The addition of monosodium glutamate and inosine monophosphate-5 to high-protein meals: effects on satiety, and energy and macronutrient intakes

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(Received 12 May 2008 – Revised 17 December 2008 – Accepted 26 January 2009 – First published online 9 March 2009)

In a fed and orally stimulated state, whether the addition of monosodium glutamate (MSG) (alone or in combination with inosine monophosphate-5 (IMP-5)) to a high-protein (HP) meal leads to early satiety and a difference in energy intake at a second course was investigated. Ten men and twelve women consumed, in random order, a first-course meal consisting of: (1) water (control); (2) a HP meal with 0·6 % MSG and 0·25 % IMP-5; (3) a HP meal with no additives; (4) a HP meal with MSG only; (5) a sham-fed meal 2 (oral-stimulation). Appetite perceptions, plasma concentrations of glucacon-like peptide 1 (GLP-1), glucose and insulin, and energy intake at a buffet (i.e. a second course) were measured before and after each condition. Changes in appetite, and in GLP-1, glucose and insulin, were similar for the three fed HP conditions and all were greater (post hoc all P<0·01) than the control and sham conditions. Energy intake was not different following the HP + MSG + IMP (1·86 (SEM 0·3) MJ) as compared with the HP + MSG-only (2·24 (SEM 0·28) MJ) condition (P=0·08), or for the HP + MSG + IMP compared with the HP no-additives condition (1·60 (SEM 0·29) MJ) (P=0·21). Following the HP + MSG-only condition, 0·64 (SEM 0·20) MJ more energy was consumed compared with the HP no-additives condition (P=0·005). We conclude that the addition of MSG to a HP meal does not influence perceptions of satiety and it may increase energy intake at a second course. Cephalic responses after the sham condition were of similar magnitude to the control and therefore just tasting food is not enough to influence appetite and energy intake.

Monosodium glutamate: Appetite: Energy intake: Satiety

Nutritionists treating individuals for weight loss continually look for new food ingredients that can further enhance the effectiveness of successful dietary treatments. Protein-rich foods have become popular for the dieting public because protein-rich meals: effects on satiety, and energy and macronutrient intakes on the body's protein requirements and stores at the time(11 – 17). In the present study, we found that subjects have also demonstrated that a preference for dietary protein (or liking of) is related, in part, to the way the body responds to MSG and inosine monophosphate (IMP) functions as a signal to regulate protein intake. A small body of scientific evidence is emerging to support this notion. Recently in thirty-six women and twenty-four men, we observed that the taste detection threshold of glutamate given orally as monosodium glutamate (MSG) and inosine monophosphate (IMP-5) is lower than that for MSG alone (P=0·005). This leads us to speculate that the taste detection threshold for MSG may relate to a preference for protein when given a choice of different foods, and ultimately, it may relate to habitual protein intake. That is, individuals with a low, rather than a high detection threshold, may more readily consume an adequate protein intake and consequently achieve a balanced nutritional status intake.

In addition to MSG and IMP-5 indicating a ‘liking’ or ‘preference’ for protein, Bellisle et al. showed that the addition of 0·6 % MSG to novel nutritionally valuable foods of medium palatability increased the spontaneous intake of them without affecting overall energy intake (due to a reduction in intake of non-MSG-enriched foods), in both a healthy elderly and diabetic elderly population(19 – 21). In thirty-two healthy individuals, Rodgers & Blundell(22) found that the addition of 20 % (dry weight) MSG to soup made it significantly more ‘pleasant’, ‘savory’ and ‘satisfying’ than a soup without MSG and, once again, there was no difference in subsequent energy intake despite there being a more rapid recovery of low concentrations (i.e. 0·1 to 0·4 %) also reported a greater liking and preference of protein-rich foods(10). Interestingly, Murphy showed that individuals who preferred higher concentrations of MSG in solution had a lower nutritional/protein status than those who preferred lower concentrations of MSG(18). This leads us to speculate that the taste detection threshold for MSG may relate to a preference for protein when given a choice of different foods, and ultimately, it may relate to habitual protein intake. That is, individuals with a low, rather than a high detection threshold, may more readily consume an adequate protein intake and consequently achieve a balanced nutritional status intake.

