easier cognitive task. During experimentally induced pain, the

 $\begin{array}{c} 22\\ 29 \end{array}$  performance of a difficult cognitive task increased CoP sway but did not change CoP complexity.

# 871 $\frac{24}{2}$ Pain intensity and counting performance

The subjects showed higher pain intensity for the hypertonic saline injection and a larger pain area 873  $^{2}$ 8 compared with the isotonic saline injection, as expected, indicating that experimental pain occurred (Hirata 874 3 et al. 2011). The McGill pain questionnaire indicated that hypertonic saline was perceived more impairing

875 3 2 than the isotonic injection in all subscales except for the affective one. It is important to note that during

876 3 4 sotonic injections subjects rated pain around 1/10, which cannot be classified as a totally pain free condition.

Counting performance requires the use of cognitive process which relies on the working memory of  $\frac{37}{3}$  the subject (Lemaire 1996), impairing motor output performance when executed simultaneously with a 879  $\frac{39}{40}$  motor task (Vuillerme and Nafati 2007). Seminowicz and Davis (2007) showed that subjects are able to  $^{41}_{
m maintain}$  performance of difficult cognitive task while experiencing different levels of pain. In this study, the  $^{42}$ 881  $^4$  painful condition did not affect the counting performance while performing a motor task (standing still)

882 4 Endicating that healthy subjects are able to engage multiple tasks (motor and cognitive) during pain without

883 4 Tompromising performance. This suggests that sufficient cognitive resources were available to manage the

884 4 gognitive process of counting forwards or backwards despite the interpretation of painful stimuli and the

885  $_5$  postural control task (Eccleston et al. 1999). Finally, education level is associate with both motor and

gerceptual performance, where higher education level is associated with better performance (Voos et al. 887  $\frac{9}{10015}$ . Since our subjects were all university students, we believe that bias due to education level did not 888  $1\frac{1}{12}$  affect the present results.

# 889 $\frac{1}{1}$ Effect of cognitive tasks on postural stability

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Our first initial hypothesis, that (i) the kind of cognitive task (more or less demanding) in a non-painful 891  $\,^17$  condition would not interfere with CoP sway or CoP complexity, was confirmed. The factor task affected the 892 1 © oP anterior-posterior velocity, indicating an increased velocity during the execution of the more difficult 893 2 task (counting backwards) in comparison to the easier task (counting forward). Nevertheless, the CoP SaEn

 $\frac{22}{2}$  was not affected by the kind of the performed cognitive task. These results indicate that enough cognitive  $\begin{array}{c} 24 \\ 25 \end{array}$  esources were available to overcome the demands of both cognitive and postural tasks, which was expected 896  $\frac{26}{27}$  ince they were young individuals without any sensory-motor alterations.

### 897 <sup>28</sup> Effect of experimental knee-related pain on postural stability

898 30 Our second initial hypothesis, that (ii) experimental pain would increase CoP sway and decrease CoP 899 32omplexity was not confirmed since the type of saline solution injected did not affect the CoP variables. 900 3 However, even though the factor injection did not show statistical differences between the different 901 3 gonditions for any of the studied CoP variables, there was a difference between total area and ML-velocity  $\frac{37}{38}$  between the control and the painful condition when the subjects were counting forwards, i.e., in conditions  $\frac{39}{40}$  where the kind of cognitive task performed was the same. Interestingly, during the counting forward, the  $\frac{41}{42}$  ype of injection resulted significant changes in postural sway (total area and ML-velocity) in opposite  $^{905}$   $^4$  directions: positive values of the difference between pre-injection and after injection of the isotonic solution, 906 45whereas after the injection of the hypertonic solution both variables showed negative values. Additionally,

907 4 no significant changes were observed in the structural variability of the CoP signal. This is contrary to the 908 4 mitial hypothesis, where an increase in postural sway and a decrease in structural variability during painful  $\frac{1}{5}$  conditions were expected. It is also in contrast with previous findings (Mazaheri et al. 2013) but may relate <sub>929</sub> 43 932 4 gesults showed an interaction between the task and injection factors for total area and ML-velocity. After 933 5 the hypertonic injection CoP total area increased and CoP ML-velocity decreased less while counting 52 53

