

1	Spinal kinematics of adolescent male rowers with back pain in comparison to matched
2	controls during ergometer rowing
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10	
11	Funding: Not applicable
12	
13	Conflict of Interest Disclosure: The authors certify that they have no affiliations with or
14	financial involvement in any organization or entity with a direct financial interest in the
15	subject matter or materials discussed in the article.
16	
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29 30	Running Title: Spinal kinematics of rowers with back pain
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32 ABSTRACT

There is a high prevalence of low back pain (LBP) in adolescent male rowers. In this study, 33 34 regional lumbar spinal kinematics and self-reported LBP intensity were compared between 35 10 adolescent rowers with moderate levels of LBP relating to rowing with 10 reporting no history of LBP during a 15-minute ergometer trial using an electromagnetic tracking system. 36 Adolescent male rowers with LBP reported increasing pain intensity during ergometer 37 38 rowing. No significant differences were detected in mean upper or lower lumbar angles 39 between rowers with and without LBP. However, compared to rowers without pain, rowers 40 with pain had: 1) relatively less excursion of the upper lumbar spine into extension over the drive phase, 2) relatively less excursion of the lower lumbar spine into extension over time, 41 3) greater variability in upper and lower lumbar angles over the 15-minute ergometer trial, 4) 42 43 positioned their upper lumbar spine closer to end range flexion for a greater proportion of the 44 drive phase, and 5) showed increased time in sustained flexion loading in the upper lumbar 45 spine. Differences in regional lumbar kinematics exist between adolescent male rowers with 46 and without LBP, which may have injury implication and intervention strategies.

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48 Keywords: athletes, spinal pain, sports, biomechanics

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- 50
- 51 Word Count: 4330
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INTRODUCTION

The World Rowing Federation has identified that Low Back Pain (LBP) is a common condition experienced by rowers of all ages.¹ Amateur adolescent rowers aged between 14 to 16 years have been shown to have a high lifetime prevalence of LBP, with reported rates of 94% in males and 65% in female rowers.²

Mechanical loading factors such as long on-water rowing time in training sessions, 58 59 repetitive lifting of the rowing shell, and ergometer rowing have been associated with LBP in rowers.³⁻⁵ More specifically, there is a growing body of evidence suggesting that specific 60 61 patterns of spinal kinematics during ergometer rowing may be particularly provocative of LBP in rowers.^{3,6,7} In support of this, studies have identified that some rowers present with 62 large magnitude of lumbar spine flexion during ergometer rowing reflecting a potential 63 mechanism for LBP.^{3,6,7} This relationship has not yet been specifically investigated in an 64 65 amateur adolescent population. Understanding LBP mechanisms adolescent sporting populations such as rowing is important, as this is the age where most rowers take up the 66 67 sport and they appear to be particularly susceptible to LBP. Further, LBP in adolescence is a known predictor of LBP in adulthood.⁸ 68

69 It has been suggested that the repetitive nature of lumbar flexion during rowing may increase lumbar excursion during rowing,^{7,9-11} and that this has been linked to back pain.^{12,13} 70 Further, end-range flexion may also be associated with back pain, ¹⁴⁻¹⁶ as it has been 71 72 proposed that position of the lumbar spine relative to the end of range, where passive 73 structures of the spine are close to being maximally loaded or stretched, may increase the risk of tissue strain and pain. ^{17,18} Previous research has identified end-range spinal flexion in 74 sitting to be related to LBP in both sporting ^{15,19} and non-sporting populations supporting a 75 pain / postural relationship.^{16,20} 76

77 Several studies have reported spinal kinematics during rowing using healthy pain free populations and speculated a link with spinal movement and LBP.^{7,9,10} These reports have 78 79 shown that rowers frequently posture their spine at the end-range of spinal flexion with the 80 magnitude of lumbar flexion increasing over time of the rowing task, which may increase the potential for back pain.^{7,9-11} However, these investigations did not consider two separate 81 82 lumbar regions (upper and lower), which is now recognized as a more appropriate method of 83 quantifying lumbar regional kinematics, as individuals are shown to control their upper and lower lumbar spine differently during functional tasks ^{14,20,21}. At present, there is a paucity of 84 85 literature that has examined regional spinal movement during rowing and to our knowledge 86 no studies have investigated rowers with LBP. This is despite a demonstrated relationship between LBP and differences in regional lumbar kinematics in non-rowing populations.^{15,19} 87

Therefore, the aims of this study were to; investigate whether there is an increase in LBP intensity in rowers with LBP, and to investigate differences in lumbar kinematics between rowers with and without LBP, during a 15-minute rowing ergometer trial. Specifically, we hypothesized that

92 1. Pain intensity levels for rowers with LBP would increase over the course of a 1593 minute rowing ergometer trial.