Abbreviations: GLP-1, glucagon-like peptide 1; HP, high-protein; IMP, inosine monophosphate; MSG, monosodium glutamate.

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hunger over 60 min after the 20 % MSG soup. Furthermore, Essed et al. (23) performed a single-blind randomised 16 weeks parallel study in elderly nursing home patients where they added 700 mg flavour and/or 300 mg MSG to the animal protein part of the cooked lunchtime meal. After 16 weeks, they found that enhancing the taste of the protein component of the meal did not lead to a higher overall energy intake at lunch, nor did it increase body weight. In this study, however, it was unclear whether there was a period of adjustment within the first day(s) of changing the diet in the acute effects of MSG on energy and protein intakes. This is of interest because in the flavour + MSG group, they observed a mean 1 kg reduction in body weight (0 kg change in the control group) which potentially could be related to the increase in total weight of protein consumed (i.e. protein may have promoted a greater thermogenesis and satiety, and therefore reduced day-long energy intake in the first few weeks of the 16-week period). In addition, the nursing home population was heterogeneous and this, too, may have masked small differences between the treatment and control group.

Oro-sensory stimulation by different food properties evokes cephalic responses that are involved in the regulation of food intake and body weight but our understanding of them remains unclear. Cephalic responses consist of vagal cholinergic activity initiated by the thought, sight, smell and taste of food. Experimentally they are evoked using modified sham feeding (i.e. tasting but not ingesting a meal). Sham feeding followed by enteric and gastric fat loads have been shown to bring about earlier responses in perceptions of satiety and in postprandial profiles of TAG, lipoprotein and insulin concentrations (24–26). Heath et al. also found that sham feeding before administering a 50 g fat load suppressed the ghrelin response (24). To our knowledge, no human studies have examined the effect of added MSG on cephalic responses evoked by the vagus nerve and whether they influence postprandial satiety and subsequent food intake.

The aim of the present study was to determine, in a fed and orally stimulated state (sham feeding), whether the addition of MSG to a HP meal leads to an early satiety and difference in energy intake at lunch, nor did it increase body weight. Following determination of the detection threshold for the taste of MSG, twenty-two subjects underwent a single-blind, randomised cross-over design. On five experimental sessions 1 week apart, subjects consumed in random order the following meals: (1) control (water); (2) a HP meal with 0-6 % MSG and 0-25 % IMP-5 added (HP + MSG + IMP); (3) a HP meal with no additives (HP no-additives); (4) a HP meal with MSG only added (HP + MSG); (5) sham-fed meal 2 (sham HP + MSG + IMP). The reason that condition 2 was included was because we had previously demonstrated that the addition of 0-25 % IMP-5 lowered the concentration of MSG detected in a soup solution from 0-33 % to 0-26 % (10). In the sham-fed condition, food was chewed and expectorated at the time when swallowing would normally occur to represent oral stimulation only. For all other conditions the water or food was completely ingested. Figure 1 depicts the study schedule for each experimental session; each session ran for approximately 1.5 h. Subjects were asked to come to the clinic at 08.30 hours after having

Subjects and methods

Subjects

Subjects were recruited from advertisements in a local newspaper and around the Maastricht University. Inclusion criteria were healthy men and women aged 18–65 years, non-smokers, on no medication (except the contraceptive pill), not known to be allergic to MSG or other foods, not diet-restrained (assessed using the Three Factor Eating Questionnaire (27)), low to moderate alcohol use (i.e. two or fewer standard drinks per d for no more than 5 d/week), a BMI between 20 and 30 kg/m² and weight stable (<3 % change over the 3 months before screening). Fat and fat-free mass was determined for each subject from the measurement of total body water using 2H dilution and the assumption that total body water occupies an approximate average of 73-2 % of the fat-free mass (28,29).

Twelve women and ten men were enrolled in the study that was approved by the Medical Ethics Committee of Maastricht University. All subjects gave written informed consent to participate.

