The relationships of lifetime physical activity and diet with salivary cell telomere length in current ultra-endurance exercisers

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- 12 Abstract.
- BACKGROUND: Physical activity and a healthy diet may delay the aging process and ultra-endurance exercise is an extreme 13
- form of physical activity. Telomeres are protective DNA sequences located at the ends of eukaryotic chromosomes which 14 shorten as we age. 15
- **OBJECTIVE:** The aim of this study was to investigate the relationships of lifetime physical activity and diet with salivary 16
- cell telomere length in current ultra-endurance exercisers (n = 49; % female = 37, age range 26–74 years). 17
- METHODS: Physical activity and dietary intake were measured using the Lifetime Physical Activity and Diet Questionnaire 18
- (LPADQ) and salivary cell telomere length was measured using quantitative polymerase chain reaction. 19
- **RESULTS:** In this group of current ultra-endurance exercisers there was no relationship between lifetime physical activity 20
- or diet (according to food category scores) and telomere length. In contrast to the expected age-related decrease in telomere 21 length, there was no relationship between age and telomere length (95% confidence interval [CI]: -38.86, 14.54, p = 0.359) 22
- in this group of current ultra-endurance exercisers. 23
- **CONCLUSIONS:** The relationships of lifetime physical activity and diet with telomere length remain uncertain. It is possible 24
- that lifetime physical activity (including ultra-endurance exercise) and lifetime diet may independently, or in combination, 25
- contribute to a decrease in the rate of age-related telomere shortening in current ultra-endurance exercisers. ultra-endurance 26 exercisers. 27
- Keywords: Aging, extreme exercise, dietary intake, ultra-marathon, ultra-cycling 28

1. Introduction 29

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Ultra-endurance exercise involves prolonged periods of physical activity covering a distance of more than the standard marathon (42.195 km) or with a duration greater than 6 hours [1]. The majority of ultra-runners are masters athletes (>35 years) with reported age of participants ranging from 18-81

years [2–4]. Currently, men make up the majority of participants in ultra-endurance events, although an increasing number of women participants over the past decade has been reported [5, 6]. Ultra-endurance exercisers tend to be well-educated and show a large range of training and running experiences [2, 4]. Worldwide participation in ultra-endurance exercise is increasing, yet there is minimal research on the individuals who engage in this form of physical activity. Specifically, there is minimal information on the physical activity and dietary habits of ultra-endurance exercisers or how this form of extreme exercise influences the aging process.

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Aging is an obligatory component of human life 48 and is thought to be determined by several fac-49 tors, including genetics, the environment and lifestyle 50 choices [7, 8]. Telomeres are non-coding repeating 51 segments of DNA (human sequence - TTAGGG) 52 located at the ends of eukaryotic chromosomes [9]. 53 In most dividing cells telomeres shorten which can 54 lead to apoptosis or replicative senescence and this 55 progressive loss of cells can contribute to chronic 56 inflammation, tissue aging and age-related diseases 57 [10]. Telomere shortening is a natural part of the 58 aging process, but shortening may be accelerated 59 by lifestyle factors such as obesity [11], inactivity, 60 smoking [12, 13], and psychological stress [14, 15]. 61 Physical activity and diet have been found to be asso-62 ciated with components of telomere biology with 63 many studies showing positive correlations of reg-64 ular exercise and healthy eating with telomere length 65 [16–21]. While other indices of biological aging have 66 been reported, such as the epigenetic clock [22], 67 telomere length has been used as a marker of bio-68 logical aging in a multitude of research topics and 69 may provide information on the aging process at the 70 molecular level [23, 24]. 71

Physical activity is commonly assessed using ques-72 tionnaires which collect information on intensity, 73 duration and/or type of exercise [25]. Although evi-74 dence suggests habitual physical activity may help 75 preserve telomere length, it is unclear if this depends 76 on the type, volume or intensity of exercise [26]. To 77 our knowledge, only five studies have investigated 78 telomere length in ultra-endurance exercisers with 79 three showing longer telomeres and two showing 80 no difference in telomere length in ultra-endurance 81 exercisers [20, 27-30]. However, these studies have 82 not assessed physical activity across the lifespan and 83 there are no data that examine the influence of diet 84 on telomere length in ultra-endurance exercisers. 85

Most studies investigating the relationship bet-86 ween diet and telomere length have used a food fre-87 quency questionnaire (FFQ) [31-37]. This tool has 88 been used to investigate diet in relation to telomere 80 length based on individual nutrients, foods and/or 90 beverages [38–42] or to examine associations with 91 dietary patterns [17, 43, 44]. However, food fre-92 quency questionnaires used in telomere research only 93 extend as far back as one year and provide dietary 94 information specific to that time period. Although 95 there are two longitudinal studies that have collected 96 dietary intake at baseline and measured telomere 97 length 10 years later [40, 43], there are no stud-98 ies that include information on lifetime diet from 99

childhood to older age. Information on lifetime diet is important considering the dynamics of telomere length changes and attrition begin at birth and extend across the lifespan [45]. Studies show higher intakes of vegetables and/or fruit and adherence to some dietary patterns, such as the Mediterranean diet, are positively associated with telomere length [17, 36]. Therefore, assessing the lifetime intake of healthy and less healthy foods could be a useful and logical approach to investigating the impact of diet on telomere length.

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Telomere length has most frequently been assessed in blood cells, particularly peripheral blood mononuclear cells (PMBC's) [46-49]. However, more recently, studies have used salivary cells to assess telomere length as they are an appropriate alternative to blood cells and provide a fast, simple to obtain, and non-invasive method for sample collection [50]. Salivary cell telomere length is highly correlated with telomere length from whole blood and leukocytes [51, 52] and, therefore, it has been suggested telomere length of salivary cells can be used as a proxy for telomere length in other tissues [53]. Furthermore, genomic DNA from salivary cells has been found to be of similar quality to that obtained from blood cells [54]. The aim of the present study was to investigate the relationships of lifetime physical activity and diet with salivary cell telomere length in current ultra-endurance exercisers.

2. Materials and methods

2.1. Study design

This study includes a subset of participants from a 131 previous study that reported on the lifetime physical 132 activity [4] and diet [55] of current ultra-endurance 133 exercisers. Ultra-endurance exercisers from the pre-134 vious study were invited via email to take part in 135 the present study. Current male and female ultra-136 endurance exercisers were recruited for this study 137 from April 2016 to September 2016. A control group 138 was not included as it was not possible to obtain a 139 comparison group for which physical activity and 140 diet were suitably controlled across the lifespan as 141 the individual patterns could not have been matched. 142 This study was conducted according to the National 143 Statement and Human Research Ethics Guidelines 144 [56] and approved by the University of the Sun-145 shine Coast Human Research Ethics Committee. 146 Lifetime physical activity and diet were investigated 147 retrospectively using an online survey hosted on
the SurveyMonkey® platform. Participants were
provided with an information sheet, and provided
informed consent by completing and submitting the
online questionnaires and by returning a written,
signed consent form with their saliva samples.

154 2.2. Participants

The details of participant recruitment have been 155 described elsewhere [4]. Briefly, participants were 156 recruited internationally through Facebook, the Ultra 157 Listserve (www.ultra@listserve.dartmouth.edu) and 158 web pages associated with ultra-endurance organi-159 zations and events worldwide, including Australia, 160 New Zealand, the United States of America and 161 Canada. Inclusion criteria were being healthy, ≥ 18 162 years of age, able to complete an online question-163 naire in English, and being a current ultra-endurance 164 exerciser. To qualify as a current ultra-endurance 165 exerciser, participants had to have completed at least 166 one ultra-endurance event within the last five years 167 and engage in, on average, at least five hours of run-168 ning or cycling per week during the past year. The cri-169 terion of completion of at least one ultra-endurance 170 event within the last five years allowed for the 171 inclusion of individuals who identified as a current 172 ultra-endurance exerciser (i.e. in training only) but 173 may not participate in events on a regular basis. For 174 the current study, sampling was restricted to Aus-175 tralian and North American participants who had 176 completed all required components of the initial study 177 (n = 86). Fifty-five participants agreed to take part in 178 the study and 50 (91%) saliva samples (33 men; 17 179 women) were received. 180

181 2.3. Questionnaires

Lifetime physical activity and diet were assessed 182 using the Lifetime Physical Activity and Diet Ques-183 tionnaire (LPADQ), details of which have been 184 described elsewhere [55]. In brief, the questionnaire 185 is a modified and combined version of the Lifetime 186 Physical Activity Questionnaire (LPAQ) and the Life-187 time Diet Questionnaire (LDQ) [57, 58]. Medical 188 health history was collected with a Medical History 189 Questionnaire (MHQ). 190

The LPAQ assesses physical activity over several
life periods ranging from childhood to older age [58].
Participants identified all activities they engaged in on
more than 10 occasions over each life period. They
then estimated the total number of years, months and

hours they engaged in the activity. Similarly, the LDQ assesses dietary intake over five life periods (5–18 years, 19–30 years, 31–45 years, 46–60 years and 61–75 years). Using a four-point scale (i.e. rarely, 2–3 times a month; 2–3 times per week; daily), individuals were asked to recall the general frequency at which they consumed certain foods during each time period. The life periods in the LPAQ were modified from the original version to match the diet history life periods [58] and the 5–18 years.

Stressful life events were measured using the List of Threatening Events (LTE) questionnaire [59]. Participants identified if they experienced any of 12 stressful events during each of six life periods (0–12 years; 13–18 years; 19–30 years, etc.). If a stressful event occurred more than once, they were instructed to select as many times as needed. Participants indicated 'yes', 'no' or 'not applicable' to each experience and a lifetime score was obtained ranging from 0–72 (one point for each 'yes' with a higher score indicating higher lifetime stress). For the current study, participants received a link to the List of Threatening Events (LTE) questionnaire by e-mail.

2.4. Assessment of lifetime physical activity and diet

Assessment of lifetime physical activity has been described in detail elsewhere [4]. Briefly, physical activity was calculated as the total number of hours spent in each activity per life period. This was determined by multiplying the number of years, by the number of months and the number of hours per week spent in each activity. Each activity was multiplied by the intensity according to the metabolic equivalent (MET) as determined by the Compendium of Physical Activities [60]. All activities were summed to obtain a total number of MET-hours of activity for each life period. Finally, a total lifetime MET-hours was calculated by summing the total MET-hours per life period up to the participant's current life period.

The assessment of lifetime diet has also been described elsewhere [55]. Briefly, there are 78 food items within the LDQ and these were grouped into two categories (healthy and less healthy). Fifty foods were included in the healthy food category: 17 vegetables, 13 fruits, 4 dairy or alternatives, 5 whole grains, 9 meat or alternatives, 1 alcohol [i.e. red wine] and 1 fat [i.e. olive oil]. Eleven foods were included in the less healthy food category: 6 sweets or sugar sweetened beverages, 1 takeaway food, 1 snack food, 196

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1 processed meat and 2 saturated fats. As such, foods 246 were grouped into categories according to healthy 247 eating guidelines (healthy) and those considered 248 other/extra/processed foods (less healthy). Seventeen 249 food items were not included as there was not enough 250 evidence to support inclusion of these items in one 251 category over the other. A separate sub-category for 252 vegetables and fruit (30 items) was created. These 253 were investigated separately as several studies have 254 reported a benefit of vegetables and/or fruit on telom-255 ere length [34, 36, 43, 44]. Each participant received 256 three scores per life period. A score per life period 257 was calculated by summing response frequencies (i.e. 258 from 1 rarely/never to 4 daily) for each food within 259 the category. Possible ranges were 50-200 (or 49-196 260 for childhood and adolescence) for the healthy food 261 category, 11-44 for the less healthy food category 262 and 30-120 for vegetables/fruit food category. Child-263 hood and adolescence had one less item in the healthy 264 food category (i.e. excludes red wine) and, therefore, 265 a weighted score was calculated by multiplying the 266 score by 200 and dividing by 196. Finally, a weighted 267 lifetime score was calculated as the sum of the scores 268 for each life period multiplied by the number of years 269 spent in the respective life period, and then dividing 270 by the total number of years spent in all life periods. 271

272 2.5. Saliva samples

Purpose-made collection kits were mailed or hand 273 delivered to participants along with instructions for 274 providing saliva samples. Each kit included two 275 30 ml collection containers with 2 ml of a stabilizing 276 solution (RNAlater® Thermo Fisher Scientific). Par-277 ticipants were instructed to provide 5 ml of saliva per 278 container and, upon receipt, samples were stored at 279 -20°C until DNA extraction. At the time of DNA 280 extraction, saliva samples were aliquoted into approx-281 imately 750 µl volumes and one was used for 282 immediate DNA extraction. The remaining aliquots 283 were stored at -20°C for use in subsequent DNA 284 extractions if the initial isolation was unsuccessful 285 (i.e. poor DNA quality/poor yield). All assays were 286 performed by the same investigator, under identical 287 conditions. 288

289 2.6. Measurement of telomere length

Saliva samples were spun for 5 minutes (× 8000 g)
 to obtain a cell pellet and to remove the supernatant.
 Genomic DNA was extracted from salivary samples using the ISOLATE II kit (Bioline) following

the genomic DNA bench top protocol (www. bioline.com/au/). During the pre-lysis step samples were incubated for up to 18 hours depending on the time required for complete lysis. Following isolation DNA samples were stored at -20° C until further analysis. DNA was assessed for quality and quantity via visualization on a 1% agarose gel and spectrophotometry (NanoDrop, Thermo Scientific).

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Relative (rTL) and absolute telomere length (aTL) were measured using Quantitative polymerase chain reaction (qPCR). Relative TL was measured as previously described by Cawthon [61] and aTL was measured using a protocol adapted from O'Callaghan and Fenech [62]. These methods calculate telomere length based on the ratio of telomere repeat copy number to a single copy reference gene 36B4 which encodes the acidic ribosomal phosphoprotein PO. The reference gene 36B4 is not expected to change expression (i.e. copy number) as a function of any of the independent variables being measured. Therefore, the reference gene provides a control level of gene expression whereby a ratio <1 (or >1) indicates the telomere copy number was higher (or lower) than the reference gene. Standard curves for each gene were created using a 10-fold serial dilution of known quantities of a synthesized 84 mer oligonucleotide telomere (TEL) standard containing 14 TTAGGG repeats and a synthesized 75 mer single copy reference gene (SCG) standard, 36B4.

To assist in the normalisation of DNA sample con-323 centration, and reduce pipetting error, each partic-324 ipant sample DNA was prepared to a consistent 325 concentration of 5 ng/µl. The SensiFAST SYBR No-326 Rox kit (Bioline) was used for qPCR according to 327 the manufacturer's instructions. A master mix was 328 prepared with a final concentration of 1 × SensiFAST 329 SYBR No-ROX mix and 400 nM each of forward 330 and reverse primers. The primers used were those 331 previously described by O'Callaghan and Fenech 332 [62] for both the telomere sequence and the 36B4333 reference gene. The telomere primers used were: 334 telomere forward primer (CGGTTTGTTTGGGTTT 335 GGGTTTGGGTTTGGGTTTGGGTT), and telom-336 ere reverse primer (GGCTTGCCTTACCCTTACC 337 CTTACCCTTACCCTTACCCT). Primers for the sin-338 gle copy reference gene amplicon (36B4) consisted 339 of: 36B4 forward primer (CAGCAAGTGGGAAG-340 GTGTAATCC) and 36B4 reverse primer (CCCA 341 TTCTATCATCAACGGGTACAA). In each sample 342 both 36B4 and TEL genes were analysed in tripli-343 cate with two no template controls (NTC) included 344 in each qPCR run. The Rotor-Gene Q (Qiagen), was 345

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used for qPCR analysis. Each run was carried out 346 in 0.1 ml strip tubes with the 72 sample Rotor-Disc, 347 using a final volume of 10 μ l (5.8 μ l of master mix and 348 4.2 µl of participant DNA) to give a final concentra-349 tion of 20 ng/µl of sample DNA. A 3-step cycle was 350 used with cycling conditions of: 3 min at 95°C, fol-351 lowed by 40 cycles of 95°C for 5 sec, 60°C for 10 sec 352 and 72°C 15 sec (with data acquisition), followed by 353 a dissociation (melt) curve ramping from 72°C to 354 95°C. The same amplification and cycling parame-355 ters were carried out for both TEL and 36B4 genes. 356 Triplicate samples all displayed a standard deviation 357 of the Ct (cycle threshold) values of <1 Ct. Samples 358 were inspected to ensure all participants' values were 359 within the linear range, resulting in two participants 360 being identified who showed unusually low amplifi-361 cations. DNA extraction and qPCR were repeated for 362 these two samples, resulting in one participant being 363 removed from further analysis due to poor amplifica-364 tion. All qPCR assays were performed by the same 365 investigator. 366

Relative TL was determined by the ratio of telom-367 ere repeat copy number (T) to the single copy gene 368 copy number (36B4) as described by Cawthon [61] 369 and presented as the telomere length to single copy 370 gene ratio (T/S ratio). Absolute telomere length was 371 measured by determining the number of TTAGGG 372 hexamer repeats as described by O'Callaghan and 373 Fenech [62] and is reported in bases. Briefly, a ratio 374 of telomere repeat to single copy gene (36B4) was 375 obtained by first dividing the 36B4 reaction value by 376 2 (as there are two copies of this gene present in every 377 cell). The telomere length in kilo base (kb) per reac-378 tion was then calculated by dividing the TEL reaction 379 value obtained in the qPCR by the 36B4 (single gene 380 reference). This was converted to length per telom-381 ere (kb) by dividing by 92 (there are 46 chromosomes 382 with a telomere at each end and, therefore, 92 telom-383 eres in total in the human genome). Telomere length 384 is reported in bases by multiplying by 1000. 385

386 2.7. Covariates

Data for eight pre-identified covariates were collected: age, sex, body mass index, smoking history, paternal age at birth, education, medical health history and stressful life events.

391 2.8. Statistical analysis

³⁹² Descriptive statistics are displayed as mean and ³⁹³ standard deviation. Total lifetime MET hours and paternal age at birth were log transformed, and a reflect and square root transformation was applied to lifetime health eating category scores, to normalize the data. Pearson's correlation was used to assess the relationship between absolute and relative salivary cell telomere length and age. Multiple linear regression was used to investigate the independent contributions of age, total lifetime MET-hours, lifetime food category scores, lifetime stressful events scores, paternal age at birth and sex on telomere length. Data were screened to ensure all assumptions for the use of multiple linear regression were met. Statistical analyses were performed using SPSS version 24.0 with an alpha level of p < 0.05.

3. Results

3.1. Salivary cell DNA quality and quantity

Considering the unique nature of the study design and population, it was deemed important to include details on the quality and quantity of the DNA extracted from saliva samples. The quality of the DNA was very good as assessed by agarose gel electrophoresis and spectrophotometry which showed no evidence of DNA degradation. The average 260/280 ratio was 2.0 (pure DNA will have a 260/280 ratio of 1.7–2.0 [62]). The average quantity of DNA, as assessed by spectrophotometry, was 42.7 ± 23.7 ng/µl (range 5.8–95 ng/µl).

3.2. Participant characteristics

Descriptive statistics were calculated and displayed as mean and standard deviation. Participant characteristics are presented in Table 1. Sixty-five percent (32 of 49) were ultra-runners, 18% (9 of 49) were ultra-cyclists and 16% (8 of 49) were a combination (ultra-runner, ultra-cyclist and/or ultratriathlete). Sixty-three percent of participants were men. Men were older and had a higher BMI than women (50.8 ± 10.1 years for men vs 45.2 ± 6.7 years for women, p = .043 and 23.7 ± 2.6 kg/m² for men vs 21.9 ± 1.7 kg/m² for women, p = 0.008).

3.3. Predictors of telomere length

Results of the multiple linear regression are presented in Tables 2 and 3. None of the measured variables were statistically significant cross-sectional

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Participant characteristics					
Ultra-endurance exercisers $(n = 49)$					
Absolute telomere length (in bases)	1602 ± 572				
Relative telomere length (T/S ratio* [†])	1.41 ± 0.05				
Age (years)	48.7 ± 9.2				
Lifetime total MET hours	133072 ± 106911				
Lifetime food category scores					
Healthy category	117 ± 5.3				
Less healthy category	23.9 ± 2.1				
Vegetables and fruit	63.9 ± 4.5				
Lifetime stressful events scores	9 ± 3.5				
Father's age at birth (years)	28.5 ± 7.4				
Education $[n, (\%)]$					
≤high school	4 (8)				
Some college, but no degree	12 (24)				
Associate or bachelor's degree	18 (37)				
Postgraduate	15 (31)				
Smoke history $[n, (\%)]$					
Current smoker	1 (2)				
Past smoker	14 (29)				
Never smoked	34 (69)				
Ethnicity [<i>n</i> , (%)]					
Caucasian	46 (94)				
Other	3 (6)				
Country [<i>n</i> , (%)]					
Australia	29 (59)				
Canada	12 (25)				
United States	8 (16)				

Table 1

Data presented as mean \pm standard deviation and *n* (%) for categorical variables. *ratio of telomere repeat copy number (T) to the single copy gene copy number (36B4). [†]Log transformed.

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predictors of absolute telomere length (p > 0.05). Sex was a statistically significant cross-sectional predictor of relative telomere length (p < 0.05) with men having longer telomeres than women (1.42 + 0.04 for men vs 1.39 + 0.06 for women, p = 0.04). There was no statistically significant difference in absolute telomere length between men and women (1643 + 517 for men vs 1532 + 666 for women, p = 0.52). Age was not statistically correlated with



Fig. 1. Scatter plot illustrating the raw uncontrolled association between age and absolute (a) and relative (b) salivary cell telomere length in ultra-endurance exercisers (n = 49; age range 26–74). Relative telomere length was log transformed.

relative or absolute telomere length for all participants (unadjusted raw data) (Fig. 1).

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4. Discussion

This is the first study to investigate relationships of lifetime physical activity and diet with telomere length in current ultra-endurance exercisers. Participants for this study were drawn from a larger sample of ultra-endurance exercisers who provided detailed information on lifetime physical activity [4] and diet [55]. The ultra-endurance exercisers engaged in a variety of physical activities across the lifespan with

Multiple regression summary statistics for absolute telomere length $(n = 49)$								
Variable	В	95% [CI]	В	Partial r	t	р		
Age	-12.16	[-38.86, 14.54]	-0.232	-0.170	-0.932	0.359		
Lifetime total MET hours [†]	-0.899	[-667.17, 665.37]	-0.001	-0.001	-0.003	0.998		
Lifetime food category scores								
Healthy category [‡]	56.15	[-334.67, 446.97]	0.106	0.054	0.294	0.771		
Less healthy category	-16.28	[-114.08, 81.51]	-0.073	-0.063	-0.341	0.736		
Vegetables and fruit	-2.50	[-92.83, 87.83]	-0.021	-0.011	-0.057	0.955		
Lifetime stressful events scores	45.68	[-15.31, 106.67]	0.315	0.274	1.532	0.136		
Paternal age at birth [†]	-928.98	[-3443.43, 1585.47]	-0.148	-0.139	-0.756	0.456		
Sex [§]	-301.63	[-801.99, 198.73]	-0.272	-0.223	-1.233	0.228		

 Table 2

 Multiple regression summary statistics for absolute telomere length (n = 49)

Significance set at p < 0.05. [†]Log transformed. [‡]Reflect and square root transformed. [§]Coding for categorical variables is male = 0; female = 1.

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Variable	В	95% [CI]	В	Partial r	Т	р
Age	-0.002	[-0.005, 0.000]	-0.414	-0.319	-1.813	0.08
Lifetime total MET hours [†]	0.020	[-0.042, 0.082]	0.129	0.124	0.675	0.505
Lifetime food category scores						
Healthy category [‡]	0.017	[-0.019, 0.054]	0.322	0.177	0.971	0.340
Less healthy category	-0.004	[-0.013, 0.005]	-0.170	-0.158	-0.861	0.396
Vegetables and fruit	0.002	[-0.006, 0.010]	0.168	0.091	0.495	0.625
Lifetime stressful events scores	0.005	[-0.001, 0.011]	0.345	0.321	1.825	0.078
Paternal age at birth [†]	-0.123	[-0.357, 0.110]	-0.194	-0.197	-1.080	0.289
Sex [§]	-0.066	[-0.112, -0.019]	-0.585	-0.473	-2.888	0.007

 Table 3

 Multiple regression summary statistics for relative telomere length[†] (n = 49)

*Indicates significant difference at p < 0.05. [†]Log transformed. [‡]Reflect and square root transformed. [§]Coding for categorical variables is male = 0; female = 1.

total volumes (MET hours) that were, on average, higher than general populations [4]. In this group of ultra-endurance exercisers, whose exercise volumes varied considerably [4], there was no relationship between total lifetime physical activity and telomere length.

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The lack of a relationship between food scores 463 and telomere length was unexpected considering the 464 evidence regarding the influence of diet, particularly 465 vegetables and fruit, on telomere length [17, 36, 43, 466 44]. Similarly, the lack of associations with the less 467 healthy category was not expected as foods that are 468 highly processed, high in saturated fat and/or sugar, 469 have been shown to be associated with shorter telom-470 eres [33, 35, 42]. It has been suggested that certain 471 foods (e.g. processed meat, saturated fat, high sugar 472 foods), many of which are found in the less healthy 473 category in the current study, are negatively asso-474 ciated with telomere length because they increase 475 oxidative stress and inflammation [63]. Although 476 speculative, it is possible that engagement in habitual 477 physical activity, which may lower oxidative stress 478 [64], could help minimize the negative influence of 479 poor dietary choices on telomere attrition. However, 480 the current study collected information on the fre-481 quency of eating and the total quantity of food or 482 energy consumed is unknown and it is possible this 483 may have affected the findings. Also, as this was 484 the first study to use food categories developed from 485 the LDQ to assess lifetime diet, it was not possible 486 to compare the food category scores of the ultra-487 endurance exercisers in the current study to general 488 populations. 489

A secondary finding from this study was the absence of a relationship between age and telomere
length for both absolute and relative telomere length.
This finding supports previous research with ultraendurance runners where a negative correlation of

telomere length with age was not observed [20, 27] and provides some evidence that participating in regular physical activity across the lifetime, that includes ultra-endurance exercise, may help protect against telomere shortening. The lack of association between age and telomere length is of interest considering participants ranged in age from 26-74 years. As expected, the older participants (> 50 years, n=23) had accumulated significantly more lifetime hours of physical activity (that included ultra-endurance exercise) than the younger ultra-endurance exercisers (<50 years, n=26). Studies have reported a positive relationship between habitual physical activity and telomere length [19, 65], particularly amongst older, active individuals [66-68]. It is possible that the accumulating lifetime physical activity of this group, in particular the older participants, had a benefit on telomere length. However, to our knowledge, there are no studies with older ultra-endurance exercisers to help explain the lack of association between age and telomere length and more research with this population is needed.

Possible mechanisms that may explain the lack of associations of lifetime physical activity and diet with telomere length within this group of current ultra-endurance exercisers are the roles of these two variables in promoting general health and wellbeing. Both physical activity and diet may influence levels of oxidative stress and inflammation which are two key underlying mechanisms linked to telomere attrition [63, 69]. Research shows health behaviours tend to cluster by which active individuals tend to eat healthier diets [70] and, therefore, it may be the cumulative effect of health behaviours that benefit telomere length rather than individual components [32, 71, 72]. As such, it is possible the lifetime habits of this group, including physical activity and diet, contributed to preserving telomere length. This study

also included a measure of lifetime stress which is 533 important to consider given the known association 534 between psychological stress and telomere length 535 [15, 73]. In the current study there was no corre-536 lation between lifetime stress and telomere length 537 which may be due to the possible role of physical 538 activity on minimising the impact of stress on telom-530 ere length. Regular physical activity may reduce the 540 impact of psychological stress as observed in a study 541 in which postmenopausal women with the healthiest 542 behaviours (had the highest score related to sleep-543 ing, eating and exercise habits) did not experience the 544 effect of major life stressors on accelerating telomere 545 attrition [74]. Engaging in regular physical activity 546 may explain the lack of relationship between lifetime 547 psychological stress and telomere length in the group 548 of ultra-endurance exercisers in this study. Further-549 more, it is possible, as reported in previous studies 550 with ultra-endurance exercisers, there is an upregu-551 lation of cellular components involved in telomere 552 maintenance that helps protect against telomere attri-553 tion [28]. Although the lack of age-related decline 554 in telomere length may suggest a protective effect of 555 physical activity and diet on telomere length, factors 556 such as small sample size, unmeasured covariates (i.e. 557 genetic variances) and life period specific cofounders 558 (e.g. health related issues, changes in lifestyle or 559 socioeconomic status), need to be considered. The 560 reason for a significant finding related to sex and rel-561 ative, but not absolute, telomere length is unclear 562 and may be due to a difference in the expression 563 of the control gene between sexes. However, more 564 research that includes both measures of telomere 565 length, within the same population, is necessary. 566

To our knowledge this is the first study to report 567 on the lifetime physical activity and diet of ultra-568 endurance exercisers in relation to telomere length. 569 It is the first study to include a measure of diet when 570 investigating telomere length in ultra-endurance exer-571 cisers. Detailed information on physical activity and 572 diet were collected across several life periods which 573 is important when investigating telomere length as 574 changes typically take place over long periods of time 575 (i.e. at least 1 year) and may be influenced by cumula-576 tive factors over a lifetime [75, 76]. This study is one 577 of two studies that have measured salivary cell telom-578 ere length in ultra-endurance exercisers and provides 579 new information on diet for this population. This is 580 the first study to describe both relative and absolute 581 telomere length within the same population of ultra-582 endurance exercisers, which allows for comparisons 583 using both measures. Furthermore, the measurement 584

of absolute telomere length in this study provides the substantial opportunity for a more direct comparison to other studies that have also measured absolute telomere length.

4.1. Limitations

This study has five known potential limitations. Firstly, this study may have lower than optimal statistical power due to the sample size and this may, in part, explain the lack of associations with telomere length. Secondly, this study collected retrospective physical activity and diet data and relied on participants' ability to recall information on habits from several decades ago. Thirdly, this study measured telomere length at one point in time and did not have repeated measures of telomere length at different time points across the lifespan. Therefore, there was no information on the rate of telomere attrition or telomere length at baseline. Fourthly, this study did not include a control group. However, it was not possible to obtain a comparison group for whom physical activity and diet were suitably controlled across the lifespan. Fifth, it was not possible to consider, and control for, all potential co-variates that could influence telomere length across the lifespan. It is possible there were unmeasured variables (i.e. genetic variances) that may have influenced the findings.

More research that assesses lifetime physical activity and diet in a larger sample of ultra-endurance exercisers would assist in explaining the role of these two variables across the lifespan on telomere length. Early life experiences can have a positive or negative effect on telomere length and future research investigating the diet of ultra-endurance exercisers and telomere length during different life stages is needed. This should include information on the type, intensity, duration and frequency of physical activity of ultra-endurance exercisers which would help explain how, and if, diet and physical activity interact in relationship to telomere length.

5. Conclusions

This study adds to the limited amount of research625investigating telomere length in ultra-endurance exer-626cisers and provides the first information on diet and627telomeres for this population. The primary results628showed no relationship of lifetime physical activ-629ity or diet with telomere length. The secondary630

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finding that showed an absence of telomere short-631 ening with age is of interest and provides some 632 evidence for a potential protective role of physical 633 activity and/or diet across the lifetime on cellular 634 aging. It is possible that engaging in various forms 635 of physical activity throughout life, which includes 636 ultra-endurance exercise, may help delay telomere 637 shortening. It is also possible the lack of age-related 638 telomere shortening is independently due to lifetime 639 diet or due to an interactive effect between physi-640 cal activity and diet. However, this process remains 641 unclear and requires further investigation. As par-642 ticipation in ultra-endurance exercise continues to 643 increase worldwide and the number of older indi-644 viduals participating in this form of physical activity 645 continues to grow, it is important to understand the 646 impact this form of extreme exercise has on cellular 647 aging and how diet may influence this process. 648

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