Rowers with LBP would posture their upper and lower lumbar spine in a greater
degree of flexion than rowers without LBP during the drive phase of ergometer
rowing. Further, the LBP group would demonstrate greater increases in flexion over
the 15 minutes period compared to the non-LBP group.

98 3. Rowers with LBP would spend a greater proportion of the drive phase of the rowing
99 stroke with their upper and lower lumbar spine near end range flexion than rowers
100 without LBP. Further, this difference would become greater over 15 minutes of
101 rowing.

METHODS

103 Twenty adolescent male rowers, aged between 14 to 19 years, with (n=10) and without 104 (n=10) LBP participated in this study (Table 1). A power calculation prior to participant 105 recruitment suggested that 10 participants in each group would provide 80% power to detect 106 a group difference of 10 degrees (assuming a standard deviation of 10 in both groups, repeated measures for 3 phases over 1,7 and 15th minute, and a within-subject correlation of 107 0.6). Participants were included if they performed rowing training for a school-rowing club or 108 109 a community rowing club at least three times per week as well as competing in rowing 110 regattas. Participants were defined as having LBP if their self-reported LBP was located between the levels of the 1st and 5th lumbar vertebrae (i.e. $L_1 - L_5$) and if this pain was 111 112 provoked by rowing with an intensity greater than 3cm (out of 10cm) as indicated by a visual 113 analogue scale (VAS) within 30 minutes of rowing training. The characteristics of the 114 participants including; age, height, mass, body mass index (BMI), self reported level of pain 115 during participant recruitment (VAS) and their self reported disability score was collected from the Roland Morris Disability Questionnaire²² and Patient Specific Functional Scale²³ are 116 117 presented in Table 1. Participants in the no pain group had no history of LBP. Rowers were 118 excluded from this study if there was a presence of specific causes of LBP such as 119 inflammatory diseases, radicular pain or neurological signs to the lower limbs, or they had 120 reported any lower limb musculoskeletal injury in the six weeks preceding data collection. 121 Further participants were excluded if they received any rowing specific postural training 122 during previous rehabilitation of their LBP, as this may influence their spinal kinematics 123 during rowing, which this study was investigating. Permission to conduct the study was 124 granted by the Institutional Human Research Ethics Committee and all subjects and their 125 parents/guardians (where necessary) provided written informed consent/assent.

Three dimensional regional lumbar angles were collected using the 3-Space FastrakTM 126 127 electromagnetic tracking system at 25 Hz (Polhemus Navigation Science Division, Kaiser Aerospace, Vermont). The FastrakTM system has been used in previous rowing studies,²⁴⁻²⁶ 128 129 and has been reported to be valid and reliable in measuring joint angles in the sagittal plane, reporting average errors of 0.4° using a wooden model positioned on a modified rowing 130 ergometer.²⁷ Three of the device's sensors were secured on the participant's skin overlying 131 the spinous processes of S2, L3 and T12 using double sided tape and Fixomull[®] such that the 132 lower lumbar angle (LLA) and the upper lumbar angle (ULA) could be derived (Figure 133 1).^{20,24,26} A rotary encoder was connected to the flywheel of the rowing ergometer to 134 135 determine the stroke length and stroke rate. Prior to every data collection trial, stroke length 136 was calibrated with the voltage on the rotary encoder and then synchronised with the FastrakTM using a customized Labview software program (Version 8.6.1, National 137 138 Instruments, Texas, USA). This stroke length was used to determine the start and the end of 139 the drive phases, stroke length is shortest at the beginning of the drive phase (catch), and 140 longest at the end of the drive phase (*finish*). Ergometer rowing was chosen for this study as it has been suggested as an aggravating factor to LBP in rowers^{3,4,12,28}, and this has the 141 142 advantage of controlling extrinsic factors such as wind and water condition during data collection.³ 143

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INSERT FIGURE 1 ABOUT HERE

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Participants' maximum slouch angles were determined in static sitting with participants instructed to place their feet flat on the ground; shoulders' width apart with their knees bent to 90°; and their arms crossed in front of the chest. They were then instructed to 'slouch as far down as possible'. They were required to hold these positions for five seconds, and this process was repeated three times with a 30 second rest period between each trial. The
maximum Lower Lumbar Angle (LLA) and Upper Lumbar Angle (ULA) were then
calculated and used to represent maximum slouch. This protocol was used in a previous study
by the authors.²⁴

