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TITLE PAGE

Sodium bicarbonate supplementation delays neuromuscular fatigue without changes in

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1 ABSTRACT

2 **Purpose:** To investigate the development of neuromuscular fatigue during a basketball game 3 simulation and ascertain whether sodium bicarbonate (NaHCO₃) supplementation attenuates any neuromuscular fatigue that persists. Methods: Ten participants ingested 0.2 g.kg⁻¹ of 4 5 NaHCO₃ (or an equimolar placebo dosage of sodium chloride [NaCl]) 90 and 60 minutes prior to commencing a basketball game simulation (ALK-T vs PLA-T). Isometric maximal voluntary 6 7 contractions of the knee extensors (MVIC) and potentiated high (100 Hz) and low (10 Hz) 8 frequency doublet twitches were recorded before and after each match quarter for both trials. 9 In addition, 15 m sprint times and layup completion (%) were recorded during each quarter. **Results:** MVIC, 100 and 10 Hz twitch forces declined progressively in both trials (P<0.05) 10 with a less pronounced decrease in MVIC during ALK-T (P<0.01). Both 100 and 10 Hz twitch 11 forces were also significantly greater in ALK-T (P<0.05). 15 m sprint time increased over the 12 course of both trials (~2%, P<0.01); however, no significant condition or time effect was found 13 for layup completion (P>0.05). Conclusion: A basketball simulation protocol induces a 14 substantial amount of neuromuscular (reduction in knee extensor MVICs) and peripheral 15 16 fatigue with a concomitant increase in 15 m sprint time over the protocol. NaHCO₃ supplementation attenuated the rate of fatigue development by protecting contractile elements 17 of the muscle fibres. Practical Applications: This study provides coaches with information 18 19 about the magnitude of fatigue induced by a simulated basketball game, and provides evidence 20 of the efficacy of NaHCO₃ in attenuating fatigue.

21

22 KEY WORDS

23 Alkalosis; muscular fatigue; peripheral fatigue; team sports

25 INTRODUCTION

Basketball matches are characterised by a large volume of short duration, high intensity 26 movements as shown via time-motion analysis (25). Simulated games can also raise mean 27 oxygen uptake ($^{\downarrow}O_2$) and heart rate (HR) values to approximately 65% and 85% of their 28 29 maximum, respectively (25, 28). Due to the elevated metabolic demand of a basketball game, 30 a build-up of deleterious metabolites (i.e. H⁺, Pi) may reduce force producing capacity of the working muscles (2). This is deemed neuromuscular fatigue, and is defined in the present work 31 as any transient, exercise-induced reduction in muscular force generating capacity (42), with 32 underpinning mechanisms of peripheral or central origins. Previous research has shown that 33 explosive power and sprint ability is reduced following basketball-related activity (9). 34 However, to our knowledge, no study has yet investigated the time course of various sites of 35 neuromuscular fatigue during a simulated basketball match, and the efficacy of a potentially 36 ergogenic supplement in ameliorating the aforementioned fatigue by reducing metabolite 37 38 accumulation.

39

40 As a result of high intensity exercise (such as basketball), extra and intra-cellular ionic concentrations are altered within the muscles, causing reduced contractile performance. For a 41 42 complete review of the processes contributing to peripheral fatigue see Allen et al (2). Examples of factors involved in impairment within the muscular contractile apparatus are reduced 43 44 intracellular potassium ion (K⁺) concentrations caused by efflux into the interstitial spaces, 45 resulting in extracellular accumulation (19). This negatively affects the capacity of the 46 sarcolemma to propagate action potentials (27). This potentially occurs as Na⁺, K⁺ ATPase activity is inhibited by the increased presence of hydrogen ions (H⁺) (acidosis). 47

48 Similarly, acidosis inhibits myofibril ATPase activity, leading to reduced calcium ion (Ca²⁺) 49 reuptake to the sarcoplasmic reticulum (SR) and consequently less Ca²⁺ released from the SR 50 when prompted by an action potential (7, 22).