Study design

As part of a separate study that has been published (10), all subjects underwent a series of ‘triangle taste tests’ to determine their sensitivity to the taste of MSG. In brief, the ‘triangle taste test’ methodology that we used involved the presentation of ten rows of triplicate cups with 8 ml soup (total of thirty cups). Within each triplicate, either one or two cups contained soup with added MSG. The remaining cups contained soup with no added MSG. Subjects were instructed to taste (but not swallow) half the volume of each soup cup and they were allowed to re-taste each sample if necessary. They were asked to ‘identify’ the soup with the ‘added MSG’ and ‘how certain they were about their choice’. The starting concentration was 0-4 % (w/w) MSG. If they could correctly detect the soup with added 0-4 % MSG for eight or more of the triplicates then the concentration was decreased (or if not, it was increased) by a step of 0-1 % until the detection threshold for the taste of MSG was determined (within the concentration range of 0-1–0-8 %). Between each cup within the ‘triangle taste test’, subjects were encouraged to rinse their mouth with water and chew on plain white bread (but not swallow) to get rid of previous tastes. Moreover, a 10–15 min break was taken between each ‘triangle taste test’ so that subjects could again rinse their mouth with water or chew on plain white bread to get rid of previous tastes.

Following determination of the detection threshold for the taste of MSG, twenty-two subjects underwent a single-blind, randomised cross-over design. On five experimental sessions 1 week apart, subjects consumed in random order the following meal conditions: (1) control (water); (2) a HP meal with 0-6 % MSG and 0-25 % IMP-5 added (HP + MSG + IMP); (3) a HP meal with no additives (HP no-additives); (4) a HP meal with MSG only added (HP + MSG); (5) sham-fed meal 2 (sham HP + MSG + IMP). The reason that condition 2 was included was because we had previously demonstrated that the addition of 0-25 % IMP-5 lowered the concentration of MSG detected in a soup solution from 0-33 % to 0-26 % (10). In the sham-fed condition, food was chewed and expectorated at the time when swallowing would normally occur to represent oral stimulation only. For all other conditions the water or food was completely ingested. Figure 1 depicts the study schedule for each experimental session; each session ran for approximately 1.5 h. Subjects were asked to come to the clinic at 08.30 hours after having
fasted for 12 h (water to prevent thirst, if necessary, was allowed overnight). Subjects were instructed to eat (or sham eat) the respective conditions over 20 min (these conditions represented a first course). Before and at several times after each first-course meal, 100 mm visual analogue scales were completed to assess pleasantness of taste of the meal and appetite. In addition, blood was taken via an intravenous catheter on the back of the hand to measure plasma markers of satiety (glucose, insulin and GLP-1). After 30 min later, subjects were allowed to eat whatever they wanted until they were ‘comfortably full/satisfied’ from a buffet containing six protein-rich and six carbohydrate-rich foods (i.e. the second course). At t = 0, 20 and 50 min, blood was taken for determination of glucose, insulin and glutamyl-like peptide 1. At t = 0, 20, 30, 40, 50 and 65 min, 100 mm visual analogue scales were completed. At 65 min, energy and macronutrient intakes eaten at the buffet meal were assessed.

