Comparison of the effects of a high- and normal-casein breakfast on satiety, ‘satiety’ hormones, plasma amino acids and subsequent energy intake

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The present study compared the effects of a high- and normal-casein-protein breakfast on satiety, ‘satiety’ hormones, plasma amino acid responses and subsequent energy intake. Twenty-five healthy subjects (BMI 23.9 (SEM 0.3) kg/m²; age 22 (SEM 1) years) received a subject-specific standardised breakfast (20% of daily energy requirements): a custard with casein as the single protein source with either 10, 55 and 35 (normal-casein breakfast) or 25, 55 and 20 (high-casein breakfast) % of energy (En%) from protein, carbohydrate and fat respectively in a randomised, single-blind design. Appetite profile (visual analogue scale; VAS), plasma glucose, insulin, glucagon-like peptide 1, ghrelin and amino acid concentrations were determined for 4 h; here the sensitive moment in time for lunch was determined. Subjects came for a second set of experiments and received the same custards for breakfast, and an ad libitum lunch was offered at 180 min after breakfast; energy intake was assessed. There were increased scores of fullness and satiety after the 25 En% casein-custard compared with the 10 En% casein-custard, particularly at 180 min (26 (SEM 4) v. 11 (SEM 5) mm VAS; P<0.01) and 240 min (13 (SEM 5) v. –1 (SEM 5) mm VAS; P<0.01). This coincided with prolonged elevated plasma amino acid concentrations; total amino acids and branched-chain amino acids were higher after the 25 En% casein-custard compared with the 10 En% casein-custard at 180 and 240 min (P<0.001). There was no difference in energy intake (3080 (SEM 229) kJ for 25 En% and 10 En% respectively; NS) from the ad libitum lunch. In conclusion, a breakfast with 25% of energy from casein is rated as being more satiating than a breakfast with 10% of energy from casein at 3 and 4 h after breakfast, coinciding with prolonged elevated concentrations of plasma amino acids, but does not reduce subsequent energy intake.

Satiety: Energy intake: Casein protein: Glucagon-like peptide 1: Ghrelin: Protein kinetics

The increasing incidence of obesity is considered as a major health problem due to its co-morbidity of a number of diseases, including type 2 diabetes mellitus, CVD, and certain types of cancer9,10. Obesity is the result of a positive energy balance due to energy intake exceeding energy expenditure. In the system of body-weight regulation several pathways are involved and therefore weight management requires a multi-factorial approach10. Recent findings suggest that a relatively elevated protein intake seems to play a role during weight loss as well as during weight maintenance thereafter10. In addition to the protein-induced satiety that has been shown after a high-protein diet, protein-induced satiety has also been shown after a single meal11–16. Several studies on different types of protein affecting satiety have been executed11–16. The question remains, however, whether the larger satiating effects of high-protein meals hold for each specific type of protein.

Abbreviations: AUC, area under the curve; En%, percentage of energy; GLP-1, glucagon-like peptide 1; VAS, visual analogue scale.

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Casein is a part of milk protein; it comprises 80% of the protein content of bovine milk17. Casein is considered as a ‘slow’ protein because it coagulates in the stomach and delays gastric emptying18. The slower digestion rate of casein results in smaller but prolonged increased postprandial plasma amino acid levels18,19. If the extent of postprandial increase in circulating amino acids influences satiety, as was hypothesised by the amino static theory of Mellinkoff et al., consumption of different levels of casein-protein in a single meal should result in differences in subsequent satiety. We investigated possible differences in satiety ratings between a high- and normal-casein-protein concentration and the mechanisms accompanying those differences. Casein was offered in a breakfast consisting of 20% of the subject-specific daily energy requirements, with amounts of casein that represent the highest allowed protein intake per d, i.e. 25% of energy (En%) from protein v. the lowest (normal)
protein intake per d, 10 % of energy from protein\textsuperscript{(21)}. Protein was exchanged with fat; carbohydrate content was kept constant at a level of 55 En% because of its effects on protein metabolism\textsuperscript{(22)}, resulting in a comparison of a high-protein–low-fat breakfast with a normal-protein–normal-fat breakfast with casein as the single protein type.