155 Prior to ergometer testing participants completed a 5-10 minute warm up involving 156 sub-maximal ergometer rowing. Participants rowed for 15 minutes at a stroke rate of 22 157 strokes per minute with a rating of perceived exertion of 17/20. This protocol was designed after consultation between the research team and coaches as this was deemed to be common 158 training practices in the adolescent rowing population. Kinematic data was collected during 159 the last 15 seconds of the 1st minute (start), 7th minute (middle) and 15th minute (end). The 160 161 15-second period equated to three to five full-completed strokes. During the ergometer trial, 162 the Numeric Pain Rating Scale (NPRS), which is an 11-point scale (0-10) to collect selfreported pain intensity,²⁹ were collected verbally at the beginning of every minute of the 163 164 ergometer trial and also at the end of the 15-minute ergometer trial. Participants were advised 165 to cease the ergometer trial if their level of pain during testing exceeded their level of pain 166 during their usual rowing training or competition.

167 A customized LabVIEW program (Version 8.6.1, National Instruments, Texas, USA) converted outputs derived from the 3-Space FastrakTM during the first three completed 168 strokes to flexion and extension angles (angles in the sagittal plane) via matrix algebra 169 procedures as described elsewhere.³⁰ From these procedures, LLA and ULA were derived ²⁴⁻ 170 ²⁶ as shown in Figure 1. For the derived angles, 0° of the LLA is reflected by L3 marker being 171 172 parallel to the S2 sensor and positive values indicated flexion (anterior rotation of the L3 173 sensor over the S2 sensor) while negative values indicated extension (posterior rotation of the L3 sensor over the S2 sensor). Similarly, 0° of the ULA is reflected by the T12 marker being 174 175 parallel to L3 sensor, where positive values indicated flexion (anterior rotation of the T12 176 sensor over the L3 sensor) and negative values indicated extension (posterior rotation of the 177 T12 sensor over the L3 sensor. Consistent with previous research, only sagittal plane angles 178 and data from the drive phase were analysed,^{7,9,11} given that the drive phase is known to be 179 when the spinal load is greatest.¹³ All data in the drive phase were time normalized, with 0% 180 defined as the *catch* and 100% defined as *finish*. Near end-range flexion was defined as 181 above 80% of the maximum slouch angle during the static sitting test.¹⁹

Independent t-tests were used to determine whether age, height, body mass and BMI differed between no pain and pain groups. A linear two level mixed-effects model was used to evaluate the change in NPRS scores reported at baseline and each minute over the 15 minutes of rowing to assess the relationship between rowing and LBP intensity over time.

Flexion angle measures taken at percentiles of the drive phase from three completed 186 187 stokes were averaged to produce a single flexion angle (for both ULA and LLA) for the early (0,10 and 20th percentile), mid (30-70th percentile) and late (80,90 and 100th percentile) drive 188 phase, at the end of the 1st, 7th and 15th minute of rowing. A linear three level mixed-effects 189 190 model was used to determine differences between pain and no pain groups, using the 9 191 repeated measures over drive phase (early/mid/late) nested in minutes (1,7 and 15). 192 Differences in flexion angle across phase and minute were examined and estimates of group 193 difference adjusted for these factors. To examine if the difference in flexion angles between 194 pain and no pain groups became larger over the 15 minutes of rowing, a groupXminute 195 interaction term was evaluated. To examine if the difference in flexion angles between pain 196 and no pain groups were different over the early, mid and late drive phase, a groupXphase 197 interaction term was evaluated.

To evaluate the proportion of drive phase near end range flexion, angular measures (for both ULA and LLA) were sampled at 25Hz for three completed strokes collected during the last 15 seconds of the 1st, 7th and 15th minute of the ergometer rowing. These values were 201 expressed as a percentage of maximum slouch sitting angle, and the proportion of drive phase measures for which this value exceed 80% was calculated then averaged over the three 202 strokes at the 1st, 7th and 15th minute. A linear two-level mixed-effects model was used to 203 204 determine differences between pain and no pain groups, using the 3 repeated measures over the 1st, 7th and 15th minute. Differences in proportion of drive phase near end range flexion 205 206 across minute were examined and the estimate of group difference adjusted for minute. To 207 examine if the difference in proportion of drive phase near end range flexion between pain 208 and no pain groups became larger over the 15 minutes of rowing, a groupXminute interaction 209 term was evaluated. The non-parametric ranks-based Mann-Whitney test was also performed on these measures to test for group difference at the 1st, 7th and 15th minute separately to 210 211 confirm findings were robust to misspecification of the linear mixed models.