First-course meal conditions

The control meal was water and contained no energy. Meal conditions 2–5 were isenergetic and the energy content for each subject was calculated using the Harris–Benedict equation to provide 20% of the daily energy requirements (2·2 (SEM 0·7) MJ) (30), the macronutrient composition was 30% energy from protein, 30% from fat and 40% from carbohydrate. All meals were of equal weight. Conditions 2 to 5 consisted of vegetable soup and one to two rolls filled with minced meat and salad (the number of rolls provided the remainder of the 20% of daily energy). The soup consisted of 400 ml soup stock (i.e. 250 g Vectra® vegetable bouillon (Naturis B.V., Harderwijk, The Netherlands) dissolved in 1 litre of water) to which 3 g Protifar® protein powder (Nutricia Clinical Care, Trowbridge, Wilts, UK), 10 g egg noodles and 20 g finely grated soup vegetables were added. The rolls each consisted of a 42 g wholemeal bread roll, 100 g prepared minced beef filling (recipe below), 5 g diet margarine, 20 g tomato, 10 g cucumber, 10 g lettuce and 7 g Protifar® protein powder (Nutricia Clinical Care). The minced beef filling was prepared in daily batches (i.e. 175 g fine minced beef, 20 g onion, 10 g olive oil and 650 g organic tomato paste). All food items used in the recipes, except when specified otherwise, were purchased from the local Albert Heintm supermarket. MSG and IMP-5 were purchased as salts from Ajinomoto Foods Germany GmbH (Hamburg, Germany) and the salts were dissolved into the soup and minced beef filling. Concentrations of 0·6% (w/w) MSG (i.e. the weight of MSG per weight of soup and roll) and 0·25 % (w/w) IMP-5 were selected because they are concentrations typically added to commercial food and are similar to levels of naturally occurring glutamate found in traditional dishes (31,32). No salt (sodium chloride) was added to the meals. The soup was served warm (55°C). Subjects were given 20 min to completely ingest each of the meals. For the sham-fed condition, subjects were asked to use 200 ml water provided to rinse their mouths and expectorate the chewed-up food into a bowl; they were asked not to drink the water. The weight of the meal expectorated was measured at the end of the session to determine how much may have been ingested.

Second-course buffet

The buffet consisted of six protein-rich and six carbohydrate-rich snack food items that were purchased from the local Albert Heintm supermarket. All twelve foods were presented simultaneously 30 min after subjects completed their first-course meal. Subjects were not informed that overall energy or macronutrient intakes consumed at the buffet were being investigated. The protein-rich food items were Albert Heintm fruit kwark, Albert Heintm natural European yoghurt, Albert Heintm unsalted peanuts, Frico® sliced hard cheese and Albert Heintm shoulder ham and portions of Slim Fast® protein-rich muesli bar. The carbohydrate-rich food items were Albert Heintm custard dessert with cream, Albert Heintm vanilla rice dessert, Euro Shopper® plain water crackers, Shutty® cheese-flavoured crackers, Lays® potato crisps and C1000® chocolate-chip cookies. All foods were presented in excess quantities in transparent plastic bowls and white plates of equal size. The protein content of the protein-rich food items that were offered at the buffet ranged from 7 to 37 g protein/100 g food whereas the carbohydrate-rich items contained between 3 and 11 g protein/100 g. Overall, the percentage of energy from protein presented for the protein-rich buffet was 45% whereas it was 9% for the carbohydrate-rich buffet items (P<0·001). Energy derived from fat for the protein-rich and carbohydrate-rich categories was not different (31 v. 35%; P=0·7). Subjects could eat whatever items they wanted from the buffet until they were ‘comfortably full’ and ‘comfortably satiated’. Food was removed after 15 min. No water was allowed over the buffet period.
Appetite ratings

Satiety, hunger, fullness, desire to eat and prospective future consumption were assessed using validated 100 mm visual analogue scales (33) completed immediately before and after the first-course meal, as well as at 10, 20 and 30 min after it (i.e. t = 0, 20, 30, 40 and 50 min; Fig. 1). The questions (for example, ‘How pleasant is the soup?’, ‘How hungry are you?’) were anchored at opposing ends of the 100 mm horizontal line by ‘not at all’ and ‘very’ and subjects were asked to mark the line to indicate how they felt at that moment.

Blood sampling

Blood was drawn immediately before and after, as well as at 50 min after, the start of the first course (i.e. t = 0, 20, 30, 40 and 50 min; Fig. 1) for the measurement of plasma glucose, insulin and glucagon-like peptide 1 (GLP-1) concentrations. Blood samples were collected in tubes containing EDTA to prevent clotting. Samples for GLP-1 (4 ml) were mixed with 40 μl of dipeptidyl peptidase IV (DPP-IV) inhibitor (Linco Research Inc., St Charles, MO, USA) to prevent degradation. Plasma was obtained by centrifugation (4°C; 3000 rpm; 10 min), immediately frozen in liquid N2 and stored at −80°C until analysed. Glucose concentrations were determined using the hexokinase method (Glucose HK 125 kit; ABX Diagnostics, Montpellier, France). Plasma concentrations of insulin were measured by RIA (Linco Research Inc., St Charles, MO, USA) to prevent degradation. Plasma active GLP-1 samples were analysed using enzyme-linked immunoradiometric assay (EGLP-35K; Linco Research Inc.).