51

52 In this study, contractile function was measured using potentiated, electrically-evoked paired 53 twitches at two different frequencies (10 and 100 Hz) in an attempt to refine the sites of 54 peripheral fatigue development. The mechanical response (twitch) to low frequency doublets (10 Hz) has been shown to be modulated by the extent of Ca^{2+} release from the SR (19, 21). 55 High frequency doublets (100 Hz), and their respective twitch amplitude, have been shown not 56 to be affected by moderate decreases in Ca^{2+} . Therefore decreases in high frequency twitch 57 amplitude reflect attenuated action potential propagation caused by extracellular K⁺ 58 accumulation (21). The ratio of low:high frequency twitch forces gives detail about the 59 aetiology of contractile decline during a fatiguing task (27). 60

61

The role acidosis plays in the development of neuromuscular fatigue during high intensity 62 exercise (such as basketball) remains under debate (10, 44). However, it is generally agreed 63 that athletes who perform high intensity exercise (such as basketball players) would likely 64 benefit from NaHCO₃ supplementation (8) as it attenuates the aforementioned negative effects 65 of acidosis. For instance, a reduction in extracellular accumulation of K⁺ during exhaustive 66 exercise has been evidenced following NaHCO₃ supplementation (40). NaHCO₃ 67 supplementation also attenuates the inhibiting effects of H^+ by increasing the intra – 68 69 extracellular pH gradient. This allows for greater efflux of deleterious metabolites outside the 70 muscle cells and attenuates their harmful effects on the contractile function (26).

72 NaHCO₃ has been shown to enhance high-intensity performance (5, 6) and delay the development of neuromuscular fatigue during a fatiguing task (38). The ergogenic effects found 73 74 in the aforementioned laboratory-based studies give rationale for the investigation of NaHCO3 75 as an ergogenic aid during high-intensity team sport activity such as basketball. Interestingly, 76 only a limited amount of studies have investigated the effect of NaHCO₃ supplementation on performance outcomes during simulated game based protocols (1, 23, 31, 34), and 77 78 neuromuscular function has never been assessed throughout a simulated basketball match. 79 Afman et al (1) recently found a beneficial effect of NaHCO₃ supplementation on 15m sprint 80 times, but not layup completions, during a modified Loughborough Intermittent Sprint Test (LIST), which was validated to replicate the demands of a 40-min basketball game. Therefore, 81 82 the present study aims to investigate the development of neuromuscular fatigue and more 83 specifically, the peripheral mechanisms during a basketball game simulation. The study also aims to ascertain whether NaHCO₃ supplementation attenuates this development. It was 84 hypothesised that there would be a significant decline in both voluntary force generating 85 capacity of the knee extensors and the amplitude of evoked paired-twitches. It was hypothesised 86 87 that this decrease in contractile function would lead to faster 15-m sprint times throughout the protocol and with smaller declines in the supplement (ALK-T) compared to the placebo (PLA-88 89 T) trial.

- 90
- 91 **METHODS**

92 Experimental Approach to the Problem

Participants visited the laboratories on three separate occasions. Three and seven days separated
 familiarisation and 1st fatiguing trial, and 1st and 2nd fatiguing trials respectively, to ensure full
 washout of the supplement/placebo (5). Familiarisation involved a neuromuscular function

96 assessment performed on an isokinetic dynamometer, followed by one block of the modified 97 LIST protocol. For the two experimental trials, participants performed four blocks of the 98 modified LIST with neuromuscular assessment prior to, and following each block of the LIST 99 (1). Participants were asked to avoid consuming any stimulants or alcohol, and to replicate food 100 intake during a 24-hour period before testing. The study was a double-blind

crossover design with exposure to supplements rando mized and counterbalanced. Each participant received extensive information, and signed an informed consent form ar a dical questionnaire after they had the opportunity to ask any questions to researcher. The protoc was approved by the University Ethics committee and adhered to the Dromation of the distance.

Subjects

Ten healthy and active male basketball players volt deered take part in the study (age 21 ± 1 years; height: 182 ± 5 cm; weight: 81.5 ± 8 d (1) participants had over 4 years of competitive basketball experience.

Procedures

Neuromuscular Fun on Assess ont

For the neuropscular as the entropy of the right knee extensors, participants sat on the Con-Trex Multi-Just science Con-Trex, Dubendorf, Switz erland) as per the published reliability study (30) (~85 mip angle; distal dynamometer's shin pad attached 2 -3 cm proximal to the lateral malleolus with a strap around the shank; straps were fastened and locked across chest and pelvis; movement resisting pad over the mid-thigh of the contracting leg). Knee angle was kept at 90° for all maximal voluntary isometric contractions (MVICs) and twitches.

105 Torque measurement was corrected to take gravity effect into account. Participants were 106 instructed to cross their arms across their chest and were provided with visual feedback of force 107 during the protocol.