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38 go the different position of the feet used in this study, which affects the postural sway (Day et al. 1993). The  $^9$  911  $_1$  tandem feet position adopted allows less displacement of the CoP due to the limited base of support  $\frac{11}{1}$  compared to side-by-side feet position, since if the subjects increase the CoP amplitude they may fall (Day  $^{913}$   $^{1}$   $^{3}$  et al. 1993). This also may reflect a voluntary strategy, requiring a greater amount of cognitive resources and  $^{1}$   $^{4}$ 914  $^{\,1}ar{s}$ ttention (Morasso and Sanguineti 2002), attempting to avoid large excursions of the body and consequent 916 1 he other tasks, also known as posture first strategy (Vuillerme and Nafati 2007). The subjects were able to 21educe the postural sway without compromising the counting performance during the easy cognitive task,  $\frac{22}{2}$  guggesting that the available cognitive resource was sufficient to perform the less challenging cognitive task  $^{24}_{^{2}}$  without compromising postural stability. Therefore, these results indicate that healthy subjects have the  $\frac{26}{27}$  capacity to perform easy cognitive tasks while ensuring postural stability (Siu and Woollacott 2007). 921  $^{2}$ Reducing postural sway might reflect a motor strategy available for healthy subjects to avoid excessive 922 <sup>3</sup> Pranslation of the body, which could lead to balance loss (Winter 1995). This strategy was also observed 923 3 Auring the control injection while counting backwards, probably indicating that a high cognitive load seems 924 340 be interpreted as a treat to postural stability. An alternative explanation for the contrast between the 925 3 present study and the previous studies with pain patients showing larger postural sway (Schulte et al. 2004;  $\frac{37}{38}$  evinger et al. 2016) might be the pain model used that is not a complete proxy to the impaired pain patients' 927  $\frac{39}{40}$  sensory-motor system. 928  $\frac{41}{m}$  mteractions between pain and cognitive load on postural stability 42Our initial third hypothesis, that (iii) the presence of experimental pain would increase CoP sway and 930 4 Diecrease CoP complexity only when performing a difficult cognitive task was partially confirmed since CoP 931 4 way increased during pain under a difficult cognitive task, but the CoP complexity did not change. ANOVA

39 packwards in comparison to counting forwards condition, corroborating our hypothesis. ANOVA results also 9  $_1$  showed an effect of the task factor on AP-velocity with post-hoc comparisons showing a difference only  $\frac{11}{12}$  during the hypertonic injection condition: while counting backwards AP-velocity also increased. Altogether  $^{13}$  these results show that CoP sway increases when performing a more demanding cognitive task in the  $^{14}$ 938  $\frac{1}{9}$  resence of experimental pain. This might reflect an interference with the information-processing capacity  $\frac{1}{6}$ 940 1 9tudies suggest that disruptions of sensory information lead to worsening of proprioception in the affected  $^{941}$   $_{2}$  area (Matre et al. 2002), further impairing postural sway (Hirata et al. 2010, 2011). The results indicate that  $\frac{22}{2}$  the posture first strategy (Vuillerme and Nafati 2007) found during the easy cognitive task during pain is no  $\begin{array}{c} 24\\ 25\\ \end{array}$  Longer feasible when a difficult cognitive task is performed during painful conditions. The increased cognitive  $\frac{26}{27}$  load in painful conditions seems to impair the motor performance maybe due to insufficient cognitive 945  $^{28}$ esource to simultaneously maintain postural stability (which requires significant amount of attention 946 3 (Morasso and Sanguineti 2002)) and execute a difficult cognitive task. These results might have important 947 3 Dew implications in understanding the mechanisms related to fall accidents. Postural stability in daily life 948 3 4 ctivities is usually performed in combination with additional tasks, for example, walking in a busy slippery 949 3 gidewalk. These daily life activities involves simultaneously competition for the cognitive resources available  $\frac{37}{3}$  (Woollacott and Shumway-Cook 2002) to evaluate the environment constrains in order to promote the best 951  $\frac{39}{40}$  motor strategy (Winter 1995). Our present results indicate that, if the subject performs a challenging 952  $\frac{4}{3}$  bostural task in pain, his/her capacity for maintain balance while exposed to a difficult cognitive task is 953 <sup>4</sup> Suboptimal, which could increase the likelihood of losing balance.

The complexity of postural sway did not show any differences between the experimental conditions. 955 4 This result is contrary to the literature finding that young healthy subjects present a more regular and less 956 4 gutomatic postural sway (decreased CoP SaEn) when the motor task is more difficult (e.g. standing with eyes 957 5 closed) and more irregular postural sway and more automatic postural sway (increased CoP SaEn) when a

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959 1 with CoP complexity may be due to the nature of both motor (standing in tandem position) and cognitive
960 11 subtraction calculus) tasks used in the experimental setup that did not interfere with the automaticity of
961 12 postural control. Besides that, pain also did not affect CoP complexity, showing that experimental knee962 13 pelated pain did not compromise the coupling between the components of the system responsible for
963 13 balance in the current experimental setup. Future studies should investigate the interaction between pain,
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964 13 pognition and on CoP complexity with different motor and cognitive demands, in addition to different

gognitive task is added (Donker et al. 2007; Stins et al. 2009). The fact that the cognitive task did not interfere

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965 2 populations.