212 Models were estimated with and without adjustment for height, weight and age to 213 check for confounding as there was evidence these factors differed between pain and no pain 214 groups. An absence of confounding was assumed if potential confounders were non-215 significant in models at $\alpha > 0.1$; in this case coefficients were estimated without adjustment for 216 these factors. Additionally, although not an a priori objective, a post-hoc comparison of error 217 variances between pain groups in the mixed-effects models was conducted as plotting of the 218 raw data displayed suggested more within-subject variability in data from those subjects with 219 pain (see Results section).

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RESULTS

The demographics of the participants showed that rowers with pain were significantly taller and heavier than rowers with no pain but no differences were found in the age and BMI between the two groups (Table 1). There were no statistically significant differences in the maximum slouch angles during the static sitting trial between groups, rowers in the pain group postured their LLA at 3.2° (17.5 °) compared to 3.7° (7.8 °) in the no pain group (95%CI: -13.2° to 12.3°, p=0.942) and their ULA at 4.6° (8.1°) compared to 2.6° (11.1°) in the no pain group (95%CI: -7.2° to 11.1°, p=0.656).

Numeric Pain Rating Scale scores increased significantly over the 15 minutes of rowing from 1.7 (95%CI: 1.0 to 2.3) at baseline to 7.8 at the 15^{th} minute (95%CI: 7.10 to 8.42), with the rate of increase estimated to be 0.41 per minute (95%CI: 0.38 to 0.44, p<.001) in rowers with LBP (Figure 2). All rowers in the no pain group reported 0 NPRS at each minute of the ergometer trial.

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INSERT TABLE 1 AND FIGURE 2 ABOUT HERE

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238 No significant differences were observed in the mean LLA between groups (Table 2). 239 Adjustment for height, weight and age revealed no confounding of group differences and 240 results are presented unadjusted for these factors to maximise precision of estimates. The LLA for each subject for the early, mid and late drive phase over the 1st, 7th and 15th minute 241 242 separately for each pain group are presented (Figure 3). Significant main effect for phase (p<.001) and no evidence of interaction between pain group and phase (p=.821), with flexion 243 244 decreasing from early, mid to late phase similarly in both groups (Table 3). A significant 245 main effect for the pain group was not detected (p=.688), although an interaction between 246 minute and pain group was detected (p=.012), with the pain group displaying more extension (adjusted for phase) in the 15th minute compared to the 1st minute, whereas the no pain group 247 248 displayed similar LLA at all three time points (Table 3). Examination of the raw data plotted 249 suggested more within-subject variability in changes over minute in the pain group, with 250 relatively large changes occurring in both directions, compared to a consistent pattern of no

251	change in the no pain group (Figure 3). Therefore, this was formally tested by comparing the
252	variance of the error terms in the mixed effects model. These were significantly different,
253	with the standard deviation for the pain group being greater $[10.6^{\circ} (95\% CI: 9.4^{\circ} \text{ to } 12.8^{\circ})]$
254	than the no pain group $[4.0^{\circ} (95\% \text{CI: } 3.4^{\circ} \text{ to } 4.7^{\circ})]$, indicating significantly greater within-
255	subject variability in the pain group data.
256	
257	INSERT FIGURE 3 ABOUT HERE
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259	INSERT TABLE 2 AND 3 ABOUT HERE
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261	No significant differences were observed in the mean ULA between groups. Analysis
262	using linear mixed effects model identified no effect for minute (p=.526) and no group by
263	minute interaction (p=.774). The means and standard deviations for ULA by phase, minute
264	and pain/no pain group are presented (Table 2). Adjustment for height, weight and age
265	revealed no confounding of group differences and results are presented unadjusted for these
266	factors to maximise precision of estimates. Raw data for ULA for each subject over the early,
267	mid and late drive phase by 1st, 7th and 15th minute, separately for each pain group are
268	presented (Figure 4). Although there was evidence that groups differed by phase (p<.001),
269	the estimated group difference was not statistically significant at any phase (table 3). There
270	was a significant interaction between phase and group, meaning the degree of change over
271	phase was estimated to differ by group, with a pattern of significantly more extension over
272	early, mid and late phase evident in both groups (Table 3), but to a significantly lesser extent
273	in the pain group. Raw data plotted in Figure 4 suggests more within-subject variability in
274	changes over phase in the pain group, with less consistent pattern of increasing extension
275	over the drive phase compared to the consistent pattern seen in the no pain group. This was

formally tested by comparing the variance of the error terms in the mixed effects model. These were significantly different, with the standard deviation for the pain group being greater (4.9° (95%CI: 4.0° to 6.0°) than the no pain group (2.8° (95%CI: 2.4° to 3.3°)), indicating significantly greater within-subject variability in the pain group data.