The aim of the present study was to compare the effects of a high v. a normal amount of casein-protein-containing breakfast on plasma amino acid concentrations, appetite profile, such as ratings of hunger, satiety, fullness, and desire to eat, plasma glucose, and possibly related plasma hormone levels of insulin, glucagon-like peptide 1 (GLP-1), and ghrelin and subsequent energy intake. In order to determine the moment in time that may be sensitive to show a possible difference in food intake we first assessed appetite profile ratings and ‘satiety’ hormones for 4 h and in the subsequent experiment energy intake was measured at the determined moment in time.

Subjects and methods

Subjects

Thirty healthy male and female volunteers (BMI 22–30 kg/m\textsuperscript{2}; age 18–40 years) were recruited by advertisements in local newspapers and on notice boards at the university. They underwent a screening including medical history, measurement of body weight and height and cognitively restrained eating using a Dutch translation of the Three Factor Eating Questionnaire\textsuperscript{(23,24)}. Twenty-five subjects (eleven male, fourteen female) were selected on being in good health, non-smokers, non-vegetarian, not cognitively dietary restrained, not using medication apart from oral contraceptives and at most moderate alcohol users. Their mean age was 22 (SEM 1·8) years, and their body weight was 74·4 (SEM 1·8) kg (BMI 23·9 (SEM 0·3) kg/m\textsuperscript{2}). A written informed consent was obtained from these participants and the study protocol was approved by the Medical Ethical Committee of the Academic Hospital Maastricht.

Study design

A randomised, single-blind, within-subject experimental study was performed. All subjects came to the university on two occasions, separated by at least 1 week. On each test day subjects received a subject-specific standardised breakfast and appetite ratings and blood parameters were obtained for 4 h after breakfast.

After 2 months, when the sensitive moment in time was determined based on appetite profile ratings and concentrations of metabolites, subjects again came to the university on two occasions in a randomised, single-blind design, separated by at least 1 week. On each test day subjects received a subject-specific standardised breakfast and an \textit{ad libitum} lunch was offered at the previously determined sensitive moment in time.

Breakfast

Breakfast was offered as a custard with casein (Calcium Caseinate S; DMV International, Veghel, The Netherlands) as the single protein source, with either protein, carbohydrate and fat at 10, 55 and 35 En% (normal protein) or protein, carbohydrate and fat at 25, 55 and 20 En% (high protein). The breakfast contained 20 % of daily energy requirements, calculated as BMR, according to the Harris–Benedict equations, multiplied by an activity index of 1·75\textsuperscript{(25)}. The mean energy content of the breakfast was 2·52 (SEM 0·07) MJ and the provided breakfasts were completely finished.

The custards were produced by NIZO Food Research b.v. (Ede, The Netherlands) and had tapioca starch (Farinex VA50T; AVEBE, Veendam, The Netherlands and Perfect-amy 3108; AVEBE, Veendam, The Netherlands) and sunflower-seed oil (Reddy; NV Vandemoortele, Roosendaal, The Netherlands) respectively as the carbohydrate and fat sources and were citrus–vanilla (Citrus, Vanilla; J.B. de Lange, Belfeld, The Netherlands) flavoured. Extensive product development and use of a taste panel led to custards that did not differ significantly in colour, taste or viscosity. The amino acid composition of the custards is presented in Table 1.

Table 1. Amino acid content of the breakfasts given as a custard with either 10 % of energy from casein-protein or 25 % of energy from casein-protein (g amino acids/100 g custard)

<table>
<thead>
<tr>
<th>Amino acid</th>
<th>Casein 10 % of energy</th>
<th>Casein 25 % of energy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glutamic acid*</td>
<td>0·447</td>
<td>1·27</td>
</tr>
<tr>
<td>Aspartic acid†</td>
<td>0·150</td>
<td>0·355</td>
</tr>
<tr>
<td>Cysteine</td>
<td>0·009</td>
<td>0·021</td>
</tr>
<tr>
<td>Serine</td>
<td>0·120</td>
<td>0·283</td>
</tr>
<tr>
<td>Histidine</td>
<td>0·064</td>
<td>0·152</td>
</tr>
<tr>
<td>Glycine</td>
<td>0·040</td>
<td>0·094</td>
</tr>
<tr>
<td>Threonine</td>
<td>0·090</td>
<td>0·214</td>
</tr>
<tr>
<td>Arginine</td>
<td>0·092</td>
<td>0·218</td>
</tr>
<tr>
<td>Alanine</td>
<td>0·064</td>
<td>0·150</td>
</tr>
<tr>
<td>Tyrosine</td>
<td>0·120</td>
<td>0·283</td>
</tr>
<tr>
<td>Valine</td>
<td>0·141</td>
<td>0·333</td>
</tr>
<tr>
<td>Methionine</td>
<td>0·064</td>
<td>0·152</td>
</tr>
<tr>
<td>Isoleucine</td>
<td>0·112</td>
<td>0·265</td>
</tr>
<tr>
<td>Phenylalanine</td>
<td>0·110</td>
<td>0·259</td>
</tr>
<tr>
<td>Tryptophan</td>
<td>0·027</td>
<td>0·064</td>
</tr>
<tr>
<td>Leucine</td>
<td>0·204</td>
<td>0·483</td>
</tr>
<tr>
<td>Lysine</td>
<td>0·172</td>
<td>0·405</td>
</tr>
<tr>
<td>Proline</td>
<td>0·230</td>
<td>0·544</td>
</tr>
</tbody>
</table>

* Glutamic acid = glutamine + glutamate.
† Aspartic acid = asparagine.
completed just before breakfast and at 20, 40, 60, 80, 100, 120, 180 and 240 min after breakfast. Blood samples for urea and amino acid determination were obtained at −5 min and subsequently at the same time points as the appetite ratings; blood samples for the determination of glucose, insulin and ghrelin concentrations were obtained before and at 40, 60, 120 and 180 min after breakfast. Venous blood samples for the determination of GLP-1 concentration were obtained separately before, and at 30, 60, 90, 120 and 180 min after breakfast by means of a Venflon catheter placed in an antecubital vein\(^{38}\). Subjects were allowed to drink maximally two glasses of water spread over the morning.

In the second set of experiments, the protocol started after an overnight fast from 22.00 hours to 08.30 hours with scoring appetite ratings. Breakfast was offered \((t = 0 \text{ min})\) and completed within 20 min. Lunch was offered at the previously determined sensitive moment in time. Subjects were allowed to drink three glasses of water spread over the entire test period.

### Measurements

**Appetite profile.** To determine the appetite profile, hunger, fullness, satiety, and desire to eat were rated on 100 mm visual analogue scales (VAS), anchored with ‘not at all’ and ‘extremely’ during the test day. VAS are often used to measure subjective appetite sensations and the validity and reproducibility have been shown in several studies\(^{27,28}\).

Subjects were instructed to rate themselves by marking the scale at the point that was most appropriate to their feeling at that time. The distance from this point to the left end of the scale was measured in mm; changes from baseline (Δ) were calculated by subtracting the baseline score (−5 min) from the score at a certain time point.  

**Taste perception.** Taste perception profiles of the custards were assessed after the first and the last bite of the breakfast using 100 mm VAS, anchored with ‘not at all’ and ‘extremely’ on the aspects of pleasantness, sweetness, sourness, saltiness, bitterness, savouriness, crispiness, and creaminess.

**Energy intake.** Lunch was weighed before and after eating and energy intake was calculated by multiplying the difference of the weight of the lunch by the energy value of the lunch as determined by the product labels (11.4 kJ/g).

**Blood parameters.** Blood was distributed into EDTA tubes for glucose, insulin and ghrelin measurement. For GLP-1 measurement blood was collected in EDTA tubes with added dipeptidyl peptidase IV inhibitor. For amino acid and urea determination, blood was collected in lithium heparin tubes. Blood samples were centrifuged at 4°C for 10 min at 3000 rpm. Hydrochloric acid and phenylmethylsulfonyl fluoride were added to plasma for active ghrelin determination. For amino acid analysis, 250 µl plasma was deproteinised by mixing it with 20 mg dry sulfosalicylic acid. For analysis of urea, 200 µl plasma was deproteinised by mixing it with 20 µl of a TCA solution (500 g/l). All samples were stored at −80°C until further analysis.

Plasma glucose concentrations were determined using the hexokinase method (Glucose HK 125 kit; ABX Diagnostics, Montpellier, France). Insulin concentrations were measured by RIA (Linco Research Inc., St Charles, MO, USA). Plasma active ghrelin concentrations were measured by ELISA (Linco Research Inc.). Plasma active GLP-1 samples were analysed using ELISA (EGLP-35K; Linco Research Inc.). Plasma concentrations of amino acids were determined with the use of a fully automated HPLC (Pharmacia, Woerden, The Netherlands), after pre-column derivatisation with o-phthalaldehyde\(^{29}\). Plasma urea was analysed spectrophotometrically on a COBAS Mira S (Roche Diagnostica, Hoffman-La Roche, Basel, Switzerland).

### Statistical analysis

Data are presented as mean changes from baseline with their standard errors, unless otherwise indicated\(^{30}\). The area under the curve (AUC) of changes from baseline over time (4 h for appetite ratings, amino acid and urea concentrations; 3 h for glucose, insulin, GLP-1 and ghrelin concentrations) was calculated using the trapezoidal method. A repeated-measures ANOVA was carried out to determine possible differences between the high- and normal-protein breakfast. After the second set of experiments, a repeated-measures ANOVA was carried out to determine possible differences in energy intake between the breakfasts. A \(P\) value < 0.05 was regarded as statistically significant. Statistical procedures were performed using StatView 5.0 (SAS Institute Inc., USA; 1998).

### Results

**Appetite profile**

Baseline ratings for appetite scores were not different among treatments (Table 2). The AUC of fullness ratings was increased after the breakfast with 25% of energy from casein compared with the breakfast with 10% of energy from casein (8522 (SEM 872) v. 5459 (SEM 974) mm VAS; \(P<0.01\); Fig. 1). Fullness ratings also were increased after a breakfast with 25 En% casein compared with a breakfast with 10 En% casein at several moments in time including at 180 and 240 min after breakfast \((P<0.01\) and \(P<0.01\); Fig. 1). Satiety ratings were increased after the breakfast with 25% of energy from casein compared with the breakfast with 10% of energy from casein at 180 and 240 min \((P<0.05\) and \(P<0.05\); Fig. 1).

<table>
<thead>
<tr>
<th>Casein 10% of energy</th>
<th>Casein 25% of energy</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Mean</strong></td>
<td><strong>SEM</strong></td>
</tr>
<tr>
<td>Satiety (mm VAS)</td>
<td>25</td>
</tr>
<tr>
<td>Fullness (mm VAS)</td>
<td>24</td>
</tr>
<tr>
<td>Hunger (mm VAS)</td>
<td>62</td>
</tr>
<tr>
<td>Desire to eat (mm VAS)</td>
<td>66</td>
</tr>
<tr>
<td>Glucose (mmol/l)</td>
<td>5.16</td>
</tr>
<tr>
<td>Insulin (mU/l)</td>
<td>12.46</td>
</tr>
<tr>
<td>GLP-1 (pmol/l)</td>
<td>4.20</td>
</tr>
<tr>
<td>Ghrelin (pmol/l)</td>
<td>9.90</td>
</tr>
</tbody>
</table>

* Repeated-measures ANOVA.
Taste perception
Ratings of taste perception profiles and of pleasantness of taste of the custards were not different between the breakfast with 25% of energy from casein and the breakfast with 10% of energy from protein (Table 3).

Glucose
Baseline plasma glucose concentrations were not different among treatments (Table 2). The glucose response expressed as AUC was increased after the breakfast with 10% of energy from casein compared with the breakfast with 25% of energy from casein at 40 min after breakfast ($P<0.05$; Fig. 2).

Insulin
Baseline plasma insulin concentrations were not different among treatments (Table 2). Insulin concentration was increased after the breakfast with 10% of energy from casein compared with the breakfast with 25% of energy from casein at 40 min after breakfast ($P<0.05$; Fig. 2).

Glucagon-like peptide 1 and ghrelin
Baseline plasma GLP-1 and ghrelin concentrations were not different among treatments (Table 2). There were no significant differences in GLP-1 or ghrelin concentrations between a high- and a normal-casein breakfast (data not shown).

Amino acids and urea
Baseline plasma amino acid and urea concentrations were not different among treatments (Table 4). The AUC of the response of glutamate, asparagine, serine, glutamine, histidine, glycine, threonine, citrulline, arginine, alanine, taurine, α-aminobutyric acid, tyrosine, valine, methionine, isoleucine, phenylalanine, tryptophan, leucine, ornithine, lysine, branched-chain amino acids, total amino acids and urea are presented in Table 4; significant differences between treatments are indicated. Compared with the breakfast with 10% of energy from casein, almost all amino acids showed a prolonged elevation with a typical pattern after the breakfast with 25% of energy from casein. Plasma amino acid concentrations rose immediately after breakfast to peak values at 40 min after breakfast.
Breakfast. Then concentrations slightly decreased, with concentrations increasing again from 80 min onwards. The second peak levels were reached at 180 min after breakfast. To illustrate this phenomenon, Fig. 4 presents the plasma amino acid concentrations over time of the branched-chain amino acids and total amino acids. The prolonged elevated concentrations were shown with nearly all amino acids. Total amino acids and branched-chain amino acids as well as glutamate, asparagine, serine, glutamine, histidine, threonine, arginine, alanine, α-aminobutyric acid, tyrosine, valine, methionine, isoleucine, phenylalanine, tryptophan, leucine, ornithine and lysine concentrations were increased at 180 and 240 min after the breakfast with 25% of energy from casein compared with the breakfast with 10% of energy from casein (P<0·05).

The urea response expressed as AUC was increased after the breakfast with 25% of energy from casein compared with a breakfast with 10% of energy from casein (P<0·001; Table 4).

### Energy intake

Based on the results of appetite profile ratings and the concentrations of amino acids the ad libitum lunch was offered at 180 min after breakfast.

Energy intake at lunch was 3133 (SEM 226) and 3080 (SEM 229) kJ after the breakfasts with 10% and 25% of energy from protein, respectively (NS).

### Discussion

Ratings of satiety and fullness were higher after a subject-specific breakfast consisting of 20% of total daily energy requirements with casein at a concentration of 25% of energy from protein compared with the breakfast with 10% of energy from casein, particularly at 3 and 4 h after breakfast. Energy intake at lunch was not different after a high- or normal-casein breakfast. Sometimes it is suggested that protein-induced satiety is partly due to specific sensory effects (8). There is, however, hardly any evidence for this suggestion, especially not in the case of amounts of protein of about 30 g in combination with carbohydrate and fat in a meal. Clearly, most amino acids evoke taste-aversive responses because they have a bitter or sour taste(31). This is why we did not use pure amino acids but applied complete proteins. Nevertheless, to avoid any specific sensory effect, food technology was involved to optimise taste and hedonic value of the breakfasts. The custards were vanilla–lemon flavoured and after being tested by a professional taste panel of NIZO Food Research taste perception and hedonic values again were evaluated by the subjects (see Table 2) and were excluded from affecting appetite profile ratings differently.

