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Protein intake induced an increase in exercise stimulated fat oxidation during stable body weight☆

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ABSTRACT

Background: Protein-rich weight-loss diets spare fat-free mass at the cost of fat mass. The objective was to examine if there is a change in stimulated fat oxidation related to protein intake during stable body weight.

Methods: Subjects’ (BMI 22 ± 2 kg/m², age 25 ± 8 years) maximal fat oxidation (Fatmax) was assessed during a graded bicycle test, before and after a 3-month dietary-intervention of 2 MJ/day supplements exchanged with 2 MJ/d of habitual energy intake. The parallel design consisted of protein-rich supplements in the protein group and an isocaloric combination of carbohydrate and fat supplements in the control group. Daily protein intake was determined according to 24-h urine nitrogen. Body composition was measured according to a 4-compartment model by a combination of underwater-weighing technique, deuterium-dilution technique and whole-body dual-energy X-ray absorptiometry (DXA).

Results: Subjects were weight stable and did not change their physical activity. The protein group (n=12) increased protein intake (11 ± 14 g, P<0.05) and had significantly higher daily protein intake vs. control (n=4) (80 ± 21 vs. 59 ± 11 g, P<0.05). Fatmax increased significantly in the protein group (0.08 ± 0.08 g/min, P<0.01). Fat-free mass increased independent of change in body weight (P<0.01), and fat mass and fat percentage decreased (P<0.05). Change in Fatmax was a function of change in protein intake (r=0.623, P=0.05), and not of changes in body composition or VO2max.

Conclusion: Increased stimulated fat oxidation was related to increased protein intake.

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1. Introduction

Obesity is a condition in which fat mass (FM) and fat percentage are increased [1] and levels of fat oxidation are suggested to be disturbed [2]. Fat and carbohydrate oxidation are mainly influenced by exercise intensity [3]. With increasing exercise intensity, fat oxidation first increases to its maximal fat oxidation rate (Fatmax) from low- to moderate-exercise intensities and then decreases from moderate- to high-exercise intensities [3]. The daily majority of energy demand is at rest or during moderate-exercise intensity. At rest and during moderate-exercise intensity, fat oxidation is the main source of energy production for the body [4]. So, moderate-exercise intensity yields the most grams of fat used for oxidation and could therefore play a role in the maintenance of or reduction in FM. The desired goal for the treatment of and the reduction in development of obesity is to decrease FM while preserving or increasing fat-free mass (FFM). Increased protein intake has shown to result in greater loss of FM and lower loss of FFM during energy restriction, and lower regain of FM and greater regain of FFM during weight regain after weight loss [5–7]. The resulted higher ratio of FFM to FM plays an important role in the maintenance of energy balance [8] and the preservation of metabolic and overall health [9,10]. Since elevated protein intake results in a more favorable body composition during weight loss and weight maintenance thereafter, and since FM is mainly reduced during moderate-exercise intensity, the question remains whether these characteristics hold when subjects are in conditions of energy balance. Therefore, the aim of this study was to investigate whether a change in dietary protein might change stimulated fat oxidation during exercise in subjects with constant body weight over time.

2. Subjects and methods

2.1. Subjects

Subjects were recruited by means of an advertisement in local newspapers and on notice boards at Maastricht University. Subjects who were willing to participate in the study were subsequently screened, by means of a detailed medical history and a physical examination. All subjects were in good health, non-smokers, at most moderate alcohol users, did not use prescription medication, and did not...
fluctuate more than 2 kg in body mass over at least the last 2 months. All subjects gave a written informed consent. The Medical Ethics committee of the University and Academic Hospital of Maastricht approved the study. Twenty-five subjects started in the study, 11 men and 14 women. Eight subjects (5 men and 3 women, and 4 subjects in each group) dropped out due to several reasons, such as personal and an inability to fulfill the schedule with visits to the university. Dropouts were not different from completers in baseline body weight, BMI and body composition, Fatmax, physical activity or protein intake. One subject of the control group had excessive protein intake at baseline (216 g at baseline and 89 g during the intervention), and was removed from the analysis. Subject characteristics (n = 16; 12 in the protein group and 4 in the control group) are given in Table 1.

2.2. Study design

The study had a randomized parallel design and consisted of a dietary intervention period of 3 months. A test day that included measurement of substrate oxidation during a graded bicycle test to exhaustion, measurements of body composition, blood sampling and completing questionnaires took place at baseline and after 3 months of intervention. Subjects were familiarized with the equipment and the procedures before the start of all measurements. Measurements were performed in the morning after an overnight fast. The bicycle-ergometer test started at the same time in the morning to avoid circadian variance. The day before both test days, subjects were asked to refrain from alcohol, refrain from indulgence in strenuous exercise and refrain from eating and drinking after 11:00 PM. Subjects were instructed to maintain their baseline body weight and to maintain their customary level of physical activity during the entire duration of the study.

2.3. Dietary intervention

Subjects were counseled to consume isocaloric diets to sustain body weight by exchanging 2 MJ of their habitual energy intake with 2 MJ of milk-proteins to be incorporated within the subjects’ habitual diets to increase daily protein intake. These protein supplements were rich in essential micronutrients and were supplied in three sachets daily containing in total of 52 g of milk-protein, dissolved in water to obtain a milk shake, pudding, soup or muesli (Modifast, Novartis Nutrition, Breda, The Netherlands). The control group received isocaloric carbohydrate-fat supplements consisting of a limonade (Karvan Cevitam, Koninklijke de Rijtzer, Zeist, The Netherlands) and of olive oil. All subjects were instructed to consume daily at least 200 g of fruit and 300 g of vegetables.

To assess dietary protein intake, subjects completed three 24-h urine collections at baseline and in weeks 6 and 12. Samples were samples were collected with 10 mL H2SO4 to prevent nitrogen loss through evaporation, stored frozen at −20 °C, and later analyzed for urinary nitrogen with a nitrogen analyzer (CHN-O-Rapid; Heraeus, Hanau, Germany).

2.4. Anthropometry

To monitor body weight stability, subjects were instructed to measure their body weight daily at home. At the University, body weight was measured 2-weekly using a digital balance (Chyo-MW-150 K, Chyo, Japan; weighing accuracy 0.02 kg) with subjects in underwear, in the fastest state and after voiding their bladder. If body weight fluctuated by > 2 kg from baseline body weight, subjects were instructed to adjust their energy intake to encourage a return to and maintenance of baseline body weight. Height was measured at baseline to the nearest 0.1 cm using a wall-mounted stadiometer (Seca, model 220, Hamburg, Germany). Body mass index (BMI) was calculated by dividing body weight by height squared (kg/m²).

2.5. Body composition

Body composition was assessed in the fastest state with the 4-compartment model of Lohman [11]. The model was used to calculate percentage fat mass (%FM) from the independently determined whole-body density (Db), total body water (TBW) and total bone mineral content (BMC). Measuring whole-body density, total body water and total bone mineral content separately increases the accuracy of FM and FM at baseline and after the intervention and is therefore more suitable to determine changes in FM and FM, especially if subjects sustain their body weight. All measurements were completed within the same

### Table 1

<table>
<thead>
<tr>
<th>Group</th>
<th>Baseline Total</th>
<th>Control N=16</th>
<th>Protein N=12</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>24.8±7.6</td>
<td>24.8±9.2</td>
<td>24.8±7.5</td>
</tr>
<tr>
<td>Body height (cm)</td>
<td>172.7±10.0</td>
<td>172.9±2.1</td>
<td>172.7±11.7</td>
</tr>
<tr>
<td>Body weight (kg)</td>
<td>66.0±8.6</td>
<td>65.5±1.7</td>
<td>66.1±9.9</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>22.1±1.7</td>
<td>21.9±1.7</td>
<td>22.1±1.8</td>
</tr>
<tr>
<td>FFM (kg)</td>
<td>52.1±10.8</td>
<td>51.9±3.4</td>
<td>52.1±12.5*</td>
</tr>
<tr>
<td>FM (%)</td>
<td>37.7±4.5</td>
<td>35.6±4.6</td>
<td>37.8±4.7*</td>
</tr>
<tr>
<td>FFA (μmol/L)</td>
<td>214.7±7.7</td>
<td>206.0±6.1</td>
<td>216.8±4.7*</td>
</tr>
<tr>
<td>Protein intake/gd</td>
<td>427±121</td>
<td>409±94</td>
<td>433±132</td>
</tr>
<tr>
<td>Protein intake/body weight (g/kg)</td>
<td>67±17</td>
<td>58±13</td>
<td>70±17*</td>
</tr>
<tr>
<td>Fatmax (g/min)</td>
<td>0.0±0.2</td>
<td>0.0±0.2</td>
<td>1.1±0.2*</td>
</tr>
<tr>
<td>VO2max (ml/min)</td>
<td>2804±835</td>
<td>2549±597</td>
<td>2889±932</td>
</tr>
<tr>
<td>VO2max/FM (ml/min/kg)</td>
<td>50.6±7.4</td>
<td>46.7±0.5</td>
<td>51.9±6.5</td>
</tr>
<tr>
<td>Baeckec</td>
<td>9.2±1.4</td>
<td>8.8±1.0</td>
<td>9.4±1.5</td>
</tr>
<tr>
<td>Sportd</td>
<td>3.5±0.9</td>
<td>3.6±0.7</td>
<td>3.5±0.9</td>
</tr>
<tr>
<td>Leisure timed</td>
<td>3.5±0.4</td>
<td>3.3±0.4</td>
<td>3.3±0.4</td>
</tr>
<tr>
<td>Workd</td>
<td>2.3±0.7</td>
<td>2.1±0.2</td>
<td>2.4±0.8</td>
</tr>
</tbody>
</table>

Mean values and standard deviations.

P<0.05, P-value of paired Student’s t-test over time, baseline compared to after 3 months.

a Body mass index (BMI kg/m²) was calculated as body weight (kg) divided by height (m) squared.

b Body composition of the 4-compartment model of Lohman; "FM = (274.7/Db – 71.4 TBW/BM + 114.6 BMC/BM - 205.03)."

c Based on 24-h urinary nitrogen content, 3-month value is the average of 1.5 and 3 months.

d Plasma concentrations after overnight fasting, n=15 at 3 months.

The Baecke total activity index and its activity subscores of sport, leisure time and work.
morning. The used equation of Lohman was %FM = \left(\frac{2.747}{\text{Db}} - 0.714 \times \text{TBW}/\text{BM} + 1.146 \times \text{BMI}/\text{BM} - 2.0503\right) \times 100 [11].

Whole-body density was measured with the underwater weighing technique. Body mass in air and underwater was determined on a digital balance, accurate to 0.01 kg (Sauer type E1200). Residual lung volume was measured simultaneously with the helium dilution technique using a spirometer (Volugraph 2000, Mijnhardt, The Netherlands).

Total body water was measured using the deuterium (\(^2\text{H}_2\text{O}\)) dilution technique according to the Maastricht protocol [12]. In the evening before the test day, subjects collected a background urine sample and then ingested a dose of deuterium-enriched water, after which they refrained from consuming fluid and food. The following morning, a urine sample from the second voiding was collected between 08:00 and 10:00 h. The concentration of deuterium in the urine samples was measured using an isotope ratio mass spectrometer (Micromass Optima, Manchester, UK). The dilution of the deuterium isotope is a measure for TBW. Total body water was obtained by dividing the measured deuterium dilution space by 1.04 to correct for exchange of the \(^2\text{H}\) label with non-aqueous hydrogen of body solids [13].

Total bone mineral content (BMC) was measured using dual-energy X-ray absorptiometry (DXA; Lunar Corp., Madison, WI) with a resolution of 4.8 × 9.6 mm (whole-body). Bone content and density were calculated by Lunar software (version 1.3z). The subjects wore loose metal-free clothing and remained in a supine position while scanning was completed. The results were compared with the Germany Total Body White Reference Population provided by the manufacturer.

2.6. Substrate oxidation during the cycle-ergometer test

Subjects performed a graded exercise test to exhaustion on an electromagnetically braked bicycle ergometer (Excalibur, Lode, Groningen, The Netherlands). Simultaneous indirect calorimetric breath-by-breath measurements enabled to measure fat oxidation rate over a wide range of intensities. Fat oxidation rate was plotted against exercise intensity, expressed as %VO\text{max}, power output and heart rate, constructing a fat oxidation curve to accurately determine maximum fat oxidation (Fat\text{max}). The cycle ergometer protocol has been validated to determine the exercise intensity that elicits Fat\text{max} [14]. Moreover, it has been shown that the early stages of this type of protocol do not influence the exercise intensity at Fat\text{max} [14].

Workload of the cycle ergometer test was calculated for each subject to standardize the cycling protocol between subjects. Subjects started cycling at a workload of 0.5 W/kg FFM. Workload was increased by 0.5 W/kg FFM every 3 min until exhaustion. Maximal workload was calculated as the last completed stage plus the fraction of time spent in the final non-completed stage multiplied by the workload increment.

Indirect calorimetric measurements were performed using an Omnicr IV gas analysis system. The gas analyzers were calibrated with a 18% \(\text{O}_2/0.8\% \text{CO}_2/81.2\% \text{N}_2\) gas mixture with an uncertainty of 1% relative (Linde Gas Benelux B.V., Dieren, The Netherlands). Average values for oxygen uptake (VO\(_2\)) and carbon dioxide production (VCO\(_2\)) were calculated over the last 2 minutes of each 3 minutes stage, during which the RER was <1. VO\text{max} was calculated as the average oxygen uptake of the 3 highest sequent oxygen uptakes during the last 60 s of the cycle test. Stages of 3 min can be used to accurately determine VO\text{max} [15]. Fat and carbohydrate oxidation and energy expenditure were calculated using stoichiometric equations and appropriate energy equivalents, with the assumption that the urinary nitrogen excretion rate was negligible during the cycle test.

Heart rate was recorded continuously during the test by using a radio telemetry heart rate monitor (Polar S610, Polar Electro Ltd., Oy, Finland).

2.7. Physical activity

To determine physical activity the Baecke questionnaire was used. This validated questionnaire explained 48% of the variation in Physical Activity Level (PAL) as measured with doubly labeled water [16]. The Baecke questionnaire consists of three components: work activity, sports activity and leisure activity.

2.8. Blood samples

Fasting venous blood samples were taken to determine concentrations of plasma FFA, TAG. The blood samples were collected in tubes containing EDTA to prevent clotting. Plasma was obtained by centrifugation (1500×g for 10 min at 4 °C), frozen in liquid nitrogen and stored at −80 °C until analysis. FFA concentrations were measured using the Wako NEFA C-kit (Wako Chemicals, Neuss, Germany).

3. Statistical analysis

Groups were compared by ANOVA. Changes over time within groups were compared by 2-tailed paired Student t-tests. Regression analyses were used to analyze the relation between fat oxidation and body composition or protein intake. All analyses were performed with the Statistical Package for the Social Sciences (SPSS) version 16.0.2 for Macintosh OS X. Differences were regarded as significant if \(P<0.05\).

4. Results

Body weight and BMI did not change significantly over the intervention period (Table 1). Protein intake determined according to 24-h urinary nitrogen was significantly different between the protein and control group during the 3-months of intervention (80±21 vs. 59±11 g; \(P<0.05\); or 1.2±0.2 vs. 0.9±0.1 g/kg body weight, \(P<0.05\)). The protein group significantly increased in protein intake (11±14 g, \(P<0.05\); and 0.1±0.2 g/kg body weight, \(P<0.05\)). Baseline protein intake did not differ between groups.

Fat\text{max} significantly increased in the protein group (0.08±0.08 g/min, \(P<0.01\)). Taking covariate change in body weight or change in VO\text{max} into account Fat\text{max} still significantly increased in the protein group (\(P<0.05\)). Taking change in FFM into account, Fat\text{max} changed as a trend (\(P=0.060\)). Fat-free mass increased, and FM and fat percentage decreased over time in the protein group (\(P<0.01\), \(P<0.05\), \(P<0.05\)), which resulted in a significant change in fat percentage over time between the protein and control group (\(P<0.05\)). Total activity index of the Baecke questionnaire and its subscores sport, leisure time and work, VO\text{max} and VO\text{max}/FFM did not change.

The increase in Fat\text{max} in the protein group was a function of the increase in protein intake (\(r=0.623, P<0.05\)). So, increased protein intake explained 39% of the variation in increased Fat\text{max}. At baseline, Fat\text{max} was a function of body weight (\(r=0.621, P<0.05\)), FFM (\(r=0.604, P<0.05\)), and VO\text{max} (\(r=0.621, P<0.05\)). Change in Fat\text{max} was not a function of change in body weight, change in FFM, or change in VO\text{max}.

At baseline, all subjects cycled on average 25 min 52 s ± 3 min 37 s and reached a maximal power output of 226 ± 67 W and maximal heart rate of 187 ± 7 beats per minute during the graded bicycle test to exhaustion. Fig. 1 illustrates subjects’ fat oxidation in function of their VO\text{max} and power output during the bicycle test at baseline. Fat\text{max} was 0.43±0.10 g/min at 51±10% VO\text{max}, and at a power output of 97±35 W and a heart rate of 130±16 beats per minute. Fat\text{max} resulted in 63±17% of energy expenditure with a carbohydrate oxidation of 0.65±0.46 g/min.