108 A 48 mm² self-adhesive cathode electrode (CF3200, Nidd Valley Medical Ltd, Harrogate, UK)

109 was placed directly over the femoral nerve in the femoral triangle with the anode placed directly

110 onto the greater trochanter of the femur (Prottens, Bio Protech Inc, Korea).

111 Percutaneous electrical stimulation was delivered by a constant-current stimulator (DS7A,

112 Digitimer, Letchworth Garden City, Great Britain). Stimulations were triggered manually using

113 a PowerLab 15T (Model ML818, AdInstruments Pty Ltd, Dunedin, New Zealand) and force

114 production was recorded using LabChart 7 software (AdInstruments Pty Ltd, Dunedin,

115 New Zealand). Sprint times (15-m) were recorded using wireless electronic timing gates (TC

116 Timing System, Brower, Utah, USA). Participants began each sprint from a standing start 10cm

117 behind the timing gates (see figure 1).

118 Familiarisation Session

Single electrical 200µs impulses were delivered to the right femoral nerve via the surface 119 electrode. Percutaneous single stimuli were delivered at 10 mA increasing by 10mA until a 120 plateau in twitch force amplitude was reached. This intensity was increased to 130%, to ensure 121 supramaximal stimulations were delivered (mean intensity: 170 ± 35 mA). This process was 122 123 repeated before each experimental visit. The MVIC familiarisation protocol then consisted of 124 2 and then 3×5 -s voluntary contractions performed at 50% and 75% of maximal subjective 125 effort, respectively. The participants then performed 3×5 -s MVICs. Each maximal contraction 126 was followed by two doublet stimulations (100 Hz and 10 Hz) in 1-s

127 intervals.

130 Experimental Protocol

- 131 Neuromuscular baseline tests were performed followed by a short standardised warm up (a
- 132 4length jog of the basketball court). After baseline and warm ups, participants completed four
- 133 blocks of 11 repetitions of the modified LIST shown in figure 1 (1), meaning 11 sprints and

layups were performed per quarter. Participants had 5 minutes rest between quarters, in which neuromuscular fatigue assessment was performed. Three 5 s MVICs with 6 between were performed with two doublet stimulation s (10 Hz and 100 Hz) for ving the contractions in 1 s intervals. Due to the time tak en to move from be actively court dynamometer following each quarter, the timing of the first MVL was a dardise to 75 s.

FIGURE 1 HERE

Supplement

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136

137

ior to commencement of the protocol in order to consume Participants arrived 90 minutes ceb o; NaHCO₃ was delivered in two separate the first half of either the pple t or with the ond dosage consumed 60 minutes prior. dosages of 0.2 g.kg NaHCO₃ was al each, totalling 0.4 g.kg ⁻¹. Sodium calorie-free cordi dissolved in 500 ml of proposed of two 0.138 g.kg⁻¹ dosages dissolved in 500 ml water and chloride (pla bo) cordial each (e. molar amount of sodium to account for alterations in Na⁺ handling; for more details, see (20)). The same amount of supplem ent/placebo was consumed 60 minutes prior to exercise. A similar ingestion protocol has been shown to benefit prolonged intermittent activity with no reported incidences of gastrointestinal disturbances following NaHCO3 supplementation, as did the present study (5).

139 Data Analysis

140 The maximum 500-ms value was recorded as maximal force for each MVIC plateau, and the 141 peak twitch amplitude was computed for each doublet stimulation. The greatest value over each 142 set of three MVICs and twitches was subsequently recorded for each time point.

- 143 Coefficient of variations between the 3 measures were $2.6 \pm 2.0\%$ for MVIC, $3.5 \pm 2.7\%$ for
- 144 100 Hz twitch, and $3.0 \pm 2.2\%$ for 10 Hz twitch. Each 15 m sprint time was recorded in seconds
- 145 (s). Successful completions for the layups were expressed as a percentage of total number of
- 146 attempts per quarter (out of 11).

147 Statistical Analysis

Normal distributions were verified (Kolmogorov-Smirnov test) and one-way (1 x 5) repeated 148 measures ANOVAs were run to assess the change in neuromuscular variables (MVC, 100Hz, 149 10 Hz twitches) and quantify the magnitude of fatigue elicited over the course of the placebo 150 trial (Baseline, Q1, Q2, Q3, Q4). A two-way (2×5) repeated measures ANOVAs was 151 performed to test for between condition (ALK-T vs PLA-T) and time differences (Baseline, 152 Q1, Q2, Q3, Q4). If sphericity assumption was violated (Mauchly's test) then Fratios were 153 adjusted according to the Greenhouse-Geisser procedure. Significant effects of ANOVAs were 154 155 followed up using the Bonferroni-corrected pairwise post hoc test.