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INSERT FIGURE 4 ABOUT HERE

281 No statistically significant differences were observed in the LLA in the proportion of 282 drive phase in near or end of range flexion between groups. The raw means and standard 283 deviations for the proportion of drive phase near end range LLA flexion by minute and pain/no pain groups (Table 4). This data are presented graphically for each subject over 1st, 284 7th and 15th minute, separately for each group (Figure 5A). Analysis using a linear mixed 285 286 effects model detected evidence of an association between a lesser proportion of drive phase 287 spent in flexion with increasing age and (weight-adjusted) height (Table 5). No effect for 288 minute (p=.872) and no group by minute interaction was observed (p=.284). The pain group 289 was estimated to spend less time of drive phase in near end range when compared to no pain 290 group, adjusted for minute, age, height and weight (-.27, 95%CI: -.59 to .04, p=.087, Table 3) 291 but this difference was not statistically significant. The raw data plotted displays suggest 292 greater degree of variability in the proportion of drive phase near end range LLA flexion in 293 the pain group (Figure 5A), with less consistent patterns over time in the pain group. Again, 294 this was formally tested by comparing the variance of the error terms in the mixed effects 295 model. These were significantly different, with the standard deviation for the pain group 296 being greater (.31 (95%CI: .23 to .42) than the no pain group (.06 (95%CI:.04 to .08), 297 indicating significantly greater within-subject variability in the pain group data. 298 Nonparametric analysis of this data also did not detect a difference in proportion of drive phase in near end range LLA in the pain group at the 1^{st} , 7^{th} or 15^{th} minute (Mann Whitney test, p= .341, .272 and .702 respectively).

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INSERT FIGURE 5A AND 5B ABOUT HERE

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INSERT TABLE 4 ABOUT HERE

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306 Rowers in the LBP group were found to spend a significantly greater proportion of 307 the drive phase near the end of range of ULA flexion compared to the no-LBP group. The 308 raw means and standard deviations for the proportion of drive phase near end range ULA 309 flexion by minute and pain/no pain groups (Table 5). This data is presented graphically for each subject over 1st, 7th and 15th minute, separately for each pain group (Figure 4B). 310 311 Analysis using a linear mixed effects model detected no evidence of an association between a 312 lesser proportion of drive phase spent in ULA flexion with increasing age (β =.00, 95%CI: -313 .06 to .06, p=.974) and (weight-adjusted) height (β =-.01, 95%CI: -.02 to .01, p=.144), unlike 314 results for LLA, and models were estimated unadjusted for these factors. No effect for minute 315 (p=.548) and no group by minute interaction were observed (p=.226). The pain group was 316 estimated to spend a greater proportion of the drive phase in near end range ULA than the no 317 pain group (.19, 95%CI: .03 to .35, p=.021, Table 3). The raw data suggests a greater degree 318 of within-subject variability generally in the proportion of drive phase near end range for 319 ULA flexion versus LLA, with more inconsistent patterns over time in both groups for ULA 320 than those for LLA (Figure 5B). The standard deviation of the residuals for the pain group 321 (.29 (95%CI: .21 to .39) were comparable to the no pain group (.19 (95%CI: .14 to .26). 322 Nonparametric analysis of this data confirmed a significantly greater proportion of drive

323	phase in near end range ULA in the pain group at the 7 th minute (Mann Whitney test, p=.002)
324	but not the 1^{st} (p=.160) or 15^{th} minute (p=.650).
325	
326	INSERT TABLE 5 ABOUT HERE
327	
328	DISCUSSION
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330	The results of this study demonstrate that 15 minutes of ergometer rowing results in
331	increasing intensity of LBP over time in male adolescent rowers with rowing reporting
332	related LBP (Figure 2). Although no significant differences were detected in the mean LLA
333	and ULA between rowers with and without LBP, rowers with pain did demonstrate less ULA
334	excursion and ULA into extension compared to rowers without pain over time.
335	This increase in pain intensity may reflect a temporal summation of pain, where a
336	repetitive stimulus on pain sensitive structures may cause a gradual increase of pain

sensation.^{31,32} A similar pattern of pain summation has been reported previously in cyclists 337 with LBP during a 2-hour cycling trial.¹⁹ There is debate regarding the underlying 338 mechanism for this phenomena, with some researchers suggesting that it reflects inhibitory 339 and facilitatory mechanisms in the central nervous system,³³ whilst other authors suggest 340 provocative movement behaviours may result in repeated stress on sensitized tissues with a 341 resultant summation of pain.^{14,15} In reality a combination of both of these factors may 342 343 interplay.

On average, rowers in the pain group maintained their ULA in flexion throughout the 344 drive phase [early (9.1°) mid (5.7°) and late (1.0°)] compared to rowers without pain who 345 moved into more extension in the late phase [early (10.5°) mid (6.4°) late (-3.0°)]. In addition, 346 rowers with LBP postured their ULA within 80% of end range flexion for a greater 347

348 proportion of the drive phase than rowers without LBP (mean diff .19, p=0.021). The 349 increased proportion of drive phase spent in flexion by the rowers with LBP in this study is 350 consistent with our hypothesis and may be reflective of a flexion loading strain mechanism for low back pain.³⁴ Previous studies have reported that both adolescent and adults with LBP 351 352 provoked by lumbar 'flexion' movements and postures have a tendency to posture their spines closer to end range flexion during sitting ^{16,35,36}. Similarly, cyclists with LBP have 353 been identified to maintain either lower lumbar spine in a more flexed position ¹⁵ or cycle 354 closer to end range of flexion in the lower lumbar spine.¹⁹ It may be that inability to maintain 355 356 the lumbar spine away from end of range leaves the spine more vulnerable to flexion loading 357 strain in sports where the lumbar spine is exposed to cyclical or sustained loading.

It was hypothesized that adolescent male rowers with LBP would posture their LLA 358 359 and ULA in more flexion than rowers without LBP during the drive phase of ergometer 360 rowing, and this difference would increase over 15 minutes of rowing. Although no 361 differences in the mean LLA and ULA were detected overall or within the early, mid and late phase or 1st, 7th and 15th minute, on examination of the raw data it was noted that rowers with 362 LBP had greater within-subject variability in LLA and ULAs compared to rowers without 363 LBP. This is a preliminary finding that was not an a priori aim of the study and therefore 364 further investigation is warranted. The within-subject variability in spinal kinematics in 365 individuals with LBP is not a new concept, with higher variability in spinal movement during 366 functional tasks reported in adults with chronic LBP compared to no-LBP. ^{37,38} This may be 367 368 due to altered peripheral and central sensory processing of the nervous system, resulting in poorer spinal position sense in adolescents and adults with LBP ^{39,40}, with a tendency to 369 370 either under or over shoot a neutral sitting posture during a lumbar spine reposition test, a 371 mechanism proposed to increase end range strain. Holt and associates (2003) have also 372 reported variations in spinal kinematics in athletes with and without LBP over a 60-minute ergometer trial,¹¹ but no direct comparisons were made between the participants with and
without a history of LBP.

375 We acknowledge the following potential limitations of this study. 1) The large variation 376 reported in the kinematics of the pain group participants may explain the lack of significant differences detected in the mean LLA and ULA between the LBP and the no-LBP group. 2) 377 378 A subjective indicator of rowing effort (RPE) was used in the study rather than an objective 379 measurement of subjects' effort throughout the trial such as power output as it was 380 commonly used in this age group to measure work rate in this group of rowers. Although 381 differences in work rate will exist, the authors feel that this would be minimal as stroke rate 382 was standardised between groups and unlikely to invalidate comparisons between groups. 3) 383 In light of the current finding regarding variability, the analysis of a larger number of strokes 384 and statistical procedures could be considered to evaluate spinal kinematics of rowers with 385 LBP. 4) It is also acknowledged that assessing end range slouch position in the LBP subjects 386 could have been influenced by the presence of pain, although there was no report of 387 discomfort or observable movement guarding during this aspect of the testing. Further, no 388 differences were detected in the maximum slouch angles between groups. Cross sectional 389 studies do not give clear insight to causation requiring the need for future longitudinal studies 390 in order to determine whether kinematic differences precede or follow low back pain in male 391 adolescent rowers.

In conclusion, rowers with LBP positioned their upper lumbar spine nearer end range flexion for a greater proportion of the drive phase and demonstrated greater individual variation in spinal movement than rowers without LBP. These findings may have implications for coaching practices and targeted interventions to improve consistency in rowing technique and avoid prolonged end of range spinal loading so as to minimize the potential for end range sensitization of spinal structures.

399 Acknowledgements

- 400 The authors would like to acknowledge the Senior Research Assistant at the School of
- 401 Physiotherapy and Exercise Science for his technical support in this project.

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TABLES

TABLE 1 – Mean and standard deviation of characteristics in each group and the mean, standard error and p-value of differences between the no pain and pain group.

Characteristic	No Pain	Pain	Mean	95% CI	P value
	(n=10)	(n=10)			
Age (years)	17.2 (1.4)	16.0 (1.2)	1.2	-0.1, 2.4	.074
Height (m)	1.85 (0.08)	1.70 (0.09)	0.15	-0.2, -0.1	<.001
Weight (kg)	78.2 (12.9)	66.8 (10.8)	11.5	-22.9, 0.0	.050
BMI (kg/m ²)	23.1 (3.4)	22.8 (3.8)	0.3	-2.7, 3.4	0.818
VAS (/10)	0 (0)	4.6 (1.1)			
PSFS (/30)	n/a	17 (6.1)			
RMDQ (/22)	n/a	3.5 (2.1)			

BMI = Body Mass Index; VAS = Visual Analogue Scale; PSFS = Patient Specific; Scale; RMDQ = Roland Morris Disability Questionnaire.

		No Pain			Pain	
Minute	Early	Mid Phase	Late	Early	Mid Phase	Late
	Phase		Phase	Phase		Phase
			Lower lumb	oar angle (°)		
1	8.8 (6.7)	3.7 (7.4)	-4.2 (11.1)	9.3 (16.2)	7.7 (10.0)	3.5 (11.5)
7	8.7 (7.0)	2.9 (7.5)	-2.8 (9.8)	11.5 (9.6)	7.6 (9.6)	1.9 (10.8)
15	8.8 (7.4)	2.9 (8.3)	-3.0 (11.1)	6.9 (21.4)	-1.1 (18.1)	-8.2 (21.9)
			Upper lumb	ar angle (°)		
1	8.6 (7.1)	5.4 (8.0)	-4.8 (7.7)	8.2 (7.2)	5.4 (7.6)	1.2 (9.3)
7	11.2 (6.1)	6.6 (6.7)	-2.4 (8.1)	9.4 (8.4)	6.3 (11.2)	1.2 (14.0)
15	11.8 (6.3)	7.1 (6.6)	-1.8 (8.2)	9.8 (10.1)	5.5 (14.7)	0.6 (17.1)

TABLE 2 - Mean and standard deviation of the lower and upper lumbar angles for drive
 phases over 1st 7th and 15th minute, for Pain and No Pain group.

		Marginal	β	95% CI	p-value
		means $(^{0})$	coefficient(⁰)		
			(i.e. contrast)		
		Lower	lumbar angle		
Group (Pain – No Pain	1)	_			
At Minute 1:	NP	2.8			
	Р	6.8	4.1	-3.8 to 12.0	.313
At Minute 7	NP	3.0			
	Р	7.0	4.0	-3.9 to 12.0	.318
At Minute 15	NP	2.9	2 7	11 (+ 4 2	250
	Р	-0.8	-3.7	-11.6 to 4.2	.358
Phase (ref to Early Ph	ase)	_			
	Farly	94			
	Mid	3.9	-5.5	-74 to -36	< 001
	Late	-2.5	-11.9	-13 8 to -10 0	< 001
	Lute	2.0	11.9	15.6 to 16.6	
Minute (ref to Minute	1)	_			
No Pain Group	Min 1	2.8			
	Min 7	3.0	0.2	-1.8 to 2.2	.857
	Min 15	2.9	0.1	-1.9 to 2.1	.903
Pain Group	Min 1	6.8			
	Min 7	7.0	0.1	-5.4 to 5.7	.961
	Min 15	-0.8	-7.7	-13.2 to -2.1	.007
Crown (Dain No Dair	•)	Upper lumb	ar angle		
Group (Pain – No Pain	1) ND	10.5			
At Phase I	NP D	10.5	1 /	8 0 to 5 2	697
At Phase 2	r ND	9.1	-1.4	-8.0 10 3.2	.082
At I hase 2	P	0.4 5 7	-0.6	-7.2 to 6.0	8/19
At Phase 3	NP	-3.0	-0.0	-7.2 to 0.0	.047
Trt Thuse 5	P	1.0	4.0	-2.6 to 10.6	.233
	-	1.0		2.0 00 10.0	
Phase (ref to Early 1)		-			
No Pain Group					
	Early	10.5			
	Mid	6.4	-4.2	-5.6 to -2.7	<.001
	Late	-3.0	-13.5	-15.0 to -12.1	<.001
Dain Carro					
Pain Group	Early (1)	0.1			
	Early (1)	9.1 5 7	2 1	50 to 10	007
	$\frac{1}{1} \frac{1}{2} \frac{1}$	J./ 1.0	-3.4 _8 1	-3.910 - 1.0 -106 to 57	.007
	Late (3)	1.0	-0.1	-10.0 10 -5.7	<u><u></u>,001</u>
Minute (ref to Minute	1)	_			
	 Min 1	3.9			
	Min 7	5.4	1.5	-1.6 to 4.6	.358
	Min 15	5.6	1.6	-1.5 to 4.7	.302

TABLE 3 - Mixed model coefficients for lower and upper lumbar angle.