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966 23 Despite interesting results regarding the effects of cognitive tasks in postural control during pain, the

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25 elevance of the findings for clinical populations should be interpreted with care. The experimental pain

968 2 model used here is convenient to assess the effect of pain without the interference of potential structural

28 or pathologies. However, extrapolating the current findings to an older population can only be done to some 29
 3 degree. Additionally, chronic pain patients may also suffer from depressive symptoms (Bair et al. 2003) or

971 3 2 2nxiety (McWilliams et al. 2003), which might increase cognitive load (Nebes et al. 2001). Furthermore,

334 ognitive impairments are often found in chronic pain patients, decreasing the possibility to maintain

973 3 performance of two or more concurrent tasks (Brauer et al. 2004), as opposed to what was observed in this

 $\frac{37}{38}$ tudy where young healthy subjects were recruited. Also, there was no recording of postural sway without

 $\frac{39}{40}$  any cognitive task. This would have allowed comparisons with a condition where neither pain nor cognitive

 $^{41}$ asks were influencing postural sway, and could have reduced type 2 errors given that multiple CoP variables  $^{42}$ 

977  $\frac{43}{4}$  were analyzed in the study. Thus, it can be considered a limitation to our interpretations.

#### Conclusions

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63 64 Pain and cognitive task interfered on postural stability, changing its patterns. During the performance

980 4 of a simple cognitive task, pain reduced postural sway, while during the performance of a more demanding

981 5 cognitive task, postural sway was increased in young healthy subjects. Since our subjects were young healthy

3 41 gubjects, the direct translation of the present results to patients suffering from pain should be done with  $^9$  983  $_1$  Gaution. However, these results may suggest that rehabilitation approaches should take into account that 984  $\frac{11}{12}$  pain not only affects directly the motor system, but may occupy cognitive resources, potentially resulting in 985  $\frac{1}{1}$  poorer performance when performing rehabilitation exercises. Additionally, rehabilitation strategies using 986  $^{1}$ 80th motor and cognitive resources need further investigation to outline the effect of interaction between 987 1 pain and cognition on the performance during activities of daily life in patients. 988 19 989 2 Compliance with ethical standards  $\frac{22}{2}$  **Funding:** Center for Neuroplasticity and Pain (CNAP) is supported by the Danish National Research 991  $\frac{24}{\text{Foundation}}$  (DNRF121). The authors thank the State of São Paulo Research Foundation (FAPESP) for the Suda 992 <sup>2</sup> §cholarship (FAPESP 2013/06123-7, 2015/00214-6). 993 **2 Conflict of Interest:** The authors declare that they have no conflict of interest. 29 994 3 **REFERENCES** 31 Andersson G, Hagman J, Talianzadeh R, et al (2002) Effect of cognitive load on postural control. Brain Res 995 996 Bull 58:135-9 997 3 & air MJ, Robinson RL, Katon W, Kroenke K (2003) Depression and Pain Comorbidity. Arch Intern Med 163:2433. doi: 10.1001/archinte.163.20.2433 998 35  $^{3}$   $^{6}$  rauer SG, Broome A, Stone C, et al (2004) Simplest tasks have greatest dual task interference with balance .000 37 in brain injured adults. Hum Mov Sci 23:489-502. doi: 10.1016/j.humov.2004.08.020 3 Bronstein AM, Buckwell D (1997) Automatic control of postural sway by visual motion parallax. Exp Brain .002 40 Res 113:243-248. doi: 10.1007/BF02450322 .003  $4\,\mathrm{Day}$  BYBL, Steiger MJ, Thompson PD, Marsden CD (1993) Human Body Motion When Standing : .004 42 Implications for. J Physiol 469:479-499 .005 <sup>43</sup> Donker SF, Roerdink M, Greven AJ, Beek PJ (2007) Regularity of center-of-pressure trajectories depends on .006 44 the amount of attention invested in postural control. Exp Brain Res 181:1-11. doi: 10.1007/s00221-.007 45 46  $\frac{1}{4}$  puarte M, Sternad D (2008) Complexity of human postural control in young and older adults during .009 48 prolonged standing. Exp Brain Res 191:265–276. doi: 10.1007/s00221-008-1521-7 .010  $\,^4\Psi$ ccleston C, Baeyens F, Helen P, et al (1999) Pain Demands Attention : A Cognitive-Affective Model of the Interruptive Function of Pain. 125:356–366 51 Fra P, Sainio P, Koskinen S, et al (2006) Postural balance in a random sample of 7,979 subjects aged 30 years and over. Gerontology 52:204–213 doi: 10.1159/20202052 .011 50 53 54 55 56 57 58