Determination of energy and macronutrient intakes at the second-course buffet

Food presented in the buffet was weighed before presentation and after the experimental session. The buffet was presented immediately after the 50 min appetite questionnaire was marked. Energy and macronutrient intakes eaten at the buffet were assessed by difference. Following the sham and three HP fed conditions, the degree of adjustment in total energy consumed at the second course was calculated using the following formula which had the control condition (i.e. no energy consumed at the first course) as the reference condition: (total energy intake at first + second course) − energy intake at second course in control condition)/energy intake at second course in control condition × 100%.

Statistical analysis

All data are presented as mean values with their standard errors unless stated otherwise. Statistical analyses were made using SPSS for Windows (version 11.5; SPSS Inc., Chicago, IL, USA), and the criterion for significance (two-tailed) was set at P<0.05. Repeated-measures ANOVA with meal condition as the between-subject factor was used to determine differences between the conditions for pleasantness of taste and appetite ratings, energy and macronutrient intakes, and plasma concentrations of glucose, insulin and GLP-1. If an overall effect of condition was found, post hoc analyses were done using the least significant difference method for multiple comparisons (equivalent to no adjustments). Pearson correlation analysis was done to determine if there was a relationship between the taste detection threshold for MSG + 0.25 % IMP-5 and energy intake at the second course.

Study power

The study had a 100 % power (α = 0.05) with twenty-two subjects to detect an overall effect of condition on energy intake at the buffet. To detect, between the three fed HP conditions, a mean difference in energy intake at the second course of 650 (SD 961) kJ, the study had 80 % power (α = 0.05) when using the least significant difference method for multiple comparisons.

Results

The characteristics of the subjects who completed the study are shown in Table 1. After the first bite of each first-course meal, ‘pleasantness of taste’ ratings were not significantly different (57·9 (SEM 3·6) mm (control) v. 60·9 (SEM 3·1) mm (HP + MSG + IMP-5) v. 52·4 (SEM 3·5) mm (HP + no additives) v. 59·3 (SEM 3·2) mm (HP + MSG only) v. 64·9 (SEM 3·7) mm (sham HP + MSG + IMP-5); overall effect of condition, P = 0·07). For the sham-fed condition the weight of the meal expectorated by subjects was on average 88 (SEM 2·2) %.

### Table 1. Characteristics of the men and women who completed the study

<table>
<thead>
<tr>
<th></th>
<th>Men (n 10)</th>
<th>Women (n 12)</th>
<th>Sex effect (P)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>44 ± 19</td>
<td>32 ± 19</td>
<td>0·06</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>26·5 ± 3·8</td>
<td>23·7 ± 4·5</td>
<td>0·05</td>
</tr>
<tr>
<td>Total body water (kg)</td>
<td>46·1 ± 7·5</td>
<td>33·8 ± 6·6</td>
<td>&lt;0·001</td>
</tr>
<tr>
<td>Body fat mass (kg)</td>
<td>21·4 ± 8·9</td>
<td>21·0 ± 12·7</td>
<td>0·90</td>
</tr>
<tr>
<td>Body fat-free mass (kg)</td>
<td>63·2 ± 10·8</td>
<td>46·3 ± 8·9</td>
<td>&lt;0·001</td>
</tr>
<tr>
<td>Dietary restraint</td>
<td>6·7 ± 4·2</td>
<td>8·0 ± 5·2</td>
<td>0·40</td>
</tr>
<tr>
<td>Lowest concentration of MSG + IMP-5 tasted*</td>
<td>0·34 ± 0·22</td>
<td>0·29 ± 0·24</td>
<td>0·64</td>
</tr>
</tbody>
</table>