579 TABLE 4 - Percentage of drive phase in greater than 80% of flexion range for Lower and580 upper angle, for Pain and No Pain group.

	Lower Lumbar Angle (%)		Upper Lumba	r Angle (%)
Minute	No Pain	Pain	No Pain	Pain
1	0.56 (0.34)	0.69 (0.36)	0.45 (0.33)	0.68 (0.36)
7	0.58 (0.34)	0.62 (0.38)	0.48 (0.17)	0.77 (0.17)
15	0.58 (0.34)	0.49 (0.46)	0.48 (0.16)	0.52 (0.38)

TABLE 5 - Mixed model results for proportion of drive phase in >80% lower and upper
 lumbar end range flexion.

		Marginal	β	95% CI	p-value
		means $(^{0})$	coefficient		
			$(^{0})$		
			(contrast)		
	Lo	wer Lumbar	Angle		
Group (Pain - No Pain	n)	_			
	NP	.72			
	Р	.45	27	59 to .04	.087
Minute (ref to Minute	_				
	Min 1	.58			
	Min 7	.59	.01	04 to .06	.647
	Min 15	.59	.01	04 to .06	.657
Covariates		-			
Age (yrs)	16.6 ^a	.59	10 ^b	20 to01	.036
Height (cm)	177.6 ^a	.59	02 ^b	04 to00	.030
Weight (Kg)	72.5 ^a	.59	.01 ^b	.00 to .02	.080
Upper Lumbar Angle					
Group (Pain - No Pain	n)				
	NP	.47			
	Р	.66	.19	.03 to .35	.021
Minute (ref to Minute	1)	_			
	Min 1	.56			
	Min 7	.60	.04	09 to .19	.509
	Min 15	.53	03	17 to .11	.668

^amean of covariate in the sample

 ^b β coefficient represents the expected change in proportion of drive phase spent in >80% end 588 range flexion with each increase of one unit in the covariate

592	Figure Caption
593	FIGURE 1 – Regional lumbar kinematics (ULA – Upper Lumbar Angle; LLA – Lower
594	Lumbar Angle)
595	
596	FIGURE 2 – Group mean and standard deviation of low back pain intensity scores
597	(measured by Numeric Pain Regional Scale) during the 15-minute rowing ergometer trial.
598	
599	FIGURE 3 – Lower lumbar angle for each subject over the 1 st , 7 th and 15 th minute, for the
600	early, mid and late drive phase separately, in pain and no pain groups separately.
601	
602	FIGURE 4 – Upper lumbar angle for each subject over the drive phase separately for 1 st , 7 th
603	and 15 th minute, in pain and no pain groups separately.
604	
605	FIGURE 5A: Proportion of drive phase lower lumbar angle in greater than 80% flexion over
606	1st, 7th and 15th minute, in pain and no pain groups separately
607	
608	FIGURE 5B: Proportion of drive phase upper lumbar angle in greater than 80% flexion over
609	1st, 7th and 15th minute, in pain and no pain groups separately
610	
611	
612	



Regional lumbar kinematics (ULA – Upper Lumbar Angle; LLA – Lower Lumbar Angle) 71x77mm (300 x 300 DPI)



Group mean and standard deviation of low back pain intensity scores (measured by Numeric Pain Regional Scale) during the 15-minute rowing ergometer trial. 60x36mm (300 x 300 DPI)

(D)



FIGURE 3 – Lower lumbar angle for each subject over the 1st, 7th and 15th minute, for the early, mid and late drive phase separately, in pain and no pain groups separately. 99x99mm (300 x 300 DPI)



FIGURE 4 – Upper lumbar angle for each subject over the drive phase separately for 1st, 7th and 15th minute, in pain and no pain groups separately. 270x270mm (72 x 72 DPI)





FIGURE 5B: Proportion of drive phase upper lumbar angle in greater than 80% flexion over 1st, 7th and 15th minute, in pain and no pain groups separately

372x270mm (72 x 72 DPI)