- 156 Significance was accepted at $P \le 0.05$ and all data is presented as mean \pm standard deviation 157 (SD). All statistical analyses were performed using SPSS (version 20, Chicago, USA).
- 158
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- 161

163 **RESULTS**

- 164 TABLE 1 HERE
- 165 FIGURE 2 HERE

166

167 MVIC force ($F_{(4,36)} = 42.0, P < 0.01$), decreased significantly over time but with no significant difference between conditions (P > 0.05) (Table 1 and Figure 2). The loss of 168 MVIC was less pronounced during ALK-T as shown by the significant time × condition 169 interaction ($F_{(4,36)} = 6.88, P < 0.01$). However, post-hoc tests did not reveal a significant 170 171 difference between trials at any time points (P > 0.05). The one way ANOVA showed that during PLA-T, the decrement in MVIC ($F_{(4,36)} = 36.9, P \le 0.01$) was progressive from 172 baseline to the 3rd quarter (P < 0.05), with a plateau occurring thereafter (P > 0.05). 173 174 100 Hz twitch ($F_{(4,36)} = 20.25$, P < 0.01) and 10 Hz twitch ($F_{(4,36)} = 24.3$, P < 0.01) also 175 decreased significantly over time. No time × condition interaction effect was found for either 176 evoked twitches (100 Hz: $F_{(4,36)} = 0.76$, P = 0.56; 10 Hz: $F_{(4,36)} = 1.30$, P = 0.29). 100 Hz and 177 10 Hz evoked twitch forces were both greater throughout the protocol in ALK-T (condition 178 effect: 100 Hz: $F_{(1,9)} = 11.8$, P < 0.01; 10 Hz: $F_{(1,9)} = 8.77$, P < 0.05). The one way ANOVA 179 showed that during PLA-T, 100 and 10 Hz twitches were not different from baseline (P >180 (0.05) until after the second quarter from which time point a reduction was significant (P < 181 182 0.05). No time or condition effect was observed for 10:100 Hz twitches ratio (P > 0.05). 183 No condition or interaction effect were found for either of the performance variables (P >184

185 0.05) but the 15-m sprint times became significantly slower over both trials (time effect: $F_{(3,27)}$

186 = 9.39, P < 0.01). The participants' sprints in both trials were systematically slower from one

187 quarter to the next (P < 0.05, Table 1). When comparing first vs last quarter sprint times, both

188 ALK-T and PLA-T were significantly longer (ALK-T: -1.7%, $F_{(3.7)} = 4.3$, P < 1.3

- 189 0.01; PLA-T -2.4%, $F_{(3.7)} = 9.3$, P < 0.05).
- 190

191 **DISCUSSION**

192 To our knowledge this is the first study reporting development of neuromuscular fatigue during 193 simulation of a basketball match. Maximal force production of the knee extensors (MVIC) 194 during PLA-T reduced throughout the first three quarters of the simulated match

(Figure 2; Table 1; ~5% loss per quarter) with no further reduction in the final quarter. Peripheral fatigue was evident from the 2^{nd} quarter of the protocol with disturbances of contractile properties. The ~15% reduction in MVIC torque recorded post 3^{rd} and 4^{th} quarter in this study is similar to the ~15% reduction reported after a 60-min squash match (12), and within the ~11% (11, 13, 29) to ~20% range (16, 17) reported for laboratory based studies investigating repeated sprint activity (4-to 10-s sprints, 8-12 repetitions, 10- to 30-s passive recovery).

202

203 To our knowledge, this is also the first study applying femoral nerve stimulations to assess 204 mechanisms of peripheral fatigue during a basketball game simulation. The ~15% reductions in evoked twitch forces from baseline for both doublet stimulations are similar to those 205 206 previously reported in laboratory-based studies following repeated sprint exercise (9-15%; (29, 207 32)). In the present study, changes in the several mechanisms of contractile function impairment seem to adopt a similar time course with a decrease after the 2nd quarter, and no 208 further decrease thereafter (apart from one occurrence: 10 Hz twitch between quarter 2 and 3, 209 P = 0.01). Perrey et al (29) found decreases of 15% in low frequency (20 Hz) twitch forces but 210

of only 8% in the high frequency (80 Hz) twitch forces following repeated sprints. Their decreased ratio of low:high frequency evoked twitches (-9%) suggested that muscle fibre excitability was the predominant cause for the impairment of the contractile function. In contrast, the present study found decreases of similar extent in low and high frequency evoked twitch forces, suggesting that a basketball simulation protocol affects both excitation and contraction mechanisms to a similar degree.