* Lowest concentration of MSG within the range of 0·1–0·8 % that could be correctly identified.
Changes in appetite ratings

Baseline values for hunger, fullness, satiety, desire to eat and prospective consumption were not significantly different between the meal conditions (all \( P > 0.1 \)). Appetite profiles for each of the five meal conditions are shown in Fig. 2. There was an overall effect of first-course meal condition for all appetite profiles (overall effect, \( P < 0.001 \)). Compared with the control, the HP meal with added MSG + IMP-5, the HP meal with no added MSG and the HP meal with only MSG all increased satiety and fullness, and decreased hunger, desire to eat and prospective consumption (all post hoc comparisons, \( P < 0.05 \)). There were no significant differences between the three HP fed conditions. For the sham-fed as compared with the control condition, there was no difference for each appetite profile. Sham feeding increased satiety and fullness, and reduced hunger, desire to eat and prospective consumption, but the changes were less than observed in each of the three HP fed conditions (all post hoc comparisons, \( P < 0.01 \)).

For fullness and satiety, there were significant time \( \times \) condition interactions at \( t = 20 \) min (\( P < 0.01 \)). For fullness, the sham condition produced the lowest post-meal ratings compared with all other conditions (all \( P < 0.04 \)). For satiety, the sham condition produced the lowest post-meal ratings compared with all conditions (post hoc \( P < 0.001 \)), except the control (post hoc \( P = 0.21 \)).

![Fig. 2.](https://doi.org/10.1017/S0007114509297212)

For hunger (a), fullness (b), satiety (c), desire to eat (d) and prospective consumption (e) ratings before and after the start (i.e. denoted by the arrow at time = 0 min) of the five first-course meal conditions that were consumed in random order over 20 min: (1) control (water, \( \circ \)); (2) fed high-protein (HP) meal with monosodium glutamate (MSG) and inosine monophosphate-5 (IMP-5) (\( \bullet \)); (3) fed HP meal without additives (\( \triangle \)); (4) fed HP meal with MSG only (\( O \)); (5) sham-fed HP meal with MSG + IMP-5 (\( A \)). Values are means for ten men and twelve women (\( n = 22 \)), with standard errors represented by vertical bars. *For hunger (\( P < 0.001 \)), fullness (\( P < 0.001 \)), satiety (\( P < 0.001 \)), desire to eat (\( P < 0.001 \)) and prospective consumption to eat (\( P < 0.001 \)) there was an overall effect of meal condition as assessed by repeated-measures ANOVA. Post hoc analyses (least significance difference adjustment for multiple comparisons) found for each appetite profile that the control and sham conditions were not significantly different from each other (\( P > 0.1 \)), the three fed conditions were not significantly different from each other (\( P > 0.1 \)) but the control and sham conditions were significantly different from the three fed conditions (\( P < 0.001 \)). † Fullness recorded at immediately after the first course for the sham condition was significantly different from all the conditions (all \( P < 0.05 \)). ‡ Satiety recorded at immediately after the first course for the sham condition was significantly different from all conditions (all \( P < 0.001 \)), except the control (post hoc \( P = 0.21 \)).
Changes in glucose, insulin and glucagon-like peptide 1

Glucose, insulin and GLP-1 are shown in Fig. 3. An overall effect of condition was observed on glucose, insulin and GLP-1 (all \( P < 0.001 \)). Glucose and insulin concentrations were raised more following the three HP fed conditions than either the sham or the control conditions (post hoc comparisons all \( P < 0.05 \)) but there was no difference in response between the three HP fed conditions with or without MSG and IMP-5. GLP-1 concentrations were raised more after the HP meal with added MSG + IMP-5 than the control (post hoc \( P = 0.013 \)), and all three HP fed conditions raised the concentrations more than the sham-fed condition (post hoc comparisons all \( P < 0.05 \)) but there was no difference in the GLP-1 responses between any of the HP conditions with or without MSG (post hoc all \( P > 0.12 \)). Glucose, insulin and GLP-1 concentrations did not differ between the sham and control conditions.

