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Etiology and Pathophysiology

Why do individuals not lose more weight from an exercise intervention at a defined dose? An energy balance analysis

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Summary

Weight loss resulting from an exercise intervention tends to be lower than predicted. Modest weight loss can arise from an increase in energy intake, physiological reductions in resting energy expenditure, an increase in lean tissue or a decrease in non-exercise activity. Lower than expected, weight loss could also arise from weak and invalidated assumptions within predictive models. To investigate these causes, we systematically reviewed studies that monitored compliance to exercise prescriptions and measured exercise-induced change in body composition. Changed body energy stores were calculated to determine the deficit between total daily energy intake and energy expenditures. This information combined with available measurements was used to critically evaluate explanations for low exercise-induced weight loss. We conclude that the small magnitude of weight loss observed from the majority of evaluated exercise interventions is primarily due to low doses of prescribed exercise energy expenditures compounded by a concomitant increase in caloric intake.

Keywords: Body composition, compensation, energy balance, exercise.

Introduction

In order to lose weight, energy expenditures must exceed energy intake. To achieve this imbalance, one can decrease energy intake, increase energy expenditures or combine a decrease in intake with an increase in expenditures. As a result, most weight loss recommendations advise combining a hypo-caloric diet with an exercise component in order to achieve a significant energy deficit (1). A long-standing consistent observation is that regular exercise by itself is prescribed in small to moderate amounts resulting in modest weight loss or in some cases weight gain (2–4).

The commonly observed low levels of weight loss have led to several questions regarding the effects of regular exercise on body weight and the reasons why the energy deficit appears to be smaller than predicted. Addressing these highly debated questions (5) requires quantitative knowledge of each term in the energy balance equation:

\[ ES = EI - EE \]  

(1)

where ES represents the rate of changed body energy stores, EI is the rate of energy intake, and EE denotes the rate of energy expenditures typically expressed in kcal d⁻¹.

In a 1985 review, Pi-Sunyer and Woo (6) described how within an exercise study, lack of simultaneous measurements of the components of energy expenditure, body composition and EI create challenges for understanding...
physiological responses to exercise. Despite the limitations of existing studies, a number of reviews have combined the exercise literature to develop conclusions on the impact of exercise on human morphology and physiology (5–11). Here, we expand on these attempts by employing a quantitative energy balance approach to:

- ascertain the actual energy imbalance generated in each exercise intervention study retained for evaluation;
- examine the underlying reasons for the discrepancy between the actual energy imbalance and prescribed exercise expenditure;
- Improve existing body composition models that describe changes in fat-free mass (FFM) during exercise-induced weight loss; and
- identify gaps in knowledge on the impact of regular exercise on the components of energy expenditure and the generation of substantial energy deficits.

Our calculations are based on data from carefully selected aerobic exercise studies targeting weight loss. We focus on aerobic exercise-induced weight loss because it is open to quantitative analysis and prediction in an area that has not been fully elucidated and in which uncertainties are limiting further development.

Methods

Search strategy and study selection

We performed a systematic review of the literature from 2005 to present within the PubMed database using the terms (‘exercise’ OR ‘aerobic exercise’ OR ‘endurance exercise’) AND (‘body composition’ OR ‘fat mass’ OR ‘fat free mass’) OR ‘weight loss’ OR ‘compensation’ OR ‘energy balance’ OR ‘resting metabolic rate’. The resulting list was screened for aerobic exercise interventions within a sedentary healthy population. Aerobic exercise was defined as any activity that increased oxygen consumption, was performed at submaximal intensity, and involved large groups of skeletal muscles.

We then screened for studies that either provided total body estimates of fat mass (FM) and FFM at baseline and end of the intervention, directly measured EI and/or EE in confined subjects at the beginning and end of the intervention, obtained simultaneously collected doubly labelled water (DLW), FM and FFM measurements at the beginning and end of the study, or reported total energy expended from the prescribed exercise program. We additionally included relevant exercise interventions directed by the authors (2–4,10–14) and studies that have been previously reviewed (5,8,12).

All selected studies prescribed a dose of exercise without a reduction in diet and established compliance to the prescribed exercise dose. In all of the reviewed studies, aerobic exercise consisted of treadmill walking, treadmill running or training on cycle ergometers with interventions ranging in duration from 3 to 64 weeks (2–4,13–25).

A comprehensive list of studies reviewed, the corresponding baseline characteristics of the study samples, prescribed exercise energy expenditures and methods of body composition measurement are summarized in Table 1.

The data were used to answer three questions: What is the energy deficit induced by exercise? Here we quantify the energy deficit by either calculating the change in body energy stores or the difference between EI and EE from direct measurements obtained from confined subjects. What mechanisms explain the discrepancy between the actual energy deficit and prescribed energy expenditure of the exercise program? Here we explore which terms in the energy balance equation could account for the gap between the actual energy deficit and prescribed energy expenditure of the exercise program. Specifically, these mechanisms are thought to be limited to a compensatory increase in EI, a decrease in resting metabolic rate (RMR) beyond expected at the reduced weight, or a decrease in spontaneous activity during the non-exercise hours of the day. What are the expected changes in body composition resulting from an aerobic exercise intervention and how can they influence the estimated energy deficit caused by the exercise program? Here we first identify the limitations of extrapolating the Forbes model (26–29) relating FFM and FM during changes in EI to the case of exposure to regular exercise. We then modify the model to reflect exercise-induced body composition changes. A schematic diagram mapping the course of our analysis appears in Fig. 1.

Studies that maintained EI at baseline levels

In order to segregate the effects of exercise on energy balance from any possible change in EI, EI needs to be maintained at carefully determined baseline levels. The study of Bouchard et al. (13) is unique in this regard because EI was maintained at baseline requirements and negative energy balance was generated in 14 men solely through exercise for a period of 93 d. Subjects in the Bouchard study were confined and supervised for the entire duration of the intervention and adherence to both the exercise and dietary prescription was rigorously monitored. Additionally, RMR was measured at baseline, 50 d, and end of study (100 d).

The study of Ross et al. (15,16) was not conducted in a confined setting, although diet was monitored daily and adherence to baseline EI was confirmed through a DLW measurement midway through the study. The Ross study generated a large magnitude of energy expended through exercise comparable to reductions in intake prescribed in typical caloric restriction studies. Additionally, the Ross study included both men (n = 16) and women (n = 17).
If prescribed energy converted to energy cost by multiplying each litre of VO2 by 5 kcal. Duration of the trial represented by averaged the amount over the length of the intervention to arrive at a prescribed energy expended through exercise. The total oxygen consumed during exercise was expenditures from exercise were not directly reported, we determined this amount by multiplying the reported exercise intensity by the duration of each session. We then study; STRRIDE, Studies of Targeted Risk Reduction Interventions Through Defined Exercise.

BC, body composition; BMI, body mass index; DREW, Dose-Response to Exercise in postmenopausal Women study; FM, fat mass; INFLAME, Inflammation and Exercise study; STRRIDE, Studies of Targeted Risk Reduction Interventions Through Defined Exercise.

*Intervention was separated into three phases each lasting a period of 19 d; sedentary, mild dose of exercise and moderate dose of exercise.

**Prescribed energy expenditures from exercise converted to energy cost by multiplying each litre of VO2 by 5 kcal. Duration of the trial represented by Δt and BC-labelled column denotes body composition method (BD, body density; DXA, dual energy x-ray absorptiometry; MRI, magnetic resonance imaging; SF, skin fold; Siri-4, Siri-four compartment model; TBK, total body K; TC, three-compartment model based on body weight, body volume, body density; UWW, underwater weighing).
Studies that confined and supervised ad libitum EI and EE
Studies (17–19) were conducted in a confined setting and although EI was not maintained at baseline levels, EI and EE were both directly measured. Both men (n = 3) and women (n = 14) were included in these studies with subjects body mass index (BMI) ranging widely (19.5–34.3 kg m$^{-2}$).

Studies that obtained doubly labelled water -measured EE
The study of Westerterp et al. (4) obtained DLW measurements of total energy expenditures at baseline and end of study in 13 of the 23 subjects.

Studies of free-living subjects
The studies (2–4,14,20–25) were conducted in free-living subjects. Although subjects in most of the studies were asked to maintain baseline EI, dietary adherence was not objectively measured. These studies are particularly important as they can be compared to results from confined studies to understand how subject measurements potentially deviate from controlled conditions.

Calculation of energy imbalance
The average energy imbalance can be calculated from the average rate of change in body energy stores from the beginning to the end of the study, $ES$ in Equation 1. If $ES$ is negative (negative energy balance), then $EE$ is larger in magnitude than $EI$. Likewise, if $EI$ is positive (positive energy balance), then $EE$ is smaller in magnitude than $EI$. Using the established energy densities (30) of 1,020 kcal kg$^{-1}$ for FFM and 9,500 kcal kg$^{-1}$ for FM, we applied Formula 2 to quantify the average rate of change in body energy stored or lost in kcal d$^{-1}$:

$$ES = 1,020 \frac{\Delta FFM}{\Delta t} + 9,500 \frac{\Delta FM}{\Delta t}$$

where $\Delta FFM$ and $\Delta FM$ represent the change in kg of FFM and FM from beginning to end of the intervention and $\Delta t$ is the time length of the trial in days. Determining energy imbalance from the change in body energy stores (Equation 2) has recently been validated experimentally in several studies (31,32).

Determination of prescribed exercise energy expenditures
If prescribed energy expenditures were not directly reported, we determined the exercise prescription level in kcal d$^{-1}$ by multiplying the reported exercise intensity by the duration of each session. We then averaged the amount over the length of the intervention to arrive at a prescribed energy expended through exercise.

Calculation of the magnitude of compensation
The existing formal definition for degree of compensation during aerobic exercise is the difference in expected weight and actual weight (14). This definition relies on the accuracy of the applied model. We revised the definition of compensation as the magnitude of difference between achieved energy imbalance and prescribed expenditures.

Expected post-intervention resting metabolic rate
The per cent metabolic adaptation is defined as the per cent decrease in RMR beyond expected at the reduced weight

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Figure 1  Diagram that maps the progression of study analysis begins with the exercise intervention which increases exercise energy expenditure. The increase will lead to an energy imbalance which may translate to an energy deficit or an energy surplus. The mechanistic influences on energy imbalance that are examined are changes to resting metabolic rate (RMR), non-exercise activity (NEAT), fat-free mass (FFM) and energy intake (EI). The impact of these changes on energy imbalance translates to changed body weight (W).
Metabolic adaptation has been observed during periods of caloric restriction (33,34). To determine whether metabolic adaptation occurs in response to an exercise program, we require longitudinal RMR data in subjects who maintained baseline EI and achieved a negative energy balance solely through exercise. Therefore, we restricted our analysis of exercise-induced changes in RMR to the study of Bouchard et al. (13), which was the only confined experiment that generated a negative energy balance from exercise and included longitudinal measurements of RMR. Calculation of metabolic adaptation followed previously published methods (33,34).

First, RMR was regressed against FFM at baseline to generate linear equations that were then used to calculate expected values of RMR at 100 d from observed values of FFM (denoted RMR	ext{e}). Differences between expected and observed RMR were analyzed at 100 d through a Bland–Altman analysis (35). Per cent metabolic adaptation was calculated as the average of the individual per cent metabolic adaptation:

\[
1 - \frac{RMR_t}{RMR_e}
\]

where RMR	ext{e} represents actual observed post-intervention RMR.

Calculation of energy intake

When DLW-measured EE is obtained in conjunction with measured body composition, then two terms of the energy balance equation are known, namely ES and EE. From here, we solved for average EI during the course of the intervention:

\[
EI = ES + EE = 1,020 \frac{\Delta FFM}{\Delta t} + 9,500 \frac{\Delta FM}{\Delta t} + EE
\]

Development of fat-free mass–fat mass relationship during aerobic exercise

Bland–Altman analysis of Forbes curve predictions

The Forbes curve (26)

\[
FFM = 10.4 \ln \left( \frac{FM}{C} \right)
\]

where the parameter, C, is a function of baseline body composition,

\[
C = \frac{FM_0}{e^{FM_0/10.47}}
\]

defines a relationship between FFM and FM, and has been successfully applied within differential equation models that predict weight change resulting from changes in EI (36–38). We remark that the curve originally defined by Forbes (26) did not include the parameter, C. The inclusion of the parameter, C, combines the effect of gender, age and race on baseline body composition in a single parameter (36). Although folding these covariates into a single parameter is not optimal, the resulting equation is technically simple to apply within a more complex dynamic model and in the case of caloric restriction provides a comparable degree of accuracy predicting ΔFFM (39).

In order to determine the quality of Forbes curve predictions during an exercise program, we conducted a Bland–Altman analysis comparing Forbes curve-predicted change in FFM (ΔFFM) to observed ΔFFM in a study where negative energy balance was induced solely through caloric restriction (40), a combination of caloric restriction and exercise (40), and solely through exercise (13). We determined whether the level of agreement between Forbes curve predictions and observed data altered with the inclusion of exercise. The statistical analysis was conducted using R 2.11.0 (GNU, 2011).

For the Bland–Altman analysis, we applied data from subjects who achieved weight loss solely through caloric restriction and caloric restriction combined with exercise from a reference database of the 24-week Comprehensive Assessment of Long-term Effects of Reducing Intake of Energy (CALERIE) Study conducted at the Pennington Biomedical Research Center (40). This study was selected because dietary adherence was monitored through multiple simultaneous measurements of ES and EE (32). For the case of weight loss achieved solely through caloric restriction, we limited our analysis to the 12 CALERIE subjects that were placed on a continuous 25% calorie-restricted diet for a period of 24 weeks. The combined diet and exercise database was also taken from the CALERIE study where we limited our analysis to the 12 subjects prescribed a combination of continuous caloric restriction (12.5% calorie restriction) and exercise, which increased energy expenditure by 12.5%. Finally, we analyzed the degree of agreement between Forbes predicted vs. observed ΔFFM in the Bouchard study (13).

The Bland–Altman analysis revealed the magnitude of observed ΔFFM in response to aerobic exercise will be less than predicted by the Forbes curve. Thus, we fit a piecewise-modified Forbes curve for the case where FM decreases below baseline levels (FM\text{b}) and the case where FM increases above FM\text{b}.

The mathematical representation of these cases can be expressed as

\[
FFM = \begin{cases} 
  a_1 \ln \left( \frac{FM}{C} \right) & \text{if } FM < FM_0, \\
  a_2 \ln \left( \frac{FM}{C} \right) & \text{if } FM > FM_0
\end{cases}
\]
To determine the parameter $a_1$, we applied the study by Ross et al. (15,16) where subjects achieved a negative energy balance equivalent to the prescribed increase in physical activity. The other branch of the modified Forbes curve required knowledge of changes in body composition for a cohort who increased intake while on an aerobic exercise intervention. To fit the parameter $a_2$ we applied data from the Inflammation and Exercise study (INFLAME) (2). Forty-two percent of the INFLAME subjects ($N = 28/66$) increased FM. Additionally, 99.9% compliance to exercise protocols was established.

The newly developed model was validated on the data from the study of Bouchard et al. (13).

**Results**

Estimated mean energy imbalance for each reviewed study was calculated using Equation 2 and the results are reported in Table 1.

**Energy imbalance validity**

We compared the amount of energy imbalance achieved calculated from changed body energy stores (Equation 2) with the directly measured energy deficit in confined studies (13,18,19). Estimated mean energy imbalance agreed with directly measured negative energy balance within 80 kcal d$^{-1}$ in the Bouchard study (13), within 145.1 kcal d$^{-1}$ for the Woo Lean study (18) and within 141.7 kcal d$^{-1}$ in the Woo Obese study (19).

Next we compared conclusions obtained from calculated energy imbalance to study conclusions arrived from alternative methods. The Dose-Response to Exercise in postmenopausal Women study (DREW) compared actual weight loss to expected weight loss using the rule that a 3,500 kcal deficit yields a lb of weight loss (14). The DREW analysis determined that subjects completing a higher dose of exercise achieved significantly less weight loss than expected compared to the subjects completing lower exercise doses. Hence, under the former definition of compensation (14), subjects who conducted higher doses of exercise compensated more than subjects who completed lower doses. Similarly, under our modified definition of compensation, the high-dose exercisers compensated more than the low-dose exercisers.

The Woo Lean and Woo Obese studies (18,19) concluded lean subjects increased their intake to remain in energy balance while the obese did not. This observation was supported by our energy imbalance calculation which demonstrated that energy imbalance for the lean subjects was close to zero while the energy imbalance for obese subjects agreed closely with prescribed energy expenditures from exercise.

Studies where the magnitude of energy imbalance agrees with the prescribed energy expended through exercise

The mean reported energy expended from exercise in the Bouchard study (13) was $-580$ kcal d$^{-1}$. When compared to the mean-achieved energy imbalance calculated from Equation 2 we arrived at $-500$ kcal d$^{-1}$. Similarly, mean-achieved energy imbalance ($-701$ kcal d$^{-1}$) closely agreed with amount of prescribed exercise energy expenditures ($-700$ kcal d$^{-1}$) in male subjects of the Ross study where dietary adherence was monitored daily (15,16). We also found that for the Woo Obese study (19), calculated mean energy imbalance was $-511$ kcal d$^{-1}$, which closely agreed with the prescribed energy expended from exercise ($-555$ kcal d$^{-1}$).

Studies where achieved energy imbalance does not agree with prescribed energy expended from exercise

From Table 1 we observe that for the majority of studies, the magnitude of achieved energy imbalance was consistently lower than prescribed energy expenditure from exercise. Because the additional energy expenditures through physical activity were rigorously supervised, the low magnitude of energy imbalance must have been a result of metabolic adaptation, decreased non-exercise activity or a compensatory increase in EI. We explored evidence for each possible response.

**Evidence for metabolic adaptation**

Several reviews have examined the effect of aerobic exercise interventions in sedentary populations on RMR (7,9–11). Several studies report increases in RMR while some report no change (7–11). However, from Table 1 we observe that the energy imbalance in the majority of studies is very low, and therefore the observed response in RMR may have resulted from an insufficient magnitude of negative energy balance. Hence, based on the previous literature, we could not conclusively determine that RMR is unchanged or increased as a result of an aerobic exercise intervention. As reported in a recent review (10), we can only conclude that RMR does not change as long as body weight is maintained. Thus, the question remains whether adaptations would exist if EI was maintained at baseline levels.

In the Bouchard experiment (13), where EI was maintained at baseline levels, FFM accounted for 67% of the variance in RMR ($RMR = 17.3$ FFM + 156.4, $R^2 = 0.67$, $P < 0.0004$), where RMR is expressed in kcal d$^{-1}$ and FFM in kg. The top panel of Fig. 2 illustrates that the observed
Figure 2 Top panel: Linear regression at baseline between resting metabolic rate (RMR) and fat-free mass (FFM, kg) \( (RMR = 17.3 \text{ FFM} + 156.4, \quad R^2 = 0.66) \) and observed RMR values (closed circles) post intervention in the study of Bouchard et al. (13). The observed values falling below the regression line, particularly at lower levels of FFM, indicate metabolic adaptation. Bottom panel: Bland–Altman plot comparing expected vs. post-intervention-observed RMR. The Bland–Altman analysis revealed a bias of \(-80.9 \text{ kcal d}^{-1}\) with 95% confidence intervals of \(-311\) and \(149\), although this bias was not consistent over levels of RMR (slope = 0.45 intercept = \(-608.6\), \(R^2 = 0.39\)), indicating that metabolic adaptation primarily occurs for individuals with lower RMRs or body mass.
RMR values at 100 d are lower than the RMR values that were predicted based on FFM (i.e. the observed RMR values or closed circles are below the regression line), although this discrepancy is largest for individuals who had lower levels of FFM. This is confirmed with the Bland–Altman analysis that revealed a mean bias of $-81 \text{kcal} \text{d}^{-1}$ (95% confidence intervals: $[-148, -13]$; one-sample $t$-test $t = -2.58, P < 0.05$) or 7% ($\pm 1%$ standard deviation), suggesting the existence of exercise-induced metabolic adaptation when baseline EI is maintained (Fig. 2, bottom panel). Further, the Bland–Altman analysis revealed that metabolic adaptation is most pronounced at the lower levels of RMR ($y = 0.5x - 608.53, R^2 = 0.386$), which supports the observations from the regression in the top panel of Fig. 2.

Evidence for a decrease in non-exercise activity

The review (12) found that in the case where body weight is maintained, activity external to prescribed physical activity did not decrease except in elderly subjects. Recent use of accelerometers confirms that no noticeable differences in physical activities not associated with the exercise intervention can be detected (4). In existing studies that collected accelerometer data, there is little change in body weight and lack of objectively measured dietary intake precludes extrapolation to a case when large negative energy balance is induced from exercise. To our knowledge, there does not exist a study that measured non-exercise movements when EI was maintained at baseline levels. The only evidence that non-exercise activity is not a major contributor to low weight loss is that the energy imbalance determined from the studies of Bouchard et al. and Ross et al. (13,15,16) does not significantly differ from prescribed increased exercise expenditures.

Evidence for dietary compensation

Because of the previously mentioned difficulty in measuring the different contributors to the terms in the energy balance equation during an exercise intervention, one cannot simply use the low magnitude of imbalance calculated from changed body energy stores and conclude the presence of a compensatory increase in dietary intake. Errors in self-reported intake prevent the use of dietary records as a reliable measure of changes in intake during an exercise intervention (41). Thus, we must restrict our analysis to studies that measured both changed body composition and energy expenditures by DLW (4).

Twelve of 13 individuals from which changed body composition and DLW-assessed energy expenditures were obtained in the study of Westerterp et al. (4) increased intake during the respective intervention. Additionally, Woo et al. (18) found that five lean women increased intake to match their increased energy expenditures during the intervention, but did not observe the same result for the six obese women (19).

Does exercise preserve lean tissue?

Fourteen of the 15 aerobic exercise intervention studies resulted in very little change in FFM despite changes in FM. From this, it may be tempting to conclude that aerobic exercise preserves lean tissue; however, as stated previously, EI in the majority of these studies was unknown. Thus, the observed preservation of FFM may be a result of increased EI.

Forbes observed FFM loss during aerobic exercise and hypothesized that the response is similar to those observed in caloric restriction interventions (27–29). Loss of FFM was similarly observed in 51% (34/66) individual subjects in the INFLAME study (2). However, the large magnitude of achieved energy imbalance in the Bouchard and Ross studies (13,15,16), resulted in no significant change in FFM. Thus, the question remains to what degree FFM will decrease in response to exercise. It has been suggested that the Forbes curve predicts changes in FFM during exercise (29,36–38) and we begin by examining this hypothesis.

The Forbes curve

The original Forbes curve is well established as a model describing changes in FFM during caloric restriction (26,36–38). This observation is reconfirmed by our Bland–Altman analysis from the Forbes curve predictions of the change in FFM resulting solely from caloric restriction in the CALERIE Phase I study (40). In this case, the Bland–Altman analysis (first row of Fig. 3) indicates good agreement between Forbes curve predictions of change in FFM and actual change in FFM (Pearson correlation coefficient, $R = 0.70, P = 0.02$) with a low bias of 0.26 kg (95% limits of agreement $-1.80, 2.34$ kg).

However, with the introduction of aerobic exercise, agreement of the predicted changes in FFM by the Forbes curve weakens (second row of Fig. 3). Our Bland–Altman analysis revealed an increased bias (1.16 kg, 95% limits of agreement $-2.02, 4.35$ kg) with caloric restriction combined with exercise and the correlation between predicted change in FFM and actual change in FFM is not significant ($R = 0.35, P = 0.30$). This bias increases further (third row of Fig. 3, bias $= 3.63$ kg, 95% limits of agreement $-0.52, 7.78$ kg) in the case of negative energy balance achieved solely through aerobic exercise, with a weaker correlation that is again not significant ($R = 0.06, P = 0.72$).

Our analysis reveals that the Forbes curve overestimates the decrease in FFM during aerobic exercise. As a result, we modified the Forbes curve to account for the smaller observed decrease.
Figure 3 The Forbes curve does not predict ΔFFM during aerobic exercise. Bland–Altman plots depicting Forbes curve predictions of ΔFFM (kg) for caloric restriction alone from Comprehensive Assessment of Long-term Effects of Reducing Intake of Energy (CALERIE) Phase I (40) (first row), caloric restriction combined with aerobic exercise from CALERIE Phase I (40) (second row) and aerobic exercise alone from the Bouchard trial (13) (third row). The Bland–Altman indicates good agreement for caloric restriction alone ($R^2 = 0.52$, $P = 0.02$ [significant correlation] with a low bias [0.3 kg]). The bias increases for caloric restriction combined with exercise (1.2 kg) and the correlation is not significant ($R^2 = 0.13$, $P = 0.30$). The bias increases further for aerobic exercise alone (3.6 kg) and the correlation is once again not significant ($R^2 = 0.01$, $P = 0.727$). FFM, fat-free mass.
The modified Forbes curve for aerobic exercise

We restricted our model development to the Ross study data (15,16) and the INFLAME subjects (2) who increased intake. Applying these two data sets, the parameters $a_1$ and $a_2$ were determined so that the mean absolute error between final-predicted FFM and model-predicted FFM in Equation 4 is minimized yielding the formula:

$$F_{FM} = \begin{cases} 7 \ln \left( \frac{FM}{C} \right) & \text{if } FM < 0, \\ 13.7 \ln \left( \frac{FM}{C} \right) & \text{if } FM > 0 \end{cases}$$ (5)

We validated the newly developed model on the Bouchard study (13) and found the mean absolute error between predicted and observed change in FFM decreased by 89% in comparison to the classic Forbes curve (Table 2).

The newly developed model predicted a smaller decrease in FFM compared to the Forbes curve. Likewise, if energy intake was increased, the new model predicted a larger increase in FFM than predicted by the Forbes curve. Figure 4 depicts how movement along the newly developed FFM-FM model differs from the traditional Forbes curve.

### Discussion

The present study applied an energy balance analysis to group and individual data from 15 different aerobic exercise studies to analyze why individuals do not lose more weight through prescribed exercise. These analyses and

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**Table 2** Modified Forbes model predictions vs. observed changes in fat-free mass

<table>
<thead>
<tr>
<th>Gender</th>
<th>Baseline FM (kg)</th>
<th>Baseline FFM (kg)</th>
<th>Final FM (kg)</th>
<th>Final FFM (kg)</th>
<th>Forbes-predicted final FFM (kg)</th>
<th>Forbes error (kg)</th>
<th>Modified Forbes-predicted FFM (kg)</th>
<th>Modified Forbes error (kg)</th>
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<tbody>
<tr>
<td>Ross et al. (15,16)</td>
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<tr>
<td>M</td>
<td>33.1</td>
<td>67.9</td>
<td>27</td>
<td>67</td>
<td>65.78156</td>
<td>-1.2</td>
<td>66.7</td>
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<td>F</td>
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<td>43.75422</td>
<td>-2.6</td>
<td>44.5</td>
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<td>INFLAME (2)</td>
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<td></td>
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<td>33.9</td>
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<td>F</td>
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<td>47.1</td>
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<tr>
<td>M</td>
<td>20.8</td>
<td>61.2</td>
<td>15.9</td>
<td>61.1</td>
<td>58.4</td>
<td>-2.7</td>
<td>59.6</td>
<td>0.3</td>
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Actual FFM (kg), Forbes and modified Forbes-predicted FFM (kg) with calculated error (kg) (difference between actual and predicted) in the Ross (15,16) and INFLAME (2) studies on which the modified model was developed. The model was validated on the Bouchard data set (13).

FFM, fat-free mass; FM, fat mass; INFLAME, Inflammation and Exercise study.

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**Figure 4** Plot of a proposed modified piecewise Forbes curve (dotted and solid, modelling body composition during aerobic exercise superimposed over the original Forbes curve [dashed] describing body composition changes during caloric restriction). The top curve models the observation of less fat-free mass (FFM) and more fat mass (FM) during weight loss and assumes greater FFM and less FM during weight gain. Extrapolation of the model beyond available data is indicated by the dotted portion of the top curve.
observations yield the following conclusions and remaining uncertainties.

What is the energy deficit induced by exercise?

We have computed the energy imbalance from change in body energy stores generated in 15 different exercise studies. We have shown that the magnitude of energy imbalance generated by 13 of the 15 studies is lower than the typical amount of caloric restriction imposed in the majority of weight loss studies (<500 kcal d⁻¹). We have shown that the achieved energy balance is equal to the prescribed energy expenditures from exercise in a confined study that maintained baseline energy intake and a study that monitored subject energy intake through daily supervision. In the remaining studies, we have shown that the magnitude of negative energy balance is lower than prescribed exercise energy expenditures.

We investigated whether the small energy imbalance is a result of physiological adaptations or increased dietary intake.

We analyzed whether a possible decrease in RMR may contribute to the low energy imbalance by examining the only study that maintained baseline energy intake for the duration of a supervised exercise intervention in confined subjects. We found mean RMR in this study to be depressed by 7% beyond weight-related reductions suggesting that there may be an adaptive response if exercise is conducted in combination with an isocaloric diet. The depressed RMR was more pronounced in lean subjects than obese; an observation that may correspond to different dietary responses during exercise observed by Woo et al. (18,19). These results indicate that RMR is reduced when exercise is increased and energy intake is held constant, particularly among individuals with lower levels of FFM and hence lower RMR. This metabolic adaptation reduces the size of the exercise-induced energy deficit and preserves body mass, primarily for leaner individuals.

Unfortunately, the lack of knowledge of how much dietary compensation may have occurred in the remaining exercise studies clouds our understanding of whether there are adaptations to RMR if intake remains unchanged from baseline. To confirm and quantify the degree of adaptation that may be present, more studies that measure RMR alongside DLW and body composition during exercise interventions are critical.

A surprising observation was the repeated finding that almost no change to RMR occurs in response to an exercise intervention with ad libitum energy intake (7). This observation, combined with the results of the Bouchard and Woo studies (13,18,19), leads to a hypothesis that increased dietary intake in response to exercise-induced energy imbalance could be driven by RMR preservation.

Finally, we remark that there does not exist non-exercise activity data collected during an exercise intervention where baseline energy intake was maintained. The evidence that energy imbalance in Bouchard (13) and Ross (13,15) studies closely match prescribed energy expenditure from exercise suggests if it exists, non-exercise activity does not change substantially during an exercise intervention. The increased use of accelerometers in combination with the aforementioned body composition and energy expenditure measurements would provide information to analyze how exercise may influence non-exercise activity, assuming that the accelerometers are sensitive enough to detect these possible changes.

Our analysis of 13 subjects whose energy intake was measured through multiple DLW and body composition measurements (4) revealed that 12 of the subjects (~92%) increased dietary intake. We note that this particular intervention involved an intense training program, which prepared previously sedentary subjects to compete in a half marathon. The increased dietary intake observed in the Westerterp subjects (4) is supported by several studies which found similar results at higher exercise doses (14,42–44). It is noteworthy that the discrepancy between achieved energy imbalance and prescribed energy expenditures is larger in magnitude with higher exercise dose (3,4,14,22) (Table 1). This suggests that an important contributor to low energy imbalance when performing a high dose of aerobic exercise is increased dietary intake.

To test whether the degree of compensation is a function of baseline body composition as observed by Woo et al. (18,19), we require additional data containing DLW-measured EE and simultaneously measured body composition in subjects with a wide range of BMI who are undergoing aerobic exercise. Unfortunately, subjects classified as obese in the only existing exercise study which obtained DLW-measured EE combined with body composition did not complete the study (4). Additional EI data are required to confirm the Woo hypothesis in free-living subjects classified as obese.

What are the expected changes in body composition resulting from an aerobic exercise intervention?

In two studies where EI was maintained at baseline levels and generated a negative energy balance solely through aerobic exercise, we found that less FFM and more FM is lost than predicted by the Forbes curve. We also found that
as EI is increased during aerobic training, more gain in FFM and less gain in FM is observed than predicted by the Forbes curve. We have developed and validated a modified piecewise-defined Forbes curve that predicts body composition changes induced by aerobic exercise. Additional data that contain EI, EE and body composition are needed to further advance and validate this model.

We conclude that the small magnitude of weight loss observed from the majority of evaluated exercise intervention studies is primarily due to low doses of prescribed exercise energy expenditures compounded by a concomitant increase in caloric intake; however, exercise has additional beneficial effects that were not considered in the present analysis.

Knowledge of how increased exercise induces physiological changes in the body will improve not only our knowledge of how to treat obesity but stands to improve population-wide health.

Conflict of interest statement

Timothy Church is a member of the Jenny Craig Medical Advisory Board. Claude Bouchard is a consultant for Weight Watchers International and a member of the Scientific Advisory Board of Pathway Genomics.

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