217

218 15-m sprint times increased by ~2% in PLA-T following the basketball simulation protocol, compared to Afman et al (1) who reported a ~5% increase during the placebo trial of the 219 modified LIST. This lies within the 2-10% decrease in sprint times of short distances (≤ 20 m) 220 typically reported for team sport activities (3, 4, 18, 24). These reductions in running 221 performance are greater than losses in 'pure' strength measurements such as MVICs mentioned 222 earlier (~10-20%). This could be explained by a possible change in sprint mechanics in a 223 fatigued state, affecting speed production as a consequence (33). For instance, in the present 224 study, participants were tightly secured on the dynamometer to avoid any extra bodily 225 226 movements other than the knee extensors, so that MVIC forces could not be affected by a 227 change in technique. Interestingly however, both evoked twitches were significantly greater in ALK-T, demonstrating the protective effect increased extracellular buffering agents have on 228 229 both potassium and calcium ion-related contractile properties of the muscle fibres of the knee extensors. This protection of the muscle force-generating capacity is further illustrated in the 230 231 present study by the attenuation in the continuous development of 232 neuromuscular fatigue (MVIC torque) during the protocol under the NaHCO₃

supplementation.

235 Several studies have to date reported the effect of NaHCO₃ on neuromuscular fatigue. This was following submaximal isometric calf muscles contractions (36), a 2-min voluntary knee 236 237 extension (35), tetanic stimulation (39), and high-intensity repeated sprint cycling (38). In 238 agreement with our findings, all found no condition effect on MVIC forces from pre- to 239 postexercise. In contrast with the present results however, the force decline was similar in both 240 alkalosis and placebo trials (35). The differences in the fatiguing protocols and measurement 241 methods might explain these discrepancies. Our basketball simulation protocol engaged a greater muscle mass and was of longer duration so that a time \times condition interaction effect 242 243 was more likely to occur. Stimulations were applied to the posterior tibial nerve to evoke force 244 in the calf muscles in Siegler et al (36), and as suggested by the authors themselves, the 245 relatively low task demand coupled with the small muscle group might have contributed to the 246 lack of pH effect.

247

The 15-m sprint times were on average 0.2% faster, and MVIC torque 3.3% greater in ALKT 248 (5 measures, n = 10). Whilst 7 out of 10 participants recorded lesser decreases in 100 and 10 249 Hz twitch amplitudes in ALK-T compared to PLA-T, the two-way ANOVA did not depict any 250 interaction effect. A meta-analysis reported for an ergogenic effect of only 1.7% on some 251 performance indicators such as mean power during repeated sprint exercise (8). Several studies 252 253 also reported a lack of condition effect on 15-m sprint times following ingestion of a buffering 254 agent compared to a placebo (1, 34). Whilst the present study focussed on mechanisms affecting 255 the contractile apparatus, it should be noted that alterations within the CNS may also be responsible for declines in voluntary force. Team sport activity has been shown to induce 256 substantial decreases in voluntary activation of quadriceps muscles(14, 43). NaHCO₃ may also 257 258 attenuate afferent feedback associated with metabolite accumulation (37).

259 Therefore, the ergogenic effects demonstrated in the present study (i.e. attenuated MVIC force 260 reduction) might not be purely due to protection of contractile mechanisms. Furthermore, 261 NaHCO₃ supplementation in the present study was limited by the absence of blood gas 262 measurements; however there is evidence that a similar supplementation protocol to the one used in this study raises $[HCO_3^-]$ levels by ~5mmol.L⁻¹ and sustains elevated blood pH and 263 264 HCO₃⁻ during a prolonged intermittent sprint protocol _ENREF_17(5). Factors such as time to 265 peak [HCO₃⁻] and [pH] also show high degrees of inter and intra-individual variability (15, 41). 266 Therefore, it is possible that the ergogenic effect seen in the present study may not have been maximal as the ingestion times were standardised to 90 and 60 minutes. 267

In conclusion, the present study shows a two-phase response in the development of fatigue over 268 269 time during a simulated basketball match, with an initial early development of neuromuscular and peripheral fatigue after just two quarters of simulated match. Beyond which no further 270 271 deleterious effect on the neuromuscular function can be seen. This occurred alongside a 272 slowing down of 15-m sprint times while layup scores remained unchanged. The second major 273 finding is that ingestion of sodium bicarbonate 90 and 60 minutes priorexercise attenuates the 274 above-mentioned development of neuromuscular fatigue. Maximal force production can be preserved until the 3rd guarter of the match. The supplementation preserved both potassium and 275 calcium ion-related contractile properties of the knee extensors so that a greater muscular force 276 generating capacity was possible in the alkaline condition when twitches were evoked using 277 paired stimulations of the femoral nerve. This could be the reason maximal force production 278 279 was preserved during the protocol.