The increased satiety after the breakfast with 25% of energy from casein compared with the breakfast with 10% of energy from casein coincided with prolonged elevated concentrations of amino acids. Since postprandial amino acid profiles are likely to reflect rates of digestion, absorption and metabolism, the prolonged elevated concentrations indicate a

### Table 3.

Taste perception profiles and hedonic values on 100 mm visual analogue scales of the breakfasts given as a custard with either 10% of energy from casein-protein or 25% of energy from casein-protein assessed in twenty-five subjects (men and women)*

(Mean values with their standard errors)

<table>
<thead>
<tr>
<th></th>
<th>Casein 10% of energy</th>
<th>Casein 25% of energy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pleasantness of taste</td>
<td>Mean 58 SEM 4</td>
<td>Mean 50 SEM 3</td>
</tr>
<tr>
<td>Sweetness</td>
<td>Mean 54 SEM 6</td>
<td>Mean 52 SEM 5</td>
</tr>
<tr>
<td>Saltiness</td>
<td>Mean 9 SEM 3</td>
<td>Mean 11 SEM 3</td>
</tr>
<tr>
<td>Bitterness</td>
<td>Mean 16 SEM 4</td>
<td>Mean 14 SEM 4</td>
</tr>
<tr>
<td>Sourness</td>
<td>Mean 16 SEM 4</td>
<td>Mean 11 SEM 3</td>
</tr>
<tr>
<td>Creaminess</td>
<td>Mean 56 SEM 6</td>
<td>Mean 53 SEM 5</td>
</tr>
<tr>
<td>Crispiness</td>
<td>Mean 2 SEM 1</td>
<td>Mean 3 SEM 1</td>
</tr>
<tr>
<td>Savouriness</td>
<td>Mean 15 SEM 4</td>
<td>Mean 19 SEM 4</td>
</tr>
</tbody>
</table>

* Repeated-measures ANOVA; no significant differences.
difference in protein kinetics between the two breakfasts.

Previously it has been shown that casein coagulates in the stomach which delays gastric emptying; therefore casein is considered as a ‘slow’ protein. The higher the casein concentration the slower the release of food into the duodenum\(^\text{[18,19,32]}\). This is reflected by the typical pattern of amino acid concentrations over time. The largest differences in amino acid concentrations between the breakfasts with 25% and 10% of energy from casein existed at 3 and 4 h after breakfast. These prolonged elevated concentrations may have contributed to the increased satiety ratings after the breakfast with 25% of energy from casein resulting in less pronounced changes in plasma amino acid pattern that is typical for a ‘slow’ protein casein breakfast. The high-casein breakfast revealed a pattern of amino acids caused by the delayed gastric emptying of casein.

Surprisingly, the insulin concentration was increased after the breakfast with 10% of energy from casein whereas the glucose response expressed as AUC also was significantly increased after the breakfast with 10% of energy from casein compared with the breakfast with 25% of energy from casein whereas the carbohydrate content of the two breakfasts was exactly the same. The slower release of food into the duodenum after the high-casein breakfast also delayed and diminished the rise of glucose and subsequently insulin concentrations in the circulation. Previously, insulin concentrations have been shown not to increase after consumption of a meal with ‘slow’ proteins in healthy young adults\(^\text{[199]}\).

Protein kinetics, reflected by changes in plasma amino acid concentrations, were different between the high- and normal-casein breakfast. The high-casein breakfast revealed a plasma amino acid pattern that is typical for a ‘slow’ protein and that was, besides glucose and insulin responses, also reflected by the changes in GLP-1 and ghrelin concentrations. The absence of significant differences in GLP-1 or ghrelin concentrations between the high- and normal-casein breakfasts may be the result of the delayed gastric emptying and thus retarded entrance of food in the intestine followed by a diminished physiological response of GLP-1 secretion and a less pronounced decrease in ghrelin concentration. In summary, the breakfast with 25% of energy from casein delayed gastric emptying more compared with the breakfast with 10% of energy from casein resulting in less pronounced changes in insulin, GLP-1 and ghrelin.

In the literature differences in ‘satiety’ hormone responses between the different macronutrients have been shown\(^\text{[12,13,33,34]}\). In a review by Cummings it is stated that protein intake does not affect ghrelin response particularly\(^\text{[35]}\). For instance, no differences in ghrelin concentrations after a high-protein (30 En% protein) compared with a normal-protein diet (10 En% protein) were observed, when the high- or normal-protein diet was given during three meals over 1 d\(^\text{[36]}\). Foster-Schubert, however, reported a stronger suppression of ghrelin by proteins compared with fat or carbohydrates\(^\text{[33]}\), with a test meal extremely high in protein with hardly any of the other macronutrients present. This makes comparisons with less extreme meals, such as in the present study a moderately high-protein meal that is representative for a relatively...
Effect on food intake. Previously, Diepvens et al. (15) reported that 15 mm VAS was not large enough to induce a significant reduction in energy intake. Apparently the difference in satiety ratings of 12–15 mm VAS in order to have a significant effect on subsequent energy intake is that previously has been shown to be related to diet-induced thermogenesis and increased satiety (9,42).

This is the first study that investigated acute differences in satiety between two concentrations of casein; previously the satiating properties of casein only have been compared with other protein types (11,14). The present study shows that a breakfast with 25 % of energy from casein is rated as being more satiating than a standard breakfast. The high-casein breakfast was rated as more satiating than the normal-casein breakfast the difference was not large enough to induce a reduction in energy intake.

Energy intake and that timing of major importance to observe significant effects on food intake (K Diepvens, J Steijns, P Zuurendonk and MS Westerterp-Plantenga, unpublished results). In the past there have been experiments that showed differences in subsequent energy intake between types of protein offered as a preload without significant differences in appetite ratings (11,37,38). In case subsequent energy intake is affected without pre-prandial indications of appetite profile ratings, it may well be that the combination of the digested food from the previous preload or meal with the new digested food in the gut may evoke uncomfortable feelings that stop further energy intake. Furthermore, differences in timing may explain different results; timing is essential in studying ad libitum energy intake after a preload or a meal as shown by Anderson et al. (37). In accordance with other studies (K Diepvens, J Steijns, P Zuurendonk and MS Westerterp-Plantenga, unpublished results), the present study shows that differences in appetite ratings thus need to be at least larger than 15 mm VAS in order to have a significant effect on subsequent energy intake. Although the high-casein breakfast was rated as more satiating than the normal-casein breakfast the difference was not large enough to induce a reduction in energy intake.

Urea concentrations were elevated more after the high-casein breakfast compared with the normal-protein breakfast with casein. The high urea concentrations reflect an excess of amino acids and a state of positive protein balance after the high-protein breakfast. Postprandial protein synthesis has high ATP costs (39) and when amino acids are given in excess of protein deposition, amino acid oxidation plays a major role in energy expenditure and protein oxidation (40,41) that previously has been shown to be related to diet-induced thermogenesis and increased satiety (9,42).

This is the first study that investigated acute differences in satiety between two concentrations of casein; previously the satiating properties of casein only have been compared with other protein types (11,14). The present study shows that a breakfast with 25 % of energy from casein is rated as being more satiating than a breakfast with 10 % of energy from casein at 3 and 4 h after breakfast, coinciding with prolonged elevated concentrations of plasma amino acids, but does not reduce subsequent energy intake.

Acknowledgements

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