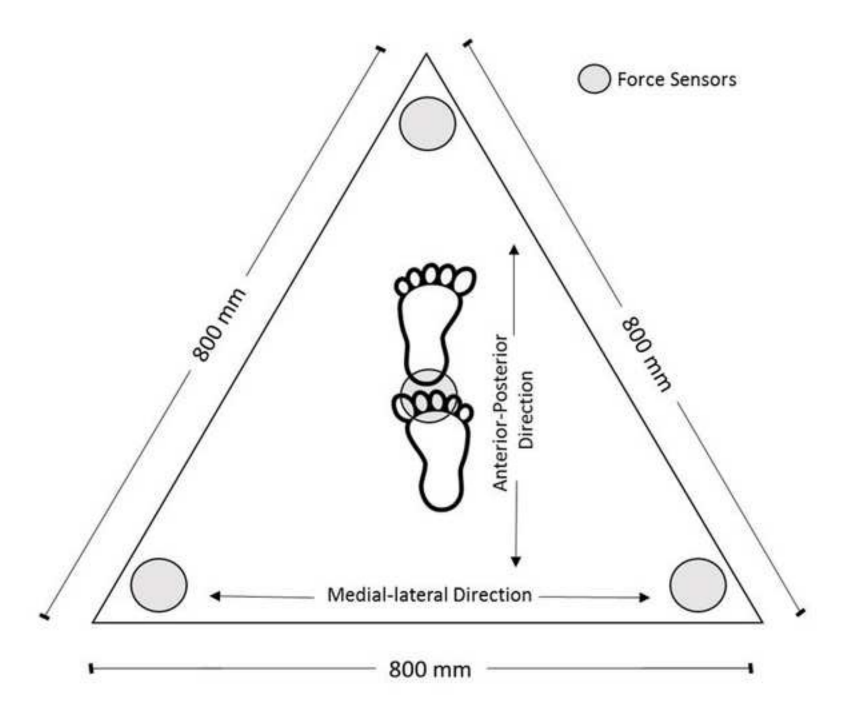
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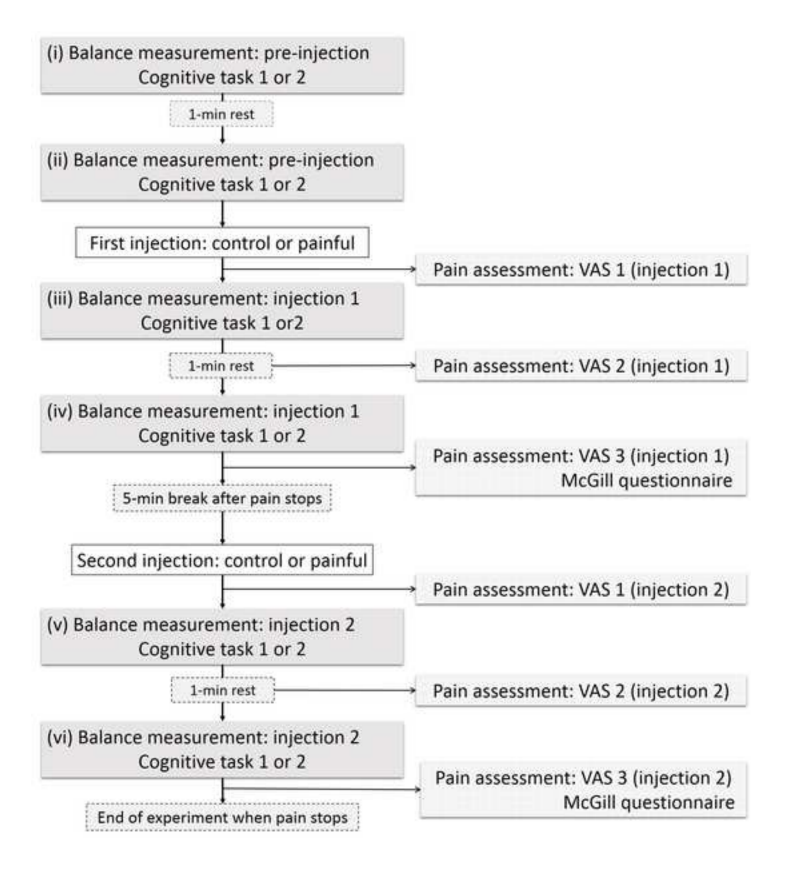
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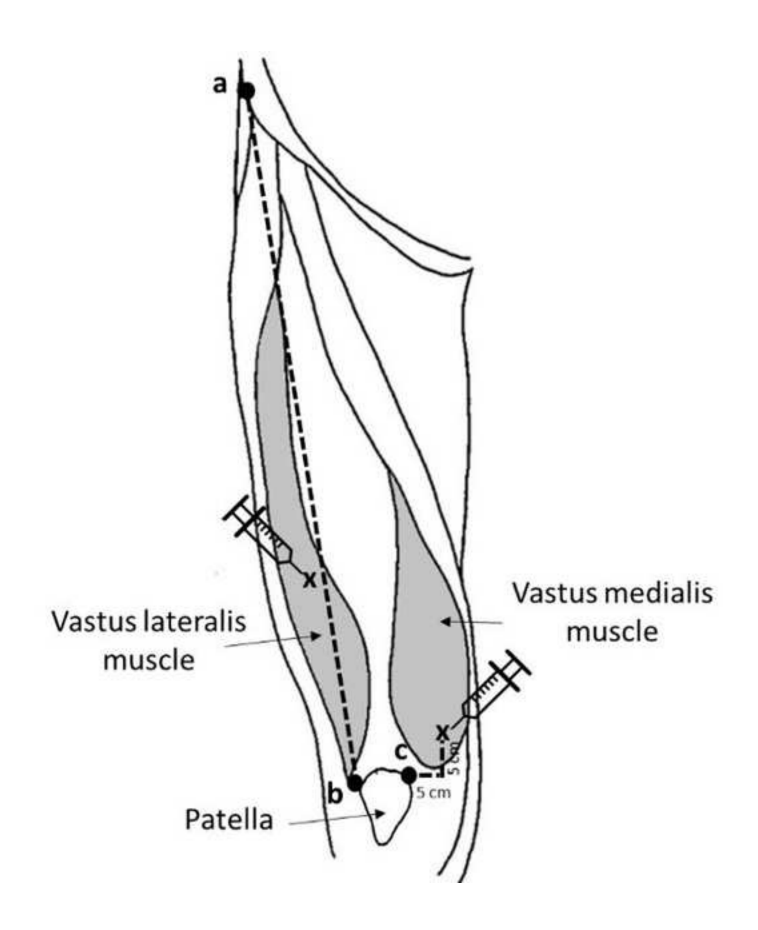
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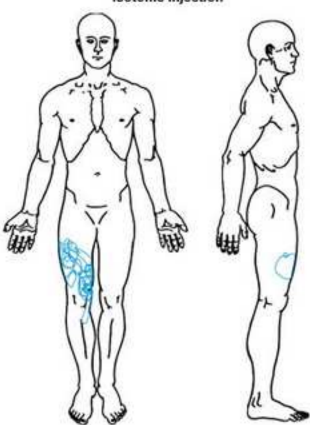
.141 8 **Figure captions** .142 10 .143  $\begin{array}{c} 1 \\ 1 \\ 2 \end{array}$  ig 1 Schematic drawing representing the force platform size, sensor locations, and the tandem position of .144  $\frac{13}{14}$  the subjects during the experiment  $\frac{14}{14}$ 145 15 .146  $\,^1$  Fig  $^2$  Study design overview: pain assessments were performed immediately after each injection and each .147 1  $\mathfrak B$ alance measurement; the order of the saline injections was randomized in a balanced way 148 21  $\frac{22}{25}$  ig 3 Injections sites for vastus lateralis muscle, performed at two thirds of the distance from the anterior  $\begin{array}{c} 24 \\ 25 \end{array}$  pina iliaca (a) to the lateral side of the patella (b); and for the vastus medialis muscle, performed 5 cm .151  $\frac{26}{27}$  proximal and 5 cm medial to the medial corner of the patella (c), .<sub>152</sub> 28 .153 3 **Fig 4** Representation of the experimental pain distribution reported areas after isotonic (top, blue in the .154 32nline version) and hypertonic (bottom, red in the online version saline injections (A); the individual .155 3 **4** istributions are superimposed in the anatomical drawings .156 36 







# Isotonic injection



## Hypertonic injection

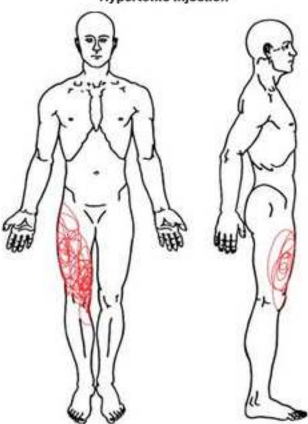


Table 1 – McGill Pain Questionnaire scores (median [Range]) for each category and total pain rating index for the pain experienced after isotonic and hypertonic injections.

McGill scores	Inject	<i>P</i> -value		
Micdili Scores	Isotonic	Hypertonic	r-value	
Sensory	1 [0-18]	8.5 [2-23]*	0.023	
Affective	0 [0-7]	0 [0-4]	0.174	
Evaluative	0 [0-1]	1.5 [0-4]*	0.001	
Miscellaneous	0 [0-7]	2.5 [0-10]*	0.004	
Total pain rating index	2.5 [0-33]	16 [5-30]*	0.001	

<sup>\*</sup>Statistically significant (*P*<0.05) higher then isotonic condition (Wilcoxon Signed Rank Test with Bonferroni correction).

Table 2 – Mean (±SD) of the cognitive tasks performances before and during both

injections type (hypertonic and isotonic) and three-way repeated measures ANOVA

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results (F; P).

Task performance		Cognitive task		ANOVA (F; P value)				
	Condition	Counting forward	Counting backward	Time	Injection	Task	Time x Injection x Task	
	Before control injection	63.3±7.5	31.3±13.5					
Total	After control injection	63.5±8.1	30.4±15.0	0.05.0.033	0.22.0.644	4 68.0; <0.001*	0.28; 0.608	
answers	Before painful injection	63.3±10.4	32.1±12.7	0.05; 0.833	0.22; 0.644			
	After painful injection	63.3±9.1	32.3±12.7					
	Before control injection	63.3±7.5	30.9±13.9					
answers B	After control injection	63.5±8.1	29.8±8.1	0.05; 0.819	0.06; 0.815	64.8; <0.001*	0.39; 0.540	
	Before painful injection	63.3±10.4	30.9±14.2					
	After painful injection	63.3±9.0	31.3±13.5					

<sup>\*</sup> Statistically significant (P<0.05).

Table 3 – Mean ( $\pm$ SD) of center of pressure (CoP) variables represented as the difference between the measures after and before each injection (isotonic injection considered as control, hypertonic injection considered as painful) and two-way repeated measures ANOVA results (F; P).

CoP Variable	Control injection		Painful injection		ANOVA (F; P value)		
	Counting forward	Counting backward	Counting forward	Counting backward	Injection	Task	Injection x task
Total area (cm²)	28.9±78.5ª	-25.1±138.7	- 84.5±145.5 <sup>a,</sup> b	12.8±53.9b	1.84; 0.196	0.75; 0.400	7.78; 0.049*
AP Velocity (cm/s)	-0.36±2.24	-0.07±1.66	-0.39±1.82	1.07±2.35	0.61; 0.446	5.92; 0.028*	1.168; 0.614
ML Velocity (cm/s)	0.98±1.93 <sup>c,</sup> d -0.73±2.23 <sup>d</sup> e		-1.71±2.61 <sup>c,</sup> e	-0.34±1.92e	3.90; 0.067	6.68; 0.697	4.69; 0.021*
AP SaEn (a. u.)	0.007±0.067	- 0.003±0.089	0.041±0.081	0.001±0.048	0.73; 0.406	1.51; 0.238	1.01; 0.331
ML SaEn (a. u.)	- 0.019±0.050	- 0.003±0.038	- 0.004±0.045	- 0.104±0.052	0.12; 0.116	0.12; 0.738	0.10; 0.755

<sup>\*</sup> Statistically significant (P<0.05). <sup>a, b, c, d, e</sup> Statistically significant difference between conditions detected in post-hoc tests (P<0.05).

Author Contribution Statement

#### **Author Contribution Statement**

RPH, TP, NV and TGN conceived and designed research. EYS and TP conducted experiments. EYS and RPH analyzed data. EYS, RPH, ICNS, TP, NV and TGN wrote the manuscript. All authors read and approved the manuscript.