Energy and macronutrient intakes at the buffet

Energy and macronutrient intakes at the second-course buffet are shown in Table 2. An overall effect of condition was observed on absolute energy intake and on intakes of protein, carbohydrate and fat (\( P < 0.001 \)). Energy intake was not different following the HP + MSG + IMP as compared with the HP + MSG only condition (\( P = 0.08 \)), or for the HP + MSG + IMP compared with HP no additives condition (\( P = 0.21 \)). However, 0.64 (SEM 0.20) MJ more energy was consumed following the HP + MSG-only compared with the HP no-additives condition (\( P = 0.005 \)). Energy intake after each of the three HP fed conditions when compared with both the control and sham-fed conditions were lower (all post hoc comparisons \( P < 0.001 \)). However, the percentage energy consumed at the second course (relative to what was consumed after the control) was greatest for the HP with MSG-only condition (59 (SEM 8) %), followed by the HP with MSG + IMP-5 condition (46 (SEM 9) %), and then the HP no-MSG condition (39 (SEM 11) %) (post hoc all \( P < 0.01 \)). This amount was significantly greater after the HP + MSG condition than after the HP no-additives condition (post hoc \( P = 0.01 \)) but was not significantly different from what was consumed in the HP + MSG + IMP-5 condition (post hoc comparison, \( P = 0.09 \)). There was no difference between the HP + MSG + IMP and HP no-additive conditions (post hoc comparison, \( P = 0.34 \)). In the sham-fed condition they under-ate as compared with the control by 1 (SEM 6) % (because recovery of food in the sham condition was not complete, i.e. 88 (SEM 2.2) %, we assumed that the energy of this condition was 10 % of meal condition 2 for

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Fig. 3. Plasma glucose (a), insulin (b) and glucagon-like peptide 1 (GLP-1) (c) profiles before and after the start of the five first-course meal conditions that were consumed in random order at time \( = 0 \) min (denoted by the arrow): (1) control (water; \( W \)); (2) fed high-protein (HP) meal with monosodium glutamate (MSG) and inosine monophosphate-5 (IMP-5) (\( X \)); (3) fed HP meal without additives (\( I \)); (4) fed HP meal with MSG only (\( A \)); (5) sham-fed HP meal with MSG + IMP-5 (\( C \)). Values are means for ten men and twelve women (\( n = 22 \), with standard errors represented by vertical bars. * For the plasma profiles of glucose (\( P < 0.001 \)), insulin (\( P < 0.001 \)) and GLP-1 (\( P < 0.001 \)) there was an overall effect of meal condition. Post hoc analyses (least significant difference adjustment for multiple comparisons) done for each analyte found that the control and sham conditions were not significantly different from each other (\( P > 0.1 \)), the three fed conditions were not significantly different from each other (\( P > 0.1 \)) but the control and sham conditions were significantly different from the three fed conditions (\( P < 0.001 \)).
Table 2. Energy and macronutrient intakes of the ten men and twelve women at the second-course buffet, 30 min after the completion of the first-course meal conditions
(Mean values with their standard errors)

<table>
<thead>
<tr>
<th>First-course meal condition</th>
<th>Energy intake (MJ) Mean SEM</th>
<th>Protein (g) Mean SEM</th>
<th>Carbohydrate (g) Mean SEM</th>
<th>Fat (g) Mean SEM</th>
<th>% En protein Mean SEM</th>
<th>% En carbohydrate Mean SEM</th>
<th>% En fat Mean SEM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>3.03 a</td>
<td>0.22</td>
<td>36.2 a</td>
<td>3.1</td>
<td>86.8 a</td>
<td>7.6</td>
<td>26.2 a</td>
</tr>
<tr>
<td>HP + MSG + IMP-5</td>
<td>1.86 b,c</td>
<td>0.31</td>
<td>21.3 b</td>
<td>3.7</td>
<td>58.7 b,c</td>
<td>10.7</td>
<td>14.2 b,c</td>
</tr>
<tr>
<td>HP no-additives</td>
<td>2.24 a</td>
<td>0.29</td>
<td>20.6 b</td>
<td>4.0</td>
<td>69.9 a</td>
<td>9.5</td>
<td>12.0 b</td>
</tr>
<tr>
<td>Sham HP + MSG + IMP-5</td>
<td>2.77 a</td>
<td>0.29</td>
<td>32.5 a</td>
<td>3.8</td>
<td>80.9 a,d</td>
<td>10.2</td>
<td>24.0 a</td>
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<tr>
<td>Overall effect (P)*</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>0.33</td>
<td>0.013</td>
<td>0.15</td>
</tr>
</tbody>
</table>