280

281 The present findings should be interpreted with caution due to a small sample size weakening 282 overall statistical power. For example, there was a condition effect for MVIC torque alongside a condition *x* time interaction effect, but with no post-hoc difference depicted. This is not uncommon in the literature surrounding NaHCO₃ supplementation (36, 38). Another limitation in this study refers to the lack of electromyography (EMG) measurements. The intensity for electrical simulation was therefore based on a plateau of twitch force with increasing current, rather than based on a plateau identified for the compound muscle action potential (M wave). As a result, there is no absolute confidence that all motor units were

innervated by the electrical stimulation. However, the mean intensity in the present study (170 mA) is comparable to that of studies using plateaus in twitch and M-wave are des for the determination of stimulation threshold in the knee extensors in similar polation. (80-170 mA,(12); ~190 mA, (14)).

PRACTICAL APPLICATIONS

A simulated basketball match protocol causes a significant mount of overall and peripheral fatigue from the 1^{st} and 2^{nd} quarter, respective ' as quarter by neuromuscular assessments. Sprint times are also slower throughout to mulation basket of match. The employment of intelligent substitution timings and the set of the negate this effect.

Supplementing two dosages of $g.kg^{-1}N$ CO₃ 90 and 60 minutes prior to a basketball simulated match protocol can significantly delay the rate of development of neuromuscular fatigue by protecting optractile operties of muscle fibres.

Acknowledgen nts

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417 **Figure Legends:**

- 418 Figure 1: Schematic of one quarter of the modified Loughborough Intermittent Sprint Test
- 419 (LIST).

Figure 2: Neuromuscular function assessment by play ing quarter. Data present in mea. SD. A: MVIC force throughout the protocol; B: 100 Hz twitch force; C: 10 Iz tw. force Significant group effect; \$ Significant time effect; *Significant intercoon effect

TABLES

Table 1: Assessment of neuromuscular fatigue and performance indicators at all stages of

both conditions (ALK-T: Alkalosis trial; PLA-T: Placebo trial). # Significant group effect; \$

Significant time effect; * Significant interaction effect. All P<0.05. All data is presented in mean \pm SD

Variable		Condition	Baseline	Quarter 1	Quarter 2	Quarter 3	Quarter 4
MVC	\$,*	ALK-T	255 ± 36	251 ± 40	244 ± 42	240 ± 43	233 ± 42
(N.m)		PLA-T	259 ± 32	247 ± 35	233 ± 41	223 ± 38	220 ± 42
100 Hz	#,\$	ALK-T	72 ± 7	69 ± 7	67 ± 9	65 ± 8	65 ± 6
(N.m)		PLA-T	70 ± 8	67 ± 10	64 ± 8	62 ± 9	61 ± 9
10 Hz	#,\$	ALK-T	71 ± 8	67 ± 8	65 ± 8	63 ± 6	63 ± 6
(N.m)		PLA-T	70 ± 7	66 ± 9	64 ± 8	61 ± 6	59 ± 7
10:100 Hz		ALK-T	98 ± 5	98 ± 3	97 ± 4	98 ± 3	96 ± 2
Ratio (%)		PLA-T	100 ± 5	99 ± 4	100 ± 3	100 ± 8	97 ± 5
15m Sprint	\$	ALK-T		2.53 ± 0.11	2.56 ± 0.12	2.58 ± 0.14	2.58 ± 0.11
(s)		PLA-T		2.54 ± 0.11	2.55 ± 0.13	2.58 ± 0.17	2.60 ± 0.16
Layup (%)		ALK-T		92.7 ± 7.2	82.7 ± 12.5	86.4 ± 13.0	87.3 ± 9.8
	2	PLA-T		88.2 ± 4.4	89.1 ± 5.7	85.4 ± 7.7	86.4 ± 10.7



<-----> ^{15m}

<-----> ^{20m}

