Energy Expenditure of Female International Standard Soccer Players: A Doubly Labeled Water Investigation

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ABSTRACT

MOREHEN, J. C., C. ROSIMUS, B. P. CAVANAGH, C. HAMBLY, J. R. SPEAKMAN, K. J. ELLIOTT-SALE, M. P. HANNON, and J. P. MORTON. Energy Expenditure of Female International Standard Soccer Players: A Doubly Labeled Water Investigation. Med. Sci. Sports Exerc., Vol. 54, No. 5, pp. 769-779, 2022. Purpose: The purpose of this study is to quantify total daily energy expenditure (TEE) of international adult female soccer players. Methods: Twenty-four professional players were studied during a 12-d period where they participated in an international training camp (also inclusive of two competitive games) representing the English national team. The TEE was assessed via the doubly labeled water method during the full 12 d as well as the initial 4-d period before game one. Energy intake was also assessed (via weighed food analysis) during the initial 4-d period to permit estimation of energy availability (EA). Results: Mean TEE did not differ $(P = 0.31)$ between the 12-d (2693 ± 432 kcal·d⁻¹; range, 2105–3507 kcal·d⁻¹; 54 ± 6 kcal·kg⁻¹ fat-free mass [FFM]) versus the 4-d assessment period (2753 ± 359 kcal·d⁻¹; range, 1942–3280 kcal·d⁻¹; 56 ± 8 kcal·kg⁻¹ FFM). Mean 4-d energy intake was 1923 ± 357 kcal·d⁻¹ (range, 1639–2172 kcal·d⁻¹) and mean activity energy expenditure was 1069 ± 278 kcal·d⁻¹ (range, 155–1549 kcal·d⁻¹). When assessed for estimated EA, 88% of players were categorized with low EA status according to the threshold of <30 kcal·kg−¹ FFM. Mean daily carbohydrate intake equated to 3.3 ± 0.7 g·kg⁻¹ body mass. **Conclusions:** When compared with previously published data from adult male players, we demonstrate that the relative daily energetic requirements of engaging in professional soccer training and match play are comparable between sexes. From a practical perspective, data suggest that practitioners should likely focus education and behavior change strategies on "fuelling" for match play and training to optimize both player health and performance. Key Words: CARBOHYDRATE, ENERGY AVAILABILITY, RED-S, NUTRITION

In a dult male professional soccer players, the physical demands of both match play $(1-3)$ and training $(4-6)$ are well documented. Such data typically demonstrate that the absolute loads completed in training are lower n adult male professional soccer players, the physical demands of both match play $(1-3)$ and training $(4-6)$ are well documented. Such data typically demonstrate that the absoenced in match play, as is the case for total distance ≤ 7 km vs \sim 10–13 km), high-speed running distance (\leq 300 m vs >900 m), sprint distance (<150 m vs >200 m), and average speed (<80 m·min⁻¹ vs ~100-120 m·min⁻¹) (7-9). When assessed during a typical in-season weekly microcycle comprising one or two games, outfield professional players typically expend 3000

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to 4000 kcal·d⁻¹ (40–60 kcal·kg⁻¹ fat-free mass [FFM]), as quantified using the criterion standard doubly labeled water method (9–11). Accordingly, evidence-based guidelines for the recommended energy and macronutrient intake to support both daily training and match play have recently been published (12). In this regard, it is suggested that daily carbohydrate (CHO) intake should equate to 3 to 8 $g \cdot kg^{-1}$ body mass to allow for flexibility between rest days, training days and match days.

In contrast to adult male players, the energetic requirements and external training loads completed by elite female players are not as well understood (13–18). This is of specific interest given recent reports documenting the prevalence of low energy availability (LEA, defined as <30 kcal·kg−¹ FFM per day) in female professional players from the English Women's Super League (13). Indeed, these researchers observed that between 50% and 70% of players were classified with LEA status on both match day and "heavy" training days where daily activity energy expenditure was >700 kcal·d⁻¹, as estimated by Global Positioning Systems (GPS). Analysis of self-reported energy intakes (EI) also demonstrated that these players consumed a consistent daily CHO intake of 3 to 3.5 g ·kg⁻¹ body mass,

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BASIC SCIENCES BASIC SCIENCES thereby failing to adjust daily CHO intake in accordance with alterations to training load or in preparation for match play. Such data build on previous observations that female players apparently "under-fuel" in relation to daily CHO intake (14–17). Given that 80% and 69% of type 1 and II muscle fibers from elite female players are classified as empty or almost empty of muscle glycogen immediately postmatch play (18), such relative CHO intakes are likely suboptimal in relation to promoting physical performance.

The reported prevalence of LEA is of particular concern given the potential for players to develop negative symptoms associated with the Female Athlete Triad (19,20) or Relative Energy Deficiency in Sport models (21). Nonetheless, despite previous assessments of activity energy expenditure and energy availability (EA) in such populations (13,15,16,22), it remains difficult to prescribe evidence based nutritional guidelines owing to the indirect methodologies employed to quantify daily total energy expenditure (TEE) (eg, activity diaries and accelerometery which may underestimate or overestimate nonexercise activity). In this regard, the doubly labeled water (DLW) method is the criterion standard method of assessing total daily energy expenditure in free-living conditions in vivo (23). Importantly, this noninvasive method allows for an assessment of energy expenditure over a 7- to 14-d period (ie, a typical in-season micro-cycle) without interfering in dayto-day activities, such as soccer training or match play (23).