% En, energy derived from the specific macronutrient expressed as a percentage of the total energy intake; HP, high-protein; MSG, monosodium glutamate; IMP-5, inosine monophosphate-5.

*a–d Mean values within a column with unlike superscript letters were significantly different (P<0.05; least significance difference method for multiple comparisons).

An overall effect of first-course meal condition was observed on energy intake, percentage of total energy consumed as fat, and on the weight of protein, carbohydrate and fat consumed. There was no significant effect on the percentage of total energy consumed as protein or carbohydrate at the second-course buffet.

Discussion

The main findings of the present study are that the addition of MSG (alone or in combination with IMP-5) to a HP meal does not influence the perceptions of appetite or satiety, or accelerate changes in the release of GLP-1, glucose or insulin, any significant increase in pleasantness with the addition of MSG. However, our data do not support the notion that MSG (alone or in combination with IMP-5) affects the taste detection threshold for MSG.

In the present study, all five test meals were rated as being similar in their 'pleasantness' upon the first taste. This is consistent with our previous study in which a 10 kJ soup without added MSG was not as 'pleasant' as a 10 kJ soup without added MSG. No other studies have reported that a 10 kJ soup preload enriched with MSG reduced absolute weight of protein, carbohydrate and fat intakes eaten at the buffet. However, when expressed as a percentage of total energy consumed, only fat intake was significantly greater in the conditions of HP with added MSG and the greatest pleasantness, although not statistically significant, was achieved with the addition of MSG and the greatest pleasantness, although not statistically significant, was achieved with the addition of MSG and HP with no additives. The lack of a statistical significance undoubtedly reflects a lack of study power (i.e. actual power is 58 %, statistically significant, was achieved with the addition of MSG and HP with no additives. The lack of a statistical significance undoubtedly reflects a lack of study power (i.e. actual power is 58 %).

The main findings of the present study are that the addition of MSG (alone or in combination with IMP-5) to a HP meal does not influence the perceptions of appetite or satiety, or accelerate changes in the release of GLP-1, glucose or insulin, any significant increase in pleasantness with the addition of MSG. However, our data do not support the notion that MSG (alone or in combination with IMP-5) affects the taste detection threshold for MSG.
the three fed HP conditions relative to the control and sham conditions most probably reflects increased carbohydrate and protein content, and increased energy. Although, there was no significance difference between the three fed HP conditions in these responses, it does appear that MSG alone and in combination with IMP-5 can increase insulin and GLP-1 concentrations marginally more than if no MSG is added to a HP meal. Further investigation may be warranted to clarify this observation. That the concentrations of insulin, glucose and GLP-1 barely increased after the control and sham conditions reflects the regulation of these analytes by nutrients. No other studies to our knowledge have examined the effect of MSG on GLP-1 or other satiety hormones. Chevassus et al. (34) did, however, examine MSG’s effect on insulin secretion. They reported that 10 g encapsulated MSG, when administered simultaneously with an oral glucose load, increased glucose-induced insulin secretion in a concentration-dependent manner in eighteen healthy volunteers. Plasma glucose was not lowered in their study, but a study in rats showed that as insulin concentrations were increased by added MSG to the diet, glucose tolerance improved (35).

Given that (i) protein is the most satiating macronutrient, (ii) Rogers & Blundell(22) observed a reduction in hunger and desire to eat following the consumption of MSG-enriched soup (albeit to an equal extent as a soup without MSG) and (iii) protein intake can be increased when MSG is added (albeit not significantly in statistical terms) (22), we formulated our hypothesis that the consