The effects of intensified training on resting metabolic rate (RMR), body composition and performance in trained cyclists

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Abstract

Background

Recent research has demonstrated decreases in resting metabolic rate (RMR), body composition and performance following a period of intensified training in elite athletes, however the underlying mechanisms of change remain unclear. Therefore, the aim of the present study was to investigate how an intensified training period, designed to elicit overreaching, affects RMR, body composition, and performance in trained endurance athletes, and to elucidate underlying mechanisms.

Method

Thirteen (n = 13) trained male cyclists completed a six-week training program consisting of a “Baseline” week (100% of regular training load), a “Build” week (~120% of Baseline load), two “Loading” weeks (~140, 150% of Baseline load, respectively) and two “Recovery” weeks (~80% of Baseline load). Training comprised of a combination of laboratory based interval sessions and on-road cycling. RMR, body composition, energy intake, appetite, heart rate variability (HRV), cycling performance, biochemical markers and mood responses were assessed at multiple time points throughout the six-week period. Data were analysed using a linear mixed modeling approach.

Results

The intensified training period elicited significant decreases in RMR ($F_{(5,123.36)} = 12.0947$, $p = <0.001$), body mass ($F_{(2,19.242)} = 4.3362$, $p = 0.03$), fat mass ($F_{(2,20.35)} = 56.2494$, $p = <0.001$) and HRV ($F_{(2,22.608)} = 6.5212$, $p = 0.005$); all of which improved following a period of recovery. A state of overreaching was induced, as identified by a reduction in anaerobic
**Conclusion**

Intensified training periods elicit greater energy demands in trained cyclists, which, if not sufficiently compensated with increased dietary intake, appears to provoke a cascade of metabolic, hormonal and neural responses in an attempt to restore homeostasis and conserve energy. The proactive monitoring of energy intake, power output, mood state, body mass and HRV during intensified training periods may alleviate fatigue and attenuate the observed decrease in RMR, providing more optimal conditions for a positive training adaptation.

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**Introduction**

Periods of intensified training are deliberately programmed to foster physiological and psychological adaptations to potentially improve physical performance. It is critical, however, to ensure that a balance between training-induced fatigue and sufficient recovery exists, in order to prevent excessive load on the athlete, and minimize the risk of maladaptation to training, illness or injury. Training-related distress can be viewed along a continuum from acute fatigue to overtraining. Short-term periods of intensified training may result in performance decrements associated with acute fatigue, which, upon appropriate recovery, can elicit an adaptive response to improve performance. This state is classically termed ‘overreaching’ or ‘functional overreaching (FOR)’, and is often employed during training camp-situations, with symptoms resolved within several days to weeks. It is important to distinguish between acute fatigue and FOR, however, since the super-compensation effect is reported to be smaller in FOR than acute fatigue [1], and FOR can elicit a greater risk for training maladaptation [2].

Progression of symptoms, and continued imbalance between training and recovery may lead to a more extreme state of severe overreaching, or ‘non-functional overreaching (NFOR)’. NFOR is typically characterized by the inability to sustain effort through intense exercise, diminished performance with maintenance or progression of the training load, and excessive fatigue both at rest and during exercise. Athletes may also present with mood disturbances, psychosocial stress, nutritional and sleep disturbances, and illness, with recovery from NFOR taking several weeks to months [3–5]. Whilst the progression from NFOR to overtraining is considered the most debilitating, the distinction between the two states is complex, since the “clinical features [of overtraining] are non-specific, anecdotal and numerous [5]”, and vary from one individual to another. Consequent long-term performance decrements from overtraining may require several months to years for recovery [3, 4, 6, 7], and should be prevented, wherever possible.

Athletic responses to intensified training periods have been studied extensively [2, 5, 8–11], but there remains no single diagnostic marker to distinguish between acute fatigue, overreaching and overtraining. Much of the applied literature has largely centered on declines in psychological and perceptual measures [1], as well as external measures such as power output, to aid in assessing the severity of an athletes’ condition. The continuum toward overtraining has also been proposed to involve disturbances at the hypothalamic-pituitary level, which may manifest in a reduced hormonal response to exercise [12–14]. In particular, previous studies suggest a disturbance in mood state, impaired race times and decreased power output may occur in athletes suffering from overreaching or overtraining [14–19].
Previous research from the present group suggests that changes in resting metabolic rate (RMR), body composition and energy intake may also be plausible indicators of training distress [20]. RMR is the minimum energy the body requires to perform its basic functions, and is principally dependent on lean mass [21]. In an applied setting, RMR can be used as an indicator of energy availability (EA); defined as the energy remaining for metabolic processes once the energy cost of exercise has been subtracted from dietary intake [22]. Sufficient energy is critical for training consistency, particularly during intensified periods, since prolonged energy restriction can lead to impaired physiological function and increased risk of fatigue, illness and injury, as well as maladaptation to the prescribed training [23]. Significant reductions in RMR, body mass and fat mass have been observed in elite rowers completing four weeks of intensified training at sea level [20], however increases and decreases in RMR have also been observed during altitude training camps in elite and highly-trained athletes, contingent on training volume and dietary practices [24, 25]. Energy homeostasis is centrally regulated, and RMR is closely linked to appetite and energy intake [26, 27]. Therefore, when energy intake is insufficient to support an intensified training load, athletes are more likely to suffer suboptimal EA and a lower RMR. Under such conditions, time trial performance has been demonstrated to decrease in an elite rowing cohort where a state of substantial fatigue and possible overreaching may have occurred [20]. It is plausible that a relationship exists between RMR, energy intake, EA and training load tolerance in endurance athletes, but further data is required to support this premise and to determine the underlying mechanisms involved. Further examination of this relationship is currently being undertaken by a subgroup of our authors.

The aim of the present study was to investigate how an intensified training period, designed to elicit overreaching, affects RMR, body composition and performance in trained endurance athletes, and to elucidate underlying mechanisms. We hypothesised that intensified training would elicit an increased energy demand, leading to reductions in RMR, body composition and performance.

**Method**

**Study design**

Thirteen trained male cyclists completed a six-week training program designed to achieve an overreached state followed by a recovery period. The study was approved by both the Australian Institute of Sport Human Ethics Committee and University of Canberra Human Research Ethics Committee. All participants provided written informed consent prior to involvement. Training was individualized based on each participant’s training history. Training consisted of a combination of monitored, laboratory-based high-intensity interval sessions, and on-road cycling. RMR, body composition, energy intake, appetite, cycling performance, heart rate variability (HRV), biochemical markers and mood responses were assessed at multiple time points throughout the six-week period (Fig 1).

**Participants**

Fourteen male cyclists were recruited from local cycling and triathlon clubs in Canberra, Australia between December 2015 and March 2016 for participation in the six-week program. One participant was unable to continue the training commitments after week 2. Characteristics of the 13 participants who completed the study were (mean ± SD, range): age 35 ± 8 years, 20–47 years; height 185 ± 7 cm, 175–195 cm; body mass 80.5 ± 7.3 kg, 66.0–94.5 kg; maximal oxygen uptake (VO$_{2max}$, absolute) 61.1 ± 6.2 ml.min$^{-1}$kg$^{-1}$, 52.9–73.0 ml.min$^{-1}$kg$^{-1}$; maximal aerobic power (MAP, absolute) 378 ± 28 W, 333–425 W; VO$_{2max}$ (absolute) 4.9 ± 0.2 L.min$^{-1}$, 4.7–5.3 L.min$^{-1}$; MAP (relative) 4.8 ± 0.6 W.kg$^{-1}$, 3.7–5.5 W.kg$^{-1}$. Participants had a consistent cycling...
training history (> 5 sessions wk⁻¹, > 10 h wk⁻¹, > 200 km wk⁻¹, > 4 years) and regularly competed in A and B-grade cycling races. Based on previous literature [28], the subjects were classified as Performance Level 3.

To determine statistically significant changes in RMR, a sample of n = 8 athletes would be required, based on a smallest worthwhile change in RMR of 8% [29], a within-subject SD of 4.3% [30], and Type I and Type II errors of 5% and 20% respectively. Due to the highly applied and demanding nature of the study, it was not possible to pair match an independent control group. We acknowledge this as a limitation to the study.
Training load

The study period was six weeks in total, consisting of a “Baseline” week (100% of regular training load; monitored for the four weeks immediately prior to the study beginning), a “Build” week (~120% of Baseline load), two “Loading” weeks (~140 and 150% of Baseline load, respectively) and finally two “Recovery” weeks (~80% of Baseline load, see Fig 1. Weekly training was prescribed individually through online software (Training Peaks, Boulder, CO), based on Training Stress Score (TSS). TSS is a training load index similar to the heart-rate based TRIMP method; taking into account the duration and intensity of the activity using power output whereby 100 TSS points is equivalent to one hour of exercise at an individual’s functional threshold power [FTP, the power output at which 4 mmol.L⁻¹ blood lactate (BLa) concentration was reached via the power-versus-lactate curve, or lactate threshold 2 [8, 31, 32]]. Participant’s baseline TSS was calculated to reflect the average of their four weeks training prior to the study beginning. All sessions were monitored and adjusted where required to reach the target TSS each week.

Preliminary testing

In the two weeks prior to the study beginning, participants completed an incremental cycling test to exhaustion using an electromagnetically braked cycle ergometer (Lode Excalibur Sport, Groningen, Netherlands) to assess \( \dot{V}\text{O}_{2}\text{max} \) and MAP, as has been described previously [33–35]. Individual training zones and FTP were subsequently calculated based on power output, heart rate (HR) and BLa values obtained for each incremental stage using in-house software [Automatic Data Analysis for Progressive Tests (ADAPT) v6.7, Canberra, Australia].

Resting metabolic rate

RMR was assessed on eleven mornings across the six-week period (Fig 1) using the criterion Douglas Bag method of indirect calorimetry, which has been described previously [30]. All athletes were overnight rested and fasted, and abstained from physical activity for at least eight hours prior to all measurements, which were each completed at the same time of day (± 1 h). Typical error (TE) for the Douglas Bag method of RMR measurement in our hands is 286.8 kJ, or 4.3% [90% confidence limits (CL): 3.1–7.2%] within days, and 455.3 kJ or 6.6% (90% CL: 4.8–11.1%) between days.

Body composition

Body composition was assessed immediately following three of the RMR measurements (Baseline, end of Loading 2, end of Recovery 2; Fig 1) via Dual-Energy X-Ray Densitometry (Lunar iDXA; GE Healthcare Asia-Pacific). Each DXA scan provided an assessment of fat mass, lean mass and bone mineral content (BMC). Fat-free mass (FFM) was calculated as lean mass plus BMC. Radiation safety approval was provided by the Radiation Safety Committee at the John James Hospital, Canberra.

Energy intake

Dietary intake was recorded either by paper diary record or iPhone application (Easy Diet Diary, Xyris Software Pty Ltd, Australia) for the three days immediately prior to each RMR measurement (Fig 1), and analysed for total energy intake and macronutrient consumption by an accredited practising dietitian using nutrient analysis software (FoodWorks Professional v7.0.3016, Xyris Software Pty Ltd, Australia).
Appetite
Subjective feelings of appetite were assessed prior to breakfast following each RMR measurement via 1–10 Likert visual analogue scale (VAS, Fig 1), adapted from [36] (S1 Fig).

Heart rate variability
HRV was assessed during the 25-minute rest period of each RMR measurement, for eleven measurements in total (Fig 1). Upon arrival to the laboratory, participants were fitted with a HR strap (Firstbeat Technologies Ltd, Jyväskylä, Finland). Upon resting supine for five minutes, a ten-minute recording was taken, which was divided into five minutes of rest followed by a five-minute measurement of inter-beat intervals. The inter-beat intervals were analysed using open source analysis software [Kubios HRV Software version 2.0; Biosignal Analysis and Medical Imaging Group, Department of Physics, University of Kuopio, Finland [37]] for time-domain analysis of the mean square root differences of the standard deviation (RMSSD) and its log (LnRMSSD).

Monitored laboratory sessions and cycling performance
Following an initial familiarization on Day 1, 12 monitored laboratory sessions were performed across the six-week period (Fig 1), inclusive of a standardised warm-up, assessment of cycling performance, and a high-intensity interval training (HIIT) session (option 1, 2 or 3) with varied work-rest ratios (Table 1). Participants were blinded to external feedback cues, and instructed to complete all efforts with maximal exertion. Peak power output was recorded immediately following the 15 s sprint. The power output data for the 5 s sprints were discarded due to concerns over the precision of the ergometer’s power output measurement and reliability of the participants’ effort. Mean power output, time to completion and Rating of Perceived Exertion (RPE, 6–20 Borg Scale [38]) were recorded immediately following the 4000 m TT, with BLa measured from capillary sample one minute later. HR was blinded, but monitored continuously throughout (Firstbeat Technologies Ltd, Jyväskylä, Finland). All sessions were performed using calibrated cycle ergometers (Wattbike Pro, Wattbike, Nottingham, UK). Each participant was assigned to the same individual bike for the entire study to ensure measurement error was minimised. Laboratory sessions were completed at the same time of day (± 1 h), with a minimum of two days between each session.

On-road cycling
On alternate days to the laboratory sessions (Fig 1), participants completed two on-road rides in their own time, with a minimum of five hours between each: 1) long duration, aerobic-based session and 2) a series of hill repeats at FTP in order to induce fatigue. Training zones were based on VO2max test results, as previously described. Power output data (Stages left arm crank: Colorado, USA; Garmin Vector: Kansas City, USA; SRM Training System: Jülich, Germany) and HR data (Garmin: Kansas City, USA) for each cyclist were uploaded to Training Peaks upon completion. Each individual’s power meter recording was standardised during 4 x on-road trials using a known distance and elevation (2.8 km, 812 m; Black Mountain, Canberra, Australia, Fig 1). For each trial, the total mass of the rider and bike were recorded, followed by the time to complete one repetition of the known course. Predicted power output was then calculated using a validated regression based on speed, mass and time to complete [39]. The difference between the predicted power and the device-recorded power was then compared to ensure consistency in the power meter recordings across time. Power comparison
data was not utilised for any other purpose than assessing for drift in the predicted-actual power relationship.

Biochemical markers (PRE-POST ergometer)

On eight occasions during the monitored laboratory sessions (Fig 1), venous blood samples (1 x 8.5 ml serum separator tube) were obtained via venipuncture from an antecubital fore-arm vein by qualified phlebotomists. Samples were taken before and after a standardised exercise, i.e. at rest (PRE) and immediately following (POST) the standardised warm-up, in an attempt to mitigate the large variability in the assessment of leptin and free thyroid

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**Table 1. Outline of the monitored laboratory sessions and assessment of cycling performance.**

| A) Standardised Warm up | Elapsed Time | Description |
|-------------------------|--------------|-------------|
| Warm Up                 | 00:00.00–06:00.00 | 6 minutes @ 60% MAP |
|                         | 06:00.00–12:00.00 | 6 minutes @ 70% MAP |
|                         | 12:00.00–15:00.00 | 3 minutes @ 80% MAP |
|                         | 15:00.00–16:00.00 | 1 minute @ 90% MAP |
|                         | 16:00.00–18:00.00 | 2 minutes @ 70% MAP |
|                         | 18:00.00–19:00.00 | 1 minute easy |
| Warm up Effort 1        | 19:00.00–19:05.00 | 5 s warm up sprint @ 80% RPE |
|                         | 19:05.00–20:00.00 | 55 s recovery |
| Warm up Effort 2        | 20:00.00–20:05.00 | 5 s warm up sprint @ 90% RPE |
|                         | 20:05.00–23:00.00 | 2 minutes 55 s recovery |

| B) Cycling Performance |
|-------------------------|
| Effort 1                |
| 23:00.00–23:05.00       | 5 s maximal sprint |
| 23:05.00–24:00.00       | 55 s recovery |
| Effort 2                |
| 24:00.00–24:05.00       | 5 s maximal sprint |
| 24:05.00–26:00.00       | 1 minute 55 s recovery |
| Effort 3                |
| 26:00.00–26:15.00       | 15 s maximal sprint (performance test) |
| 26:15.00–32:00.00       | 5 minutes 45 s recovery |
| Effort 4                |
| 00:00.00–06:00.00       | 4000 m maximal TT (performance test) |
|                         | 6 minutes recovery |

| C) HIIT Session         |
|-------------------------|
| Option                  |
| 1                       |
| 4 x (15 s on/45 s off)  | 3 minutes, 45 s Repeat x 3 |
| 12 x (5 s on/15 s off)  | 3 minutes, 15 s |
| 6 x (10 s on/30 s off)  | 3 minutes |
| 2                       |
| 6 x (10 s on/10 s off)  | 3 minutes, 10 s Repeat x 3 |
| 4 x (15 s on/30 s off)  | 3 minutes |
| 10 x (5 s on/15 s off)  | 2 minutes, 25 s |
| 3 x (20 s on/40 s off)  | 3 minutes |
| 3                       |
| 4 x (20 s on/40 s off)  | 3 minutes Repeat x 3 |
| 4 x (15 s on/45 s off)  | 3 minutes, 25 s |
| 6 x (10 s on/10 s off)  | 2 minutes, 10 s |
| 5 x (5 s on/15 s off)   | 3 minutes |

A) Standardised warm-up, B) assessment of cycling performance, and C) one of three high-intensity interval training (HIIT) session options.

https://doi.org/10.1371/journal.pone.0191644.t001
hormone (triiodothyronine, fT3). External analysis was conducted via immunoassay (Cardinal Bioresearch, Queensland, Australia): Leptin was assayed using a DuoSet ELISA kit (R&D Systems Inc, Minneapolis, USA), and fT3 on the Siemens ADVIA Centaur automated instrument (Siemens Healthcare Diagnostics Ltd, NY, USA) as per manufacturer’s recommendations. Raw data were then assessed as the percentage change between PRE and POST, per session.

Mood questionnaires

Two mood questionnaires, the Multicomponent Training Distress Scale (MTDS) [40] and the Recovery Stress Questionnaire for Athletes (RESTQ-52 Sport) [41], were administered at the same time of day (between 0900 and 1100) according to authors’ instructions on fourteen occasions throughout the six-week period to assess training-related mood disturbance (Fig 1).

Data analysis

The present study design involved repeated measures of multiple variables at specific time points, and a number of proposed inter-variable relationships. A multivariate structural equation model (SEM) was initially employed, however the complexity of the study design and irregularity of measurement points meant that the SEM did not achieve convergence. A linear mixed modelling approach was thus utilised, with independent regressions defined based on the previously predicted SEM relationships. These models allowed us to investigate the time evolution of the dependent variables, associations with other variables (covariates), as well as modelling the substantial amount of heterogeneity amongst subjects and varying baseline levels. All analyses were carried out using the lme4 package [39] in R [40]. The technical specifications of the models are: 1) inclusion of a random intercept for participants, 2) Restricted Maximum Likelihood (REML) estimation, and 3) significance testing of the fixed effects using Type II F tests with Kenward-Roger degrees of freedom approximation. The selection of independent variables included in the models was initially based on a visual assessment of descriptive plots assessing the relationship between the dependent and independent variables. Only those variables that presented the strongest relationship with the dependent variable were included as fixed effects in the linear mixed models. This procedure was adopted to avoid issues with multi-collinearity (e.g. including similar variables that highly correlate) and to avoid over-testing, thus minimizing inflated Type I errors. After fitting an initial full model, a backward model selection procedure was carried out to remove non-significant variables, which helped in the interpretation of the models. Each of the models included evolution over time as a fixed effect (i.e. Training Block), regardless of whether there were any visible changes over time in the visual assessment.

Linear mixed model data are available in Supporting Information Tables 1 to 7 (S1–S7 Tables), and presented as the F-statistic and p-value, with significance set at 0.05. 95% bootstrapping confidence intervals (95% CI) are also presented for those effects that reached statistical significance. Raw data are available in Supporting Information Tables 8 to 18 (S8–S18 Tables), and presented as individual values for each time point, and group mean ± SD.

Results

Training load

Group TSS scores (mean ± SD) for each week throughout the training period were: Baseline = 766 ± 249, Build = 921 ± 234, Loading 1 = 1077 ± 351, Loading 2 = 1121 ± 277, Recovery...
1 = 601 ± 186 and Recovery 2 = 560 ± 192; which corresponded to percentage loadings (compared to Baseline) of 120%, 141%, 147%, 79% and 73%, respectively (Fig 2).

**Linear mixed models**

*Resting metabolic rate.* Absolute RMR and relative RMR were significantly related to the training block (p < 0.05; Table 2), with reductions observed from Baseline to Loading 2, before returning toward Baseline levels in Recovery 2.

*Body composition.* Body mass significantly decreased from Baseline to Loading 2 [95% CI = -1.395; -0.162], and remained low thereafter [95% CI = -1.439; -0.123] (S1 Table).

*Energy intake.* Total energy intake, fat and protein were not significantly related to the training block (p > 0.05). However, CHO consumption increased from Baseline to Loading 2 [95% CI = 21.011; 132.436], and returned toward baseline levels by Recovery 2 [95% CI = -97.030; 50.099] (S2 Table).

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**Table 2. Linear mixed model data for the resting metabolic rate (RMR) model.**

| Training Block | Training Stress Score (TSS) | Total energy intake (mJ.day⁻¹) | HRV (LnRMSSD) |
|----------------|----------------------------|-------------------------------|---------------|
| Absolute RMR (kJ.day⁻¹) | F(5, 123.36) = 12.0947, p = < 0.001 *** | F(1, 127.4) = 5.3509, p = 0.022** (-) | F(1, 107.06) = 0.7349, p = 0.39 |
| Relative RMR (kJ.kg.FFM⁻¹) | F(2, 23.93) = 6.824, p = < 0.001 ** | F(1, 28.786) = 5.4759, p = 0.033* (+) | F(1, 105.45) = 0.0035, p = 0.95 |

FFM = fat-free mass; HRV = heart rate variability

Data are presented as the F-statistic and p-value, and a +/- symbol to denote a positive or negative linear association over time, where relevant. Where a significant linear relationship is observed,

* denotes p < 0.05,

** denotes p < 0.01,

*** denotes p < 0.001.
Appetite. Pre-breakfast sensations of ‘how much the participants felt they could eat’ were decreased between Baseline and Loading 2 [95% CI = -1.816; -0.595], and returned towards baseline levels by Recovery 2 [95% CI = -1.013; 0.372] (S3 Table).

Biochemical markers. Leptin and fT3 were not significantly related to the training block, TSS or absolute RMR (p > 0.05; S4 Table).

Heart rate variability. A significant positive association was observed between HRV and fT3 levels [F(1, 22.122) = 4.5974, p = 0.04]. An interaction effect on HRV was also observed between relative RMR and the training block [F(1, 22.608) = 6.5212, p = 0.005], whereby the higher the relative RMR, the higher the HRV [95% CI = -0.171; 4.385] (S5 Table).

Cycling performance. Peak power output for the 15 s sprint decreased in 12 participants by the end of Loading 2, and returned toward baseline levels by Recovery 2 (S6 Table). Mean power output for the 4000 m TT decreased in 9 participants by the end of Loading 2, and returned toward baseline levels by Recovery 2 [95% CI: -2.294; 93.578]. An interaction effect on mean power output was also observed between TSS and RESTQ-52 Total Stress [F(5, 118.51) = 2.4486, p = 0.04], whereby the higher the stress and TSS, the lower the power output [95% CI = -0.097; -0.032]. Peak HR and RPE during the 4000 m TT decreased from Baseline to Loading 2 [HR: 95% CI = -6.555; -0.608; RPE: -1.545; -0.097], and returned toward baseline levels by Recovery 2 [HR: 95% CI = -2.292; 4.583; RPE: -0.613; 1.176] (S6 Table).

Mood questionnaires. Increases in both MTDS Total Mood Disturbance (TMD) and RESTQ-52 Total Stress were significantly associated with the training block. Responses increased from Baseline to Loading 2 [TMD: 95% CI = 1.701; 4.562; RESTQ: 95% CI = 0.171; 0.929], and returned toward baseline levels by Recovery 2 [TMD: 95% CI = -2.091; 1.436; RESTQ: 95% CI = -0.334; 0.519] (S7 Table).

Time course of change
Raw data comparisons for each variable across the study period as a percentage change from Day 1 are presented in Fig 3.

Discussion
Main findings
The present period of intensified training elicited a state of overreaching in trained male cyclists, and significantly decreased both absolute and relative RMR, body mass, fat mass and HRV, with concomitant increases in mood disturbance, and declines in anaerobic performance, aerobic performance and associated peak HR; all of which improved following a period of recovery. It is likely that the increased energetic demands of training, coupled with insufficient energy intake, are contributing factors to these results; supporting recent evidence from elite rowers that significant decreases in RMR, body composition and performance can occur with heavy training loads when energy intake does not keep up with a greater energy output [20, 24]. The present data do not support the notion that RMR might be a useful marker to monitor training adaptation. Instead, we advocate the proactive monitoring of validated markers of training distress, including subjective wellness, energy intake, power output, body mass and HRV to attenuate fatigue and the potential for a decline in RMR; promoting athlete health, wellbeing and training ability.

RMR, energy availability and intensified training
Relative RMR decreased in the present participants from ~122 to 107 kJ.kg.FFM.day⁻¹ (~29 to 26 cal.kg.FFM.day⁻¹) at the end of the intensified training weeks, supporting a likely decrease
Fig 3. Percentage change in measured variables from baseline in relation to training load across the study duration for A) RMR, B) Body mass, C) Total energy intake, D) Appetite, E) Mood disturbance, F) Biochemical markers leptin and fT3, G) Heart rate variability (LnRMS SD), and H) Cycling performance. The left y-axis depicts Δ% in each of the measured variables, with Δ% in training load on the right y-axis, shaded beneath the curve.

https://doi.org/10.1371/journal.pone.0191644.g003
in EA as a result of the training load, and an increased risk of physiological dysfunction. This notion is supported by the negative linear relationship between both absolute and relative RMR and training load, whereby the greater the training load, the lower the RMR. Much of the previous literature on lowered EA responses to exercise has focused on female athletes who have also demonstrated symptoms of the formerly-known 'Female Athlete Triad' including menstrual dysfunction, disordered eating and impaired bone health [22, 42–44]. The novel data in the present study demonstrate that male athletes can also suffer a low relative RMR and potentially a low EA, which is supported by previous data from our group [20, 24]. Our data also agrees with recent work advocating that male athletes may be susceptible to similar adverse health effects associated with energy restriction as females [45], and confirms the recent notion of ‘Relative Energy Deficiency in Sport’ as a condition applicable to all athletes [23]. These data suggest that there is potential benefit in monitoring RMR within the daily training environment. However, we acknowledge that the measurement of RMR requires specialist equipment and so might only be undertaken where a more extensive investigation of an athlete’s training maladaptation is warranted. By understanding an athlete’s ‘normal’ RMR, and their energy demands at rest, practitioners would be better able to recognize the individual threshold at which values begin to deviate in conjunction with training load. Such knowledge, along with the proactive monitoring of energy intake and body mass, may help to ensure that athletes do not suffer energy restriction from a mismatch between energy intake and output during heavy training, and promote a more optimal EA. Importantly, by maintaining a more optimal RMR and EA, athletes are more likely to have sufficient energy for training as well as crucial physiological functions including bone health, growth and repair, cardiovascular, gastrointestinal and haematological function; ultimately promoting athletic performance [23].

Evidence that overreaching occurred

Performance decline. Training distress in the present cohort was demonstrated by small but significant reductions in both aerobic (4000 m TT, -1.1%) and anaerobic (15 s, -21.1%) performance by the end of the loading weeks, coupled with a decrease in peak HR values. These data agree with other studies reporting performance decrements [16–18, 46–48], however a handful of studies have observed either no decline [49], or even improvements in TT performance in highly-trained and elite cyclists following an overload training period [50, 51]. Such discrepancies may relate to the degree of overload imposed and the training status of the participants; with more highly trained participants being more resilient to increased training volume and intensity. It must also be acknowledged that whilst fatigue is more than likely the driving factor for the observed decreases in peak HR values, a simple explanation could be that these lowered values are directly related to the lowered peak power output from the performance trials. Further investigation is required to ascertain the mechanisms for such changes.

Interestingly, as shown in Fig 3H, the decline in 15 s peak power output occurred prior to the decline in mean power output for the 4000 m TT, and was of a greater magnitude. As we did not indirectly measure muscle activation by integrated electromyography activity nor undertake specific measurements examining changes in neuromuscular function we can only speculate if this decline in anaerobic performance was due to i) peripheral fatigue ii) central fatigue, or iii) a combination of both. We can also only speculate whether participants made a conscious decision to increasingly reduce their effort on this task during the intensified training weeks to conserve energy/themselves for the 4000 m TT. That said, the earlier decline in 15 s peak power output data in the current investigation might indicate that predominantly anaerobic efforts are a more sensitive marker of training distress than a short-term endurance effort such as a 4000 m TT. Rietjens and Kuipers [52] have proposed that a decline in reaction
time to a finger pre-cuing test was strongly suggestive of central fatigue preceding peripheral fatigue. In that study, the training load was significantly increased from baseline for two weeks, however no changes in hormonal responses, body composition or physiological assessments (power output, HR, BlA) were observed, which may imply that their participants did not suffer sufficient training-induced distress to stimulate both central and peripheral fatigue. The present data suggests that regular monitoring of power output during both aerobic and anaerobic efforts may aid in the assessment of training-related distress within the daily training environment.

**Mood disturbance.** Present perceptions of stress and recovery were consistent with increased training volume, and participants demonstrated a worsened mood state through the loading weeks. The perceptual responses provide additional confirmation that the training prescription was sufficient to induce a state of overreaching. These findings are not unique, but rather support recent research in elite rowers from the present group [20] and others [1, 16, 17, 19, 46, 49, 53–55]. Interestingly, RPE for the 4000m TT decreased through the loading weeks, which may suggest that, even though participants were instructed to complete a maximal effort, they were unable or less motivated to do so as a result of their fatigue state. RPE is also well-correlated with HR during steady-state and high-intensity cycle training [8], and so the reduction in RPE might be related to the lowered maximal HR values observed. De Koning et al [56] has postulated that RPE in a closed-loop trial is dependent on the magnitude and rate of homeostatic disturbance, as well as the knowledge of duration or distance remaining. It is plausible that participants experienced a greater homeostatic disturbance earlier in the 4000m TT during the loading weeks and subconsciously adjusted their pacing, which led to a subtle reduction in power output, heart rate and RPE. However, it should also be noted that post-exercise RPE scores are also prone to variability as physiological feedback is diminishing as soon as exercise is terminated and so there can be significant measurement error [57].

A number of statistical associations were also observed between mood disturbances, perceived recovery and HRV, providing a potential link between training load, mood responses and autonomic nervous system activity. Being some of the earliest to change, these data further reinforce the importance of subjective assessments (like RPE) as some of the easiest and more reliable markers to monitor athlete wellbeing and training adaptation, particularly within ecological situations such as training camps [1, 46, 58, 59].

**Possible mechanisms for the observed changes in RMR**

**Body composition.** An individual’s FFM is the greatest determinant of RMR, thus a greater amount of FFM results in a higher energy requirement due to a greater proportion of metabolically active tissue [60, 61]. Previous research has largely demonstrated increases in RMR following exercise, possibly related to increases in FFM [62, 63], increased metabolic demand in response to exercise-induced muscle damage [64–68], and excess post-exercise O₂ consumption (EPOC), which may elevate energy expenditure for up to 24 hours following training [69, 70]. In the present study, we suggest that the small but significant changes in FFM between Baseline and the end of the loading period (-1.3%) are likely to have only partially affected RMR to the extent observed (-12.1%). In addition, participants would have demonstrated some muscle damage and EPOC during the intensified training periods but they did not demonstrate an increase in RMR. A possible explanation for these contradictory findings might be due to the timing of training on the day prior to the RMR measurement, however, in our study, training activity was standardised, and so we are confident our results were not affected in this way. We propose that the decreases in both absolute and relative RMR were
due to a compensatory response to the intensified training load or insufficient energy intake, or both.

In addition, participants’ body mass and fat mass decreased by the end of the loading weeks, suggesting an energy imbalance. Taken together with the finding of a reduction in both absolute and relative RMR, these data support earlier studies which have suggested energy conservation under intensified training circumstances [20, 71]. One contrasting study found no change in body mass or fat mass in competitive cyclists undertaking two weeks of intensified training [18], however the study estimated body composition from skinfold measurements, which typically have lower test-retest reliability than the DXA method used in the present study, and so might account for the disparity in the findings. Nonetheless, our findings emphasize the critical nature of maintaining energy intake, independent of feelings of appetite (which might be relatively insensitive), in order to maintain body mass and RMR; each of which are strongly linked [72]. This notion is particularly important for athletes who cannot afford to lose lean mass, risking a decline in performance from a decrease in muscular strength and power capabilities.

**Energy intake and appetite.** Supplementary CHO ingestion throughout a training cycle has been reported to assist in alleviating the symptoms of overreaching [73, 74], and may mitigate the stress hormone response to exercise [75]. If total energy intake is insufficient, however, acute ingestion of CHO immediately before and after a training session may not provide an attenuation of fatigue-induced decreases in maximal power output or immunological disturbance [76]. The present cohort attempted to increase their CHO intake by the end of the loading weeks; however such compensation appears not to have been sufficient to attenuate a reduction in RMR or fatigue. It is plausible that changes in participants’ appetite responses were delayed in relation to the changes in energy output, and so an energy imbalance occurred. However, we acknowledge that individual appetite responses were highly variable, and so these findings must be interpreted with caution.

Leptin is a hormone secreted by the adipose tissue, and is reported to regulate neuroendocrine function, appetite perception and energy homeostasis through a series of complex interactions within the hypothalamus, the mesolimbic dopamine system and hindbrain [77–81]. High leptin levels are associated with increased satiety and energy expenditure, whilst low leptin levels, as seen in the present cohort, are consistent with low levels of body fat and chronic energy restriction [77, 81–85]. In particular, leptin has been suggested as a marker of training stress in male rowers [86], and is widely reported to decrease following heavy training periods [71, 87, 88]. In contrast to previous research, leptin levels in the present study tended to increase through the loading weeks, indicating greater satiety; however the responses were highly variable between individuals and so not statistically significant. Pre-breakfast perceptions of ‘how much the participants felt they could eat’ were lower in the loading weeks, further supporting an increase in satiety or decrease in hunger. Anecdotal reports from athletes within the Australian Institute of Sport cite a loss of appetite with heavy training, but these reflections, and our data, are not consistent with the literature. Another explanation of our findings might relate to dietary intake. In overweight and obese populations, overfeeding is reported to increase circulating levels of leptin [81]. More applicable to the present context, perhaps, is that leptin levels are highly correlated with carbohydrate intake [89], and can be influenced by circulating insulin and pro-inflammatory cytokines such as tumor necrosis factor and interleukin-6 [81], so it is possible that the observed trend of increased carbohydrate intake during intensified training had some effect. Perhaps another confounding factor in the observed leptin response was the timing of the blood sampling, which was undertaken prior to and immediately following a physical activity task, and may have been influenced by participants’ acute energy intake (such as glucose-rich sweets) prior to the blood sampling, as well as their feeding
across the day. Despite this, the present data suggest that, in a practical sense, it is crucial for athletes to maintain sufficient energy intake to support their training load. It is possible that athletes should be instructed to eat in relation to the training undertaken, rather than appetite, to fuel optimal performance and recovery.

**Thyroid hormone.** Free triiodothyronine (fT3) has been proposed as a key regulator of metabolic rate and overall energy expenditure by modulating a number of regulatory pathways in skeletal muscle and other tissues [90–92]. Increases in circulating thyroid hormones are broadly associated with an increase in RMR, with the opposite trend occurring in response to lowered hormone levels [89]. Total T3 tended to decrease in response to chronic energy restriction and high-energy expenditure in a military setting [93]; and in females, T3 is lower in association with an increased severity of exercise-associated menstrual disturbances, reflective of energy conservation [85]. In the present study, the percentage change in fT3 demonstrated varied responses throughout the loading and recovery weeks, which did not result in statistical significance. Nonetheless, the substantial changes illustrated in Fig 3F might indicate an altered thyroid and hypothalamic–pituitary–thyroid (HPT)-axis activity as a result of the intervention, which may have practical implications for energy production and thermogenesis, nutrient metabolism, and the regular functioning of the cardiovascular system [94]. We were unable to measure these axes directly, however, and so this notion remains speculative and requires further investigation.

**Heart rate variability.** The observed reduction in LnRMSSD might be attributed to accumulated fatigue as a result of the training load, and may reflect the decreased ability of the ANS to respond to exercise training, stress and illness [95]. Reductions in LnRMSSD may further indicate parasympathetic hyperactivity (or saturation) and reduced sympathetic tone [96] if accompanied by increases in inter-beat intervals [97], which has been reported in response to periods of intensive training in elite and well-trained endurance athletes [97–100]. We propose that alterations in ANS activity might have influenced metabolic activity, as evidenced by the similar pattern of RMR and HRV responses, and the statistical association between fT3 and HRV. Fig 3 illustrates a decrease in RMR immediately prior to a decrease in HRV, so it is possible that an increase in parasympathetic activity, with ensuing reduction in sympathetic activity, may influence (or be influenced by) changes in RMR. Further research is needed to fully understand this potential association.

**Limitations**

The present investigation was applied in nature, and whilst scientific rigour was paramount, there remain some limitations that must be acknowledged. Firstly, we acknowledge that our findings need to be interpreted with caution given that individuals, when training intensively, can exhibit highly variable responses, and also the statistically significant changes lay close to both the technical error of measurement and normal day-to-day variability. The study design consisted of multiple measurements across a number of time points, which resulted in difficulty in applying a statistical model; the power of which would have been improved with both a greater number of participants, as well as simultaneous measurements. The combination of biological and measurement error further adds complexity, and as such we have focused on the broad trends observed between variables. We also acknowledge the lack of an independent pair-matched control group, however the difficulty in retaining participants for the course of the six weeks meant it was not possible to recruit a separate cohort for comparison. Whilst this means that it is difficult to conclude with certainty that the changes observed are truly due to the training intervention applied, we are confident that by monitoring the participants for four weeks prior to the study beginning, we were able to gauge an accurate representation of their
routine training. We are thus confident that the physiological changes observed during the study period can indeed be attributed to the increased training load. We also acknowledge the possibility that some of the unexpected responses to intensive training may be due to the trained status of our participants, who, given their routine volume of training might have been better able to adapt to the ‘predictable’ stress of the training imposed. As such, a number of different central responses might have been produced which we were not able to predict and subsequently assess. Finally, we recognize that the participants were free-living, trained cyclists, but not elite athletes. As such, they were subject to stressors outside of our control including work and study commitments, family duties, and lifestyle factors which may have added to the imposed training load.

Practical application

The present data suggest that during periods of intensified training, practitioners should employ a series of monitoring tools—early, and often—to avoid detrimental levels of training-related distress and ensure sufficient energy intake to support the greater energetic demands. In the daily training environment, athletes should specifically be encouraged to increase their energy intake in relation to training load, rather than appetite, to support a more optimal EA. The proactive monitoring of subjective wellness, energy intake, power output, body mass and HRV during intensified training may further support athlete health, wellbeing and training ability before a detrimental decline in RMR, and likely EA, becomes apparent. Importantly, a more optimal RMR and EA will, in turn, ensure sufficient energy is available for training, recovery and adaptation, and ultimately, athletic performance.

Conclusion

Athletes often undertake periods of intensified training in order to improve performance following a period of recovery. The present study demonstrates, however that exercising with an increased training load, without sufficient energy intake, can risk significant reductions in both absolute and relative RMR, body mass, HRV and performance, and increased mood disturbance. Such physiological disturbance and maladaptation to training may be problematic in athletes who cannot afford to lose mass, or those undertaking intense training prior to competition. We propose that a cascade of changes in metabolic, neural and hormonal mechanisms results from the body’s attempt to conserve energy and maintain homeostasis when energy demands are increased. The proactive monitoring of subjective wellness, energy intake, power output, body mass and HRV during intensified training periods may alleviate fatigue and attenuate any decreases in RMR, and subsequently provide more optimal conditions for a positive training adaptation.

Supporting information

S1 Fig. Subjective feelings of appetite assessment via 1–10 Likert visual analogue scale. (JPG)

S1 Table. Linear mixed model data for the body composition model. Data are presented as the F-statistic and p-value, and a +/- symbol to denote a positive or negative linear association over time, where relevant. Where a significant linear relationship is observed, * denotes p < 0.05, ** denotes p < 0.01, *** denotes p < 0.001. (DOCX)

S2 Table. Linear mixed model data for the energy intake model. Data are presented as the F-statistic and p-value, and a +/- symbol to denote a positive or negative linear association over
time, where relevant. Where a significant linear relationship is observed, * denotes $p < 0.05$, ** denotes $p < 0.01$, *** denotes $p < 0.001$.

(SDOCX)

S3 Table. Linear mixed model data for the appetite model. Data are presented as the F-statistic and p-value, and a +/- symbol to denote a positive or negative linear association over time, where relevant. Where a significant linear relationship is observed, * denotes $p < 0.05$, ** denotes $p < 0.01$, *** denotes $p < 0.001$.

(SDOCX)

S4 Table. Linear mixed model data for the biochemical markers model. Data are presented as the F-statistic and p-value, and a +/- symbol to denote a positive or negative linear association over time, where relevant. Where a significant linear relationship is observed, * denotes $p < 0.05$, ** denotes $p < 0.01$, *** denotes $p < 0.001$.

(SDOCX)

S5 Table. Linear mixed model data for the heart rate variability model. Data are presented as the F-statistic and p-value, and a +/- symbol to denote a positive or negative linear association over time, where relevant. Where a significant linear relationship is observed, * denotes $p < 0.05$, ** denotes $p < 0.01$, *** denotes $p < 0.001$.

(SDOCX)

S6 Table. Linear mixed model data for the cycling performance model. Data are presented as the F-statistic and p-value, and a +/- symbol to denote a positive or negative linear association over time, where relevant for the Modified Power Profile sprints and 4000m TT. Where a significant linear relationship is observed, * denotes $p < 0.05$, ** denotes $p < 0.01$, *** denotes $p < 0.001$. From the initial full model, variables considered non-significant following a backward model selection procedure and subsequently removed are denoted by #.

(SDOCX)

S7 Table. Linear mixed model data for the mood questionnaire responses model. Data are presented as the F-statistic and p-value, and a +/- symbol to denote a positive or negative linear association over time, where relevant for a) the MTDS and b) RESTQ-52 Sport. Where a significant linear relationship is observed, * denotes $p < 0.05$, ** denotes $p < 0.01$, *** denotes $p < 0.001$.

(SDOCX)

S8 Table. Raw data: Absolute RMR. Data are presented as individual values for each time point, and group mean ± SD.

(SDOCX)

S9 Table. Raw data: Relative RMR. Data are presented as individual values for each time point, and group mean ± SD.

(SDOCX)

S10 Table. Raw data: Minute ventilation $[VE_{STPD}]$. Data are presented as individual values for each time point, and group mean ± SD.

(SDOCX)

S11 Table. Raw data: Body composition. Data are presented as individual values for each time point, and group mean ± SD.

(SDOCX)
S12a-d Tables. **Raw data: Energy intake.** Data are presented as individual values for each time point, and group mean ± SD.

(DOCX)

S13a-d Tables. **Raw data: Appetite.** Data are presented as individual values for each time point, and group mean ± SD.

(DOCX)

S14a-b Tables. **Raw data: Biochemical markers PRE-POST ergometer warm-up.** Data are presented as individual values for each time point, and group mean ± SD.

(DOCX)

S15a-b Tables. **Raw data: Heart rate variability.** Data are presented as individual values for each time point, and group mean ± SD.

(DOCX)

S16a-e Tables. **Raw data: Cycling performance.** Data are presented as individual values for each time point, and group mean ± SD.

(DOCX)

S17 Table. **Raw data: Mood questionnaires—Multicomponent training distress scale.** Data are presented as individual values for each time point, and group mean ± SD.

(DOCX)

S18 Table. **Raw data: Mood questionnaires—RESTQ-52 sport.** Data are presented as individual values for each time point, and group mean ± SD.

(DOCX)

**Acknowledgments**

We would like to sincerely thank the athletes for their participation in the study, and the staff and students from AIS Physiology, AIS Nutrition, and UCRiSE for their assistance with testing sessions. We would also like to thank Professor Romain Meeusen, Professor Peter Hassmen and Dr Nathan Versey for your advice in designing the study, John Cardinal and Victor Vuong for your assistance with biochemical analysis and Jamie Plowman for your technical expertise.

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