Accordingly, the primary aim of the present study was to therefore assess TEE of female soccer players via the criterion standard DLW method. To this end, we studied 24 English female soccer players during a 12-d period where players participated in an international training camp (also inclusive of two competitive games) representing the English national team. As a secondary measure, we also assessed EI (via weighed food analysis) during the initial 4 d of the assessment period to allow for an estimation of EA. Given that this cohort represents players of the highest standard, it is hoped that these data may provide a platform for which to develop evidence-based nutritional guidelines that optimize the health and performance of female players.

METHODS

Participants. Twenty-four female professional international soccer players volunteered to take part in the study. Cohort participant characteristics (also categorized according to playing position) are presented in Table 1. All players remained injury free for the duration of the study. All experimental procedures and associated risks were explained to players and written informed consent was obtained. The study was conducted according to the Declaration of Helsinki and was approved by the University Ethics Committee of Liverpool John Moores University.

Overview of study design. An overview of the experimental protocol is shown in Figure 1. All players completed a 9-d international training camp in November 2019 comprising four training days, one rest day, two travel days, and two match days. Players completed the training prescribed by the national team's coaching staff and were available for team selection to play in two competitive international matches on days 5 (home game) and 8 (away game) during the study period. Three players did not play in either match and where appropriate, these players' data are not reported (indicated accordingly). Total energy expenditure was assessed during a 12-d (9-d camp followed by 3 d at home) and 4-d assessment period using the DLW method, whereas EI was also assessed during the 4 d before match one. Total energy expenditure was assessed over 12 d (as opposed to 9 d) due to logistical challenges of urine collection on days 9 to 11 of the study. Players completed the second international football match abroad in Croatia on day 8. On day 9, players traveled back to the UK and were then driven from the airport to their homes. This resulted in no opportunity to collect urine samples on this day. It was decided between international staff and domestic club staff that players were to rest at home on days 10 and 11 without any interruptions. On day 12, players arrived back at their respective clubs for duty, allowing a final urine sample to be collected. External loading was quantified from all pitchbased training sessions and games. To compare data across time, days are expressed in proximity to the match, for example, 1 d before the game is referred to as match day (MD) minus one (ie, MD-1) whereas the day after the game is referred to as $MD + 1$, and so on.

Baseline measures. Because of logistical issues associated with player availability, body composition was assessed for 18 players only, occurring 2 to 4 wk before the training camp via whole-body dual-energy X-ray absorptiometry (DXA) (Hologic QDR Series, Discovery A, Bedford, MA), where the effective radiation dose was 0.01 mSv per person.

TABLE 1. Baseline player characteristics of elite English female soccer players competing at international level.

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Position	Goalkeepers	Defenders	Midfielders	Attackers	Sauad
Stature (cm)	174.3 ± 0.5 (n = 3)	$169.7 \pm 2.4 (n = 9)$	$168.2 \pm 9.2 (n = 4)$	$163.0 \pm 3.5 (n = 8)$	168.1 ± 5.9 (n = 24)
Body mass (kg)	67.0 ± 8.7 (n = 3)	62.4 \pm 3.2 (n = 9)	60.4 ± 5.0 (n = 4)	60.1 ± 1.1 (n = 8)	62.1 \pm 4.7 (n = 24)
FFM (kg)	45.5 ± 3.5 (n = 3)	44.1 ± 3.6 (n = 6)	42.8 ± 3.9 (n = 4)	41.6 ± 2.1 (n = 5)	43.2 ± 3.4 (n = 18)
Fat mass (kg)	14.4 ± 5.1 (n = 3)	11.1 ± 1.3 (n = 6)	10.3 ± 3.0 (n = 4)	12.2 ± 1.4 (n = 5)	11.8 ± 2.7 (n = 18)
Percent body fat (%)	22.9 ± 5.2 (n = 3)	19.5 ± 2.6 (n = 6)	18.6 ± 4.6 (n = 4)	20 ± 2.7 (n = 5)	20.6 ± 3.7 (n = 18)
Whole body bone mineral content (g)	$2808 \pm 361 (n = 3)$	2837 ± 158 (n = 6)	$2803 \pm 236 (n = 4)$	$2637 \pm 165 (n = 5)$	2766 ± 213 (n = 18)
Whole body bone mineral density $(q \cdot cm^{-2})$	1.26 ± 0.12 (n = 3)	1.33 ± 0.06 (n = 6)	1.35 ± 0.11 (n = 4)	1.26 ± 0.10 (n = 5)	1.31 ± 0.10 (n = 18)
Pelvis bone mineral density $(q \cdot cm^{-2})$	1.37 ± 0.19 (n = 3)	1.28 ± 0.11 (n = 6)	1.35 ± 0.19 (n = 4)	1.42 ± 0.11 (n = 5)	1.38 ± 0.13 (n = 18)
Whole body Z-score	2.7 ± 1.0 (n = 3)	2.4 ± 0.5 (n = 6)	2.7 ± 1.2 (n = 4)	2.1 ± 0.5 (n = 5)	2.4 ± 0.7 (n = 18)
Predicted RMR (kcal- d^{-1})	1549 ± 56 (<i>n</i> = 3)	1515 ± 71 (n = 6)	$1494 \pm 95 (n = 4)$	$1449 \pm 46 (n = 5)$	1486 ± 66 (<i>n</i> = 18)

Stature, body mass, FFM, fat mass and percent body fat values are presented according to playing position. Stature and body mass $n = 24$. Fat-free mass, fat mass, percent body fat, bone mineral content, bone mineral density, pelvis bone mineral density, Z score derived from DXA $n = 18$. Predicted RMR ($n = 18$). Predicted RMR = 120.81 + (4.88 \times stature[cm]) + 8.24 \times FFM $[kg]$) + (5.71 \times age[years]) (24).

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FIGURE 1—Schematic overview of the 12-d study period including the 9-d national training camp. Total energy expenditure was assessed over 12 and 4 d (as opposed to 9 d) due to logistical challenges of urine collection on days 9 to 11 of the study. Day 6 and days 9 to 12 represented rest days during which no scheduled training took place.

All scans were performed and analyzed by the same trained operator in accordance with best practice procedures (25). Resting metabolic rate (RMR) was estimated for each player using a recent female athlete specific predictive equation (24). This equation (RMR = $120.81 + (4.88 \times \text{stature}[\text{cm}]) +$ $8.24 \times FFM[kg]) + (5.71 \times age[years])$ was selected as it was developed using healthy female athletes of a similar age-range and FFM to those in the present study. On the morning of day 1 of the training camp, all players (ie, $n = 24$) were assessed for body mass and stature. Under standardized conditions (>8 h overnight fast), measurement of stature (SECA, model-217, Hamburg Germany) and body mass (SECA, model-875, Hamburg, Germany) were measured to the nearest 0.1 cm and 0.1 kg, respectively according to the International Society for the Advancement of Kinanthropometry guidelines (26) by an International Society for the Advancement of Kinanthropometry Level-1 practitioner.

Quantification of external training and match load. The decision to wear GPS units during training was left to the players (goalkeepers do not wear these units). As such, 13 outfield players who completed all training sessions and matches wore the same portable global GPS units (Apex; STATSports, Newry, Northern Ireland) for all pitch-based training sessions and both matches. Pitch-based sessions were monitored using the GPS units as previously described in professional soccer players (4,27,28). The GPS unit was placed inside a custommade manufacturer provided vest (Apex; STATSports) that held the unit on the upper back between both scapulae, allowing clear exposure of the GPS antennae to acquire a clear satellite connection. External load variables selected for analysis from the training and match data were duration of activity (min), total distance covered (km) and high-speed running (defined as >5.30 to 6.30 m·s⁻¹, >19.08 to 22.68 km·h⁻¹).

Measurement of energy expenditure using the DLW method. Twenty-four players were available for assessment of TEE. Energy expenditure was determined via the DLW method (the criterion standard method of measuring energy expenditure in free-living conditions) which we have previously used in professional team sport athletes (9,11,29). During the evening of day 0, between the hours of 18:00–20:00, players provided a background urine sample. Players then consumed a single bolus oral dose weighed to four d.p. of deuterium (^{2}H) and oxygen (^{18}O) stable isotopes in the form of water $(^{2}H_{2}^{18}O)$, with a desired enrichment of 10% ¹⁸O and 5% ²H₂ using the calculation:

$$
dose (mL) = 0.65 (body mass, g) \times DIE/IE,
$$

where 0.65 is the approximate proportion of the body comprised of water, DIE is the desired initial enrichment (DIE = 618.923 \times body mass (kg)^{-0.305}) and IE is the initial enrichment (10%) 100,000 parts per million (30) dosed according to body weight 2 to 3 wk before the national camp. To ensure the whole dose was administered, participants were observed consuming each bolus dose and each glass vial was refilled with additional water which players were asked to consume. Time of dosing was recorded. Isotopes were purchased from Sercon (Cheshire, UK).

During the morning of day 1 (07:00–10:00), body mass was assessed (SECA, model-875, Hamburg, Germany), and participants were asked to provide a urine sample, collected in a 50-mL tube. This allowed initial isotope enrichment to be determined following total body water equilibrium (30). Thereafter, body mass was collected during the morning of days 2, 3, 4, 5, 6, and 12 and urine samples (second pass of the day) were collected on days 2, 3, 4, 5, 6, 7, 11, and 12 (in line with logistical constraints), to determine elimination rates of both isotopes via the multipoint method (23).

For the DLW analysis, urine was encapsulated into capillaries, which were then vacuum distilled (31), and water from the resulting distillate was used. This water was analyzed using a liquid water analyzer (Los Gatos Research; (32)). Samples were run alongside three laboratory standards for each isotope and three International standards (Standard Light Artic Precipitate, Standard Mean Ocean Water and Greenland Ice Sheet Precipitation; (30,33)) to account for machine day to day variation and correct delta values to parts per million. Isotope elimination rates were converted to EE using an updated two-pool model equation (34) and a mean calculated food quotient of 0.85 ± 0.2 . The results from the energy expenditure data are expressed as a daily average from the 12-d data collection period and also the initial 4-d collection period. Physical activity level (PAL) was also calculated for each player by dividing TEE by RMR. The PAL data are provided for 18 players only, given that 6 players were not available for DXA assessment (hence predicted RMR was not calculated for these players).

Assessment of energy and macronutrient intake. All 24 players on camp completed assessment of dietary and EI. Dietary intake was assessed for the first 4 d of the study via weighed food inventory. A 4-d assessment period was chosen due to logistical issues with overseas travel for the rest of the study. This method of EI assessment has previously been used alongside DLW with athletes (35). All main meals were consumed (ie, breakfast, lunch, and dinner) in the presence of the research team. Any snacks consumed outside of these meals were reported to the research team via the remote food photography method, as described previously (9,11,36). All players were free to self-select food choices and had received no prior education on nutrition strategies for training days. As such, players were asked to continue with their habitual nutritional practices through the study period. The information gained from this study was then used to produce individualized education and behavior change strategies. Weighed food intake was assessed using an identified weighing station for main meals only, which included four separate calibrated weighing scales (Salter 1160 BKDR, Tonbridge, Kent, UK) placed on top of four separate A3 1-cm cubed template place mats. The members of research team operating the stations during breakfast, lunch, and dinner included three Sport and Exercise Register registered performance nutritionists. Once participants had selected their first item of food, they arrived at the weighing station, placed their plate on the scale and informed the registered nutritionist the weight of the plate. This

number was then populated into a predesigned spreadsheet with a description of the food item underneath their name. For example, the participant would tell the member of staff the weight of their food item, that is, 762 g of white pasta, to inform both the weight and item of food. The participant would then place their second chosen item of food on the plate, for example chicken, and would return to the weighing station to re-weigh their plate, by calling out the weight and food item to the member of staff. Participants would follow the same process of calling out the new total weight and food item to one of the three nutritionists who again would populate the spreadsheet. The spreadsheet was predesigned to subtract the weight of the plate from the initial food item to allow quantification of food item number 1. Subsequently, as each food item was then added to the participant plate, the spreadsheet would automatically subtract the previous food item away from the measured food item so quantification of each food item could be calculated independently. This process was repeated until all participants had completed their total meal choice, at which point a photographic picture was captured of the complete final meal and weight and stored for later analysis. If players had finished eating and still had food left on their plate, they were asked to return to the weighing station to see a member of the research team who would subtract any food items left off the original completed meal total via the spreadsheet. In addition to weighing food, the remote food photographic method was used (11) to understand and retrieve information on what players consumed away from the three main mealtimes. This included EI consumed during "snack windows" provided on camp and EI consumed in hotel rooms. Players were asked to provide a photograph of the food or drink that they consumed and were sent to the research team on a smart phone via WhatsApp messaging service, as described previously (36). Third, to further enhance reliability and ensure that participants missed no food or drink consumption, six random 24-h food recalls were also performed by two members of the research team to cross-check methods one and two. To obtain energy and macronutrient composition, professional dietary analysis software (Nutritics Ltd, Ireland) was used by a Sport and Exercise Nutrition register accredited practitioner with experience working with Nutritics Ltd. All EI is reported in kilocalories (kcal) and kilocalories per kilogram of total body mass (kcal·kg⁻¹). Macronutrient intakes were also analyzed and reported in grams (g) and grams per kilogram of body mass $(g \cdot kg^{-1})$.

Menu construction and the preparation of meals and snacks were undertaken by the national team's professional chef and performance nutrition team and developed in line with the demands of the training camp and consideration of proximity to each game. Throughout the duration of EI assessment, meals were consumed at the base camp hotel for the squad with menus provided on a buffet style basis. Breakfast options available daily included: eggs, beans, toast, porridge, muesli, fruits and yoghurts. Lunch and dinner had different options that included one red meat option, one poultry option, one fish option, three to four CHO options (eg, pasta, rice, potatoes, quinoa), three vegetable options alongside a salad bar and snacks such as yoghurts, nuts, cereal bars and condiments. During training sessions, players were provided with low calorie isotonic sports drinks (Lucozade Lite), water and upon request, isotonic energy gels (Science in Sport, GO Isotonic Gels, UK). Protein drinks (Science in Sport, Whey Protein, UK) were provided after training sessions. All CHO provided during training were optional and consumed ad libitum as opposed to individualized prescription to players.

Estimation of EA. Given that FFM was known for 18 players only (due to completion of DXA assessment), EA was initially estimated for this cohort. However, due to a sample error with the urine sample provided by one player on day 4, this player's 4-d analysis of TEE was not completed, hence EA is estimated for 17 players. The thermic effect of food (TEF) was assumed to be 10% of EI for all individuals (37), subsequently enabling estimations of activity energy expenditure ($AEE = TEE - [RMR + TEF]$) and EA ($EA = EI - [AEE]$) FFM]) (38) during the initial 4 d of the training camp. Energy availability was defined using the following thresholds: optimal $($ >45 kcal·kg⁻¹·FFM⁻¹·d⁻¹), reduced (30–45 kcal·kg⁻¹·FFM⁻¹·d⁻¹) and low $($30 \text{ kcal} \cdot \text{kg}^{-1} \cdot \text{FFM}^{-1} \cdot \text{d}^{-1}$) (20).$

Statistical analysis. All data were initially assessed for normality of distribution using Shapiro–Wilk's test. Differences in training load, match load and EI across days were analyzed using a one-way repeated-measures ANOVA. Where significant main effects were present, Tukey post hoc analysis was conducted to locate specific differences. Comparisons between EI and expenditure were analyzed using a paired t-test. Ninetyfive percent confidence intervals (95% CI) for the differences are also presented. Relationships between TEE and body mass, FFM, stature, RMR and 4-d AEE were assessed using Pearson's correlation. All statistical analyses were completed using SPSS (version 27, SPSS, Chicago, IL) where $P \leq 0.05$ is indicative of statistical significance. Data are presented as mean \pm SD.

RESULTS

Baseline characteristics. Player characteristics including stature, body mass, FFM, fat mass, percent body fat, bone mineral content and bone mineral density are presented in Table 1. Data are presented for the full cohort, as well as mean data from positional groups.

Training and match load. External loading variables are presented for $n = 13$ in accordance with those players who wore GPS monitors across all training sessions and games. Training duration (Fig. 2A) was longer on MD-4 (89 \pm 4 min) compared with MD-1 for match one (61 \pm 2 min; 95% CI, 22–32 min; $P < 0.01$) and MD-1 for match two $(63 \pm 7 \text{ min})$; 95% CI, 17–34 min; $P < 0.01$). Similarly, MD-3 training duration (89 \pm 5 min) was also longer than MD-1 training duration for match one (95% CI, 21–33 min; $P < 0.01$) and match two (95% CI, 18–33 min; $P < 0.01$). In contrast, no difference was apparent for the duration of match one $(64 \pm 33 \text{ min})$ and match two $(73 \pm 31 \text{ min})$ compared with the remaining training days $(P > 0.05)$.

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FIGURE 2—A, Training and match–play duration, (B) total distance, and (C) high speed running distance during an international training camp from female soccer players. White bars represent training days, denoted as days away from match day (MD), that is, MD-5, etc., and gray bars represent match day. No training was completed on days with no data bars. "Significant difference from MD-4, $P < 0.05$. "Significant difference from MD-3, $P < 0.05$. "Significant difference from MD-1 before match 1, $P < 0.05$. "Significant difference from MD one, $P < 0.05$. f Significant difference from MD two, $P < 0.05$. Black circles represent individual players. All data are representative of $n = 13$ in accordance with players who wore GPS monitors.

In accordance with exercise duration, more distance (Fig. 2B) was covered on MD-4 (6020 ± 620 m) compared with MD-1 for match one (2927 \pm 862 km; 95% CI, 2090–4095 km; $P < 0.01$) and MD-1 for match two $(4063 \pm 540 \text{ m}; 95\% \text{ CI}, 1177-2736 \text{ m};$ $P < 0.01$). Similarly, MD-3 distance covered (6340 \pm 537 m) was

greater than MD-1 distance covered for match one (95% CI, 2264–4562 m; $P < 0.01$) and match two (95% CI, 1721– 2833 m; $P < 0.01$). The distance covered on MD-1 for match one was significantly lower than both the distance covered on MD-1 for match two ($P = 0.012$) and the distance covered in match two (7430 ± 3237 m; 95% CI, −7734 to −1272 m; $P = 0.004$). There was no significant difference in distance covered between match day 1 (6243 \pm 340 m) and all other days ($P > 0.05$).

High-speed running distance (Fig. 2C) was significantly greater during match one (361 \pm 183 m) compared with MD-4 (126 \pm 85 m; 95% CI, 73–395 m; P < 0.01), MD-1 for match one (85 \pm 79 m; 95% CI, 102–450 m; P < 0.01) and MD-1 for match two $(77 \pm 41 \text{ m}; 95\% \text{ CI}, 107\text{--}460 \text{ m}; P \le 0.01)$. Highspeed running distance was significantly greater during match two $(337 \pm 197 \text{ m})$ when compared with MD-1 for both match one ($P < 0.01$) and match two ($P = 0.013$), although no significant difference was apparent with other training days or match one ($P > 0.05$). There was no significant difference in highspeed running distance between other training days ($P > 0.05$).

Energy expenditure. Mean TEE for the whole cohort $(n = 24)$ across the full 12-d period was 2693 ± 432 kcal·d⁻¹ (range: 2105–3507 kcal·d⁻¹), 43 ± 6 kcal·kg⁻¹ (range, 33–55 kcal·kg⁻¹) and 54 ± 6 kcal·kg⁻¹ FFM (range: 45–68 kcal·kg⁻¹ FFM). Mean 4-d TEE ($n = 23$) was 2753 ± 359 kcal·d⁻¹ (range, 1942–3280 kcal·d−¹), 44 ± 7 kcal·kg−¹ (range, 29–55 kcal·kg−¹) and 56 ± 8 kcal·kg⁻¹ FFM (range, 37–68 kcal·kg⁻¹ FFM). There was no significant difference between 12-d TEE and 4-d absolute TEE ($P = 0.307$). Mean 4-d AEE ($n = 23$) was 1058 ± 352 kcal·d⁻¹ (range, 155–1549 kcal·d⁻¹) and mean PAL values ($n = 18$) was 1.79 ± 0.24 (range, 1.4–2.2). For illustrative purposes, individual data points (where players are represented within their positional groups) are displayed in Figures 3A–D.

EI and macronutrient intake. Mean EI $(n = 24)$ during the 4-d assessment period was 1923 ± 232 kcal·d⁻¹ (range, 1639–2172 kcal·d⁻¹). Both absolute ($P < 0.01$) and relative $(P < 0.01)$ mean EI (Figs. 4A and B) was significantly different between training days. In absolute terms, players consumed less energy on MD-3 (1639 \pm 285 kcal·d⁻¹) compared with MD-4 (2172 \pm 373 kcal·d⁻¹, 95% CI, -807 to -259 kcal·d⁻¹; $P < 0.01$), MD-2 (1919 \pm 319 kcal·d⁻¹; 95% CI, -554 to -5 kcal·d⁻¹; $P = 0.04$), and MD-1 (1962 ± 452 kcal·d⁻¹; 95% CI, -597 to -48 kcal·d⁻¹; $P = 0.01$). In contrast, there was no difference between the MD-4 and MD-2 ($P = 0.80$) or MD-1 ($P = 0.19$) and between MD-2 and MD-1 ($P = 0.97$). In relative terms, players consumed less energy on MD-3 (26 ± 5 kcal·kg⁻¹·d⁻¹) compared with MD-4 $(34 \pm 6 \text{ kcal} \cdot \text{kg}^{-1} \cdot \text{d}^{-1})$; 95% CI, 34–13 kcal·kg⁻¹·d⁻¹; $P < 0.01$) and MD-1 (31 \pm 8 kcal·kg⁻¹·d⁻¹; 95% CI, -10 to 1 kcal·kg⁻¹·d⁻¹; $P = 0.02$). In contrast, no difference was apparent between MD-3 and MD-2 (30 \pm 6 kcal·kg⁻¹·d⁻¹, $P = 0.07$), MD-4 and MD-2 ($P = 0.11$) or MD-1 ($P = 0.25$) and between MD-2 and MD-1 ($P = 0.97$).

Mean absolute CHO intake (Fig. 4C) was similar ($P = 0.37$) between MD-4 (218 ± 56 g·d⁻¹), MD-3 (203 ± 57 g·d⁻¹), MD-2 (192 ± 45 $g \cdot d^{-1}$), and MD-1 (203 ± 71 $g \cdot d^{-1}$). Similarly, mean relative CHO intake (Fig. 4D) was similar ($P = 0.38$) between MD-4 (3.5 ± 0.9 g·kg⁻¹·d⁻¹), MD-3 (3.2 ± 1.0 g·kg⁻¹·d⁻¹), MD-2 $(3.0 \pm 0.7 \text{ g} \cdot \text{kg}^{-1} \cdot \text{d}^{-1})$, and MD-1 $(3.2 \pm 1.1 \text{ g} \cdot \text{kg}^{-1} \cdot \text{d}^{-1})$.

Mean absolute protein intake was significantly different $(P < 0.01$; Fig. 4E) between training days such that on MD- $4 (123 \pm 21 \text{ g} \cdot \text{d}^{-1})$, MD-3 (120 ± 33 g·d⁻¹), and MD-1 $(135 \pm 24 \text{ g} \cdot \text{d}^{-1})$ more protein was consumed than on MD-2

FIGURE 3—A, Mean 12 daily total energy expenditure ($n = 24$), (B) mean 4-d total energy expenditure ($n = 23$), (C) mean 4-d activity energy expenditure $(n = 23)$, (D) physical activity level $(n = 18)$ within each positional group. *Black circles* represent individual players.

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FIGURE 4—A, Absolute and (B) relative EI, (C) absolute and (D) relative CHO intake, (E) absolute and (F) relative protein intake and (G) absolute and (H) relative fat intake across the initial 4-d assessment period ($n = 24$ for all variables). Black circles represent individual players. a Significant difference from MD-4, b Significant difference from MD-3, c Significant difference from MD-2, d Significance difference from MD-1.

 $(100 \pm 23 \text{ g} \cdot \text{d}^{-1}; 95\% \text{ CI}, 5-41 \text{ g} \cdot \text{d}^{-1}; P < 0.01; 95\% \text{ CI},$ 2–39 g·d⁻¹; $P = 0.02$ and 95% CI, 18–52 g·d⁻¹; $P < 0.01$, respectively). No difference was observed between MD-4, MD-3, and MD-1 ($P > 0.05$). Mean relative protein intake was significantly different ($P < 0.01$; Fig. 4F) between training days such that on MD-4 $(1.9 \pm 0.2 \text{ g/kg}^{-1} \text{d}^{-1})$, MD-3 $(1.9 \pm 0.4 \text{ g} \cdot \text{kg}^{-1} \cdot \text{d}^{-1})$, and MD-1 $(2.1 \pm 0.4 \text{ g} \cdot \text{kg}^{-1} \cdot \text{d}^{-1})$ more protein was consumed than on MD-2 (1.6 ± 0.4 g·kg⁻¹·d⁻¹; 95% CI, 0.0–0.6 g·kg⁻¹·d⁻¹; *P* < 0.01; 95% CI, 0.0–0.5 g·

 $\text{kg}^{-1} \cdot \text{d}^{-1};$ $P = 0.03;$ and 95% CI, 0.3–0.8 g $\text{kg}^{-1} \cdot \text{d}^{-1};$ $P < 0.01$, respectively).

Mean absolute fat intake was significantly different ($P < 0.01$; Fig. 4G) between training days such that on MD-4 (90 ± 21 g·d⁻¹), more fat was consumed than on MD-3 (38 ± 14 g·d⁻¹; 95% CI, 37–66 $g d^{-1}$; $P < 0.01$) and MD-1 (67 \pm 24 $g d^{-1}$; 95% CI, 3– 42 $g d^{-1}$; $P < 0.01$). Similarly, more fat was consumed on MD-2 $(87 \pm 33 \text{ g} \cdot \text{d}^{-1}; 95\% \text{ CI}, 28 - 69 \text{ g} \cdot \text{d}^{-1}; P < 0.01)$ than MD-3 and MD-1 (67 ± 24 g·d⁻¹; 95% CI, 15–43 g·d⁻¹; $P < 0.01$) compared with MD-3. Mean relative fat intake was significantly different $(P < 0.01$; Fig. 4H) between training days such that on MD-4 $(1.4 \pm 0.3 \text{ g} \cdot \text{kg}^{-1} \cdot \text{d}^{-1})$, more fat was consumed compared with MD-3 $(0.6 \pm 0.2 \text{ g} \cdot \text{kg}^{-1} \cdot \text{d}^{-1})$; 95% CI, 0.5–1.0 $\text{g} \cdot \text{kg}^{-1} \cdot \text{d}^{-1}$; $P < 0.01$) and MD-1 (1.0 ± 0.4 g·kg⁻¹·d⁻¹; 95% CI, 0.0- $0.6 \text{ g} \cdot \text{kg}^{-1} \cdot \text{d}^{-1}$; $P < 0.01$). Similarly, more fat was consumed on MD-2 (1.3 \pm 0.5 g⋅kg⁻¹⋅d⁻¹)</sup> when compared with MD-3 (95% CI, 0.4–1.1 $g \text{·kg}^{-1} \text{·d}^{-1}$; $P < 0.01$) and on MD-1 when compared with MD-3 (95% CI, 0.2–0.6 g·kg⁻¹·d⁻¹; $P < 0.01$).

EI versus energy expenditure ($n = 24$) and EA ($n = 17$). In relation to the initial 4-d assessment period, there was a significant difference between EI and TEE $(-825 \pm 419 \text{ kcal} \cdot \text{d}^{-1}; 95\%$ CI, -1006 to -643 kcal·d⁻¹; $P < 0.01$) (see Fig. 5A). However, despite significant differences in EI and TEE, body mass did not change across this period (see Fig. 5B) $(0.01 \pm 1.16 \text{ kg})$; 95% CI, -0.48 to 0.51 kg; $P = 0.95$). Mean daily ($n = 17$) estimated EA was 18 ± 9 kcal·kg⁻¹·FFM⁻¹·d⁻¹ (range: 2– 36 kcal·kg−¹ ·FFM−¹ ·d−¹). Overall, 88% of players assessed for EA represented with <30 kcal·kg⁻¹·FFM⁻¹·d⁻¹ (see Fig. 5C).

Factors affecting TEE and AEE. There was a significant positive relationship between 12-d TEE and body mass $(r^2 = 0.56; P < 0.01)$, FFM $(r^2 = 0.65; P < 0.01)$ and predicted RMR ($r^2 = 0.51$; $P < 0.01$). There was also a significant positive relationship between 4-d TEE and 4-d AEE ($r^2 = 0.97$; $P < 0.01$). There was no significant relationship between TEE and stature $(r^2 = 0.15; P > 0.05)$. Data are presented in Figure 6.

DISCUSSION

In using the DLW method, we provide the first direct assessment of total daily energy expenditure of adult female professional soccer players. Our measurements were obtained from players of the highest standard and were collected over a 12-d period when players were representing their national team. When compared with previously published data from adult male players, we demonstrate that the relative daily energetic requirements of engaging in professional soccer training and match play are comparable between sexes. As such, these data now provide a platform for which to develop evidence

based nutritional guidelines for this population. From a practical perspective, our data suggest that practitioners should likely focus education and behavior change strategies (at least for the present cohort) on "fuelling" for match play and training to optimize both player health and performance.

Previous assessments of daily TEE and AEE in female soccer players have been quantified using a combination of indirect methods such as accelerometers, heart rate monitors, activity logs and prediction equations (19,24,25,39,40). In absolute terms, such studies report that the TEE of female soccer players ranges from ~2400 to 2700 kcal·d⁻¹ (22,41,42). In using the DLW method, we observed comparable mean 4-d (three training days, one rest day) TEE of 2753 \pm 359 kcal·d⁻¹ (range, 1942–3280 kcal·d−¹), whereas mean TEE from the full 12-d assessment period was 2693 ± 423 kcal·d−¹ (range, 2105–3507 kcal·d⁻¹). In absolute terms, our data demonstrate a lower TEE to that previously observed in adult male professional players where mean expenditure was approximately 3500 kcal·d⁻¹ (9–11). Nonetheless, when expressed in relative terms (alongside comparable PAL values of 1.4–2.2), it is therefore apparent that the daily energetic requirements of both men and women engaging in professional soccer training and match play typically equates to 40 to 60 kcal⋅kg⁻¹ FFM.

Notwithstanding the limitations of comparing indirect and direct assessment methods, the present data also suggest that the energy requirements of competing and training at an "international" level may be higher than that associated with the players' respective domestic level competition. For example, when compared with players from the English Women's Super League, assessments of the AEE of the goalkeepers $(924 \pm 133 \text{ kcal} \cdot d^{-1})$, defenders $(964 \pm 436 \text{ kcal} \cdot d^{-1})$, midfielders $(1318 \pm 195 \text{ kcal} \cdot d^{-1})$, and attackers $(1073 \pm 348 \text{ kcal} \cdot d^{-1})$ studied here is greater than the mean AEE (418 kcal·d⁻¹) of those players training within the domestic Women's Super League (13). It is noteworthy, however, that the DLW derived assessment of AEE documented here is inclusive of all activity "outside" of pitch-based training such as strength-based sessions undertaken in the gym, recovery swimming pool sessions, as well as nontraining related activity such as walking to and from the

FIGURE 5—A, Difference between TEE and EI ($n = 23$), (B) changes in body mass ($n = 24$) and (C) mean estimated daily EA ($n = 17$) when assessed across the initial 4-d assessment period. Black circles represent individual players.

FIGURE 6—The relationship between mean 12-d TEE and (A) body mass $(P < 0.01)$, (B) FFM $(P < 0.01)$, (C) stature $(P > 0.05)$, predicted RMR $(P < 0.01)$ and (E) 4-d TEE vs 4-d AEE ($P < 0.01$). Black circles represent individual players.

training center and hotel and walking up and down stairs etc. In contrast, the AEE quantified by Moss et al. (13) is derived from a combination of metabolic equivalents and/or accelerometers worn during training, matches and strength and conditioning sessions only. In addition, the training loads completed by Moss et al. (13) was completed in the final month of the season (May), a time when training loads are typically reduced in comparison to other phases of the season.

The external training and match loads observed here are lower than the respective loads associated with other international and domestic level soccer match play (43–45). For example, total distance and high speed running distance covered by outfield players is lower in our study $(8.8 \pm 1.4 \text{ km and})$ 0.35 ± 0.18 km, respectively) compared with other international $(9.9 \pm 1.8 \text{ km and } 1.5 \pm 0.1 \text{ km}$, respectively) and domestic $(9.7 \pm 1.4 \text{ km and } 1.3 \pm 0.9, \text{ respectively})$ soccer matches (44). Difference between studies are most likely due to variation in methods used to collect match load data, where in previous studies, distance covered and high-speed running was estimated from time motion analysis as opposed to GPS adopted here. In addition, the thresholds used for high-speed running in previous studies (>18 km·h⁻¹) is lower than this study (>19 km·h⁻¹) and makes it difficult to compare between studies. Such challenges in the lack of a definitive approach to identify high-intensity actions and the subsequent ambiguity in this area have recently been documented (39).

In relation to EI, previous studies in female soccer players have reported estimated EI of 2124 \pm 444 kcal·d⁻¹ (13), 2226 ± 368 kcal·d⁻¹ (41) and 2387 ± 177 kcal·d⁻¹ (16). In contrast, we report estimated EI that are approximately 200 to

300 kcal·d⁻¹ lower (mean of 4 d, 1923 ± 357 kcal·d⁻¹), a finding that may be due, in part, to the differing methods employed (eg, self-reported food diaries vs researcher supervised weighed food intakes, the latter which may have influenced player food choices toward underconsumption of foods). In agreement with recent observations from players from the English Women's Super League (13), we also observed minimal CHO periodization with players reporting comparable and consistent daily CHO intakes of 3.0 to 3.5 $g \cdot kg^{-1}$. Notably, only one player consumed the recommended range of 6 to 8 g·kg−¹ on the day before the match (12), thus it is likely that players commenced the first game with sub-optimal muscle glycogen stores (18). In contrast, mean protein intake across all training days $(1.8 \pm 0.4 \text{ g} \cdot \text{kg}^{-1} \cdot \text{d}^{-1})$; range, 1.6–2.1 g $\text{ kg}^{-1} \cdot \text{d}^{-1}$) was aligned to supporting training adaptations (46) and in accordance with recommendations for professional soccer players (12). When taken together, it, therefore, appears that female soccer players may not consume (or periodize) sufficient CHO intake to meet the demands of training and competition, a factor that could lead to chronically LEA and symptoms associated with the female athlete triad (21) or Relative Energy Deficiency in Sport models (19). Unfortunately, we are limited in that we do not currently provide any data assessing the impacts of the EI reported here on health and performance outcomes. Nonetheless, from a practical perspective, our data suggest that practitioners should likely target education and behavior change strategies on "fuelling" for match play and training to optimize both player health and performance. Based on our assessment of TEE, it is suggested that relative intakes of CHO, fat and protein corresponding to four to eight (to account for rest days, training days, match day minus 1, match day, etc), 1.5 to 2 and 1.6–2 $g \text{ kg}^{-1} \text{ d}^{-1}$ body mass would provide a reasonable starting point for which to meet the daily energy requirements of female soccer players of professional standard.

Although we readily acknowledge the difficulties in assessing EA (40) as well as the limitation of our 4-d assessment period via weighed food inventory (ie, players may alter food intake because of researcher presence), it is noteworthy that the estimated prevalence of LEA observed here (ie, 88%, 15 of 17, players presented with LEA <30 kcal·kg⁻¹ FFM) is greater than previous reports where 70%, 24%, and 65% of players presented with LEA in English (13), American (14), and Polish national leagues (42), respectively. The lower absolute EI reported here coupled with the potentially increased physical demands associated with com-

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BASIC SCIENCES BASIC SCIENCES peting at international level (when compared with domestic level competition) may be a contributing factor. Although we also acknowledge the limitations (35,36) associated with dietary assessment and potential underreporting (as evidenced by the lack of statistical change in body mass), further work is required to ascertain whether players' chosen dietary choices were an unconscious or conscious decision that is based on beliefs surrounding optimal nutritional practices. We also acknowledge that the classification of LEA status as <30 kcal·kg−¹ FFM is based on laboratory studies that typically adopt short-term periods of "consistent" daily EI, EE and therefore EA. For example, studies that established EA concepts did so over short (4–7 d) periods, where careful but artificial control of diet and exercise, were prescribed (20). The application of such a threshold to real world situations is likely limited by the fact that daily energy expenditure fluctuates day-to-day in accordance with alterations to eating schedules, training load, and competitive demands. Accordingly, the prevalence of LEA status in the present study (and associated longterm physiological implications) may be over-estimated. Further studies are required to evaluate the prevalence of LEA using longer assessment timeframes. Furthermore, assessment of within-day and between-day EA combined with screening tools (21,47,48) and clinical markers would help gain greater accuracy with current assessments of EA in female athletes in the applied field.

CONCLUSIONS

In summary, we provide the first report to directly assess total daily energy expenditure in a cohort of adult female professional soccer players of international standard. Our data suggest that the relative daily energetic requirements of engaging in professional soccer training and match play are comparable in men and women. From a practical perspective, our data suggest that individualized education and behavior change strategies should focus on "fuelling" (ie, increasing daily CHO intake) for match play and training to optimize health and performance.

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