5. Discussion

Increase in protein intake, without an increase in physical activity or fitness, increased stimulated fat oxidation and FFM independently of body weight over a 3-month intervention period. The change in protein intake explained 39% of the variation in change in stimulated fat oxidation.
oxidation during the incremental bicycle test. The variation in this Fat\textsubscript{max} was mainly determined by protein intake and FFM.

An increase in protein content of the diet results into a shift in 24-h substrate balances [17]. Protein balance shifts toward a positive balance and fat balance shifts toward a negative balance. In our subjects these shifts in macronutrient balances resulted in an increase in FFM and a decrease in FM and fat percentage without a change in body weight over a period of 3 months. The negative fat balance and subsequent decrease in FM imply that overall more fatty acids were transported out of the FM into the circulation instead of being transported into the FM as a fat-deposit. In line with this perspective, the dietary fat oxidation was previously inversely related to fat percentage and BMI, with lean subjects having the highest and obese subjects the lowest values of dietary fat oxidation [18]. Therefore, lean subjects could take up less dietary fat in their FM as a fat-deposit. Moreover, a lower ratio of FM to FFM results in a higher daily energy expenditure within a subject, as FFM is determining the resting metabolic rate [19] as well as 24-h energy expenditure [20]. The relationship between a positive protein balance, muscle protein synthesis, and change in FFM has only been suggested but has not been determined quantitatively. Part of this relationship has been revealed; protein rich diets result into net muscle-protein synthesis [10,21,22], and increase energy expenditure due to the high dietary fat in their FM as a fat-deposit. In line with this perspective, the dietary fat oxidation was previously inversely related to fat percentage and BMI, with lean subjects having the highest and obese subjects the lowest values of dietary fat oxidation [18]. Therefore, lean subjects could take up less dietary fat in their FM as a fat-deposit. Moreover, a lower ratio of FM to FFM results in a higher daily energy expenditure within a subject, as FFM is determining the resting metabolic rate [19] as well as 24-h energy expenditure [20]. The relationship between a positive protein balance, muscle protein synthesis, and change in FFM has only been suggested but has not been determined quantitatively. Part of this relationship has been revealed; protein rich diets result into net muscle-protein synthesis [10,21,22], and increase energy expenditure due to the high dietary fat in their FM as a fat-deposit. Moreover, a lower ratio of FM to FFM results in a higher daily energy expenditure within a subject, as FFM is determining the resting metabolic rate [19] as well as 24-h energy expenditure [20].

Body composition, stimulated fat oxidation and protein intake have been determined with high accuracy. Combination of densitometry with measurements of total body water and bone mineral content determined the 4-compartment model of subjects’ body composition. Furthermore, we used 24-h urine nitrogen concentrations, which are more applicable to quantify protein intakes because dietary record methods are prone to misreporting [27]. The graded bicycle test with continuous indirect calorimetric measurements allowed precise determination of maximal fat oxidation.

The relative exercise intensity that elicited maximal fat oxidation of 51%VO\textsubscript{2}\text{max} after an overnight fast in our subjects is comparable with 43–64%VO\textsubscript{2}\text{max} observed in previous studies after an overnight fast in trained and untrained men [28,29] and with 48–56%VO\textsubscript{2}\text{max} in a post-absorptive state 3–4 h after a meal in men and women [30,31].

The 4-compartment method to determine the body composition used, the biomarker nitrogen used to determine the amount of daily protein intake, and the graded exercise test to exhaustion with simultaneous indirect calorimetric breath-by-breath measurements used to determine the maximum fat oxidation are the strengths of this study. Regardless of the significant findings presented, the limited amount of subjects is a limitation of the study, which limits the possibility to examine whether the association between change in Fat\textsubscript{max} and change in protein intake is mediated and modified by change in body composition and change in physical activity, respectively.

In conclusion, in subjects with increased protein intake stimulated fat oxidation was increased. Stimulated fat oxidation was a function of protein intake, body composition and VO\textsubscript{2}\text{max}. Yet, change in stimulated fat oxidation was mainly a function of change in protein intake.

**Acknowledgments**

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**References**


**Fig. 1.** Fat oxidation in function of VO\textsubscript{2}\text{max} and power output during a graded bicycle test to exhaustion. CHO\textsubscript{ox} is the amount of carbohydrate oxidized in kJ/min. Fat\textsubscript{ox} is the amount of fat oxidized in kJ/min. E\textsubscript{Eox} is the amount of total energy expenditure in kJ/min. HR is heart rate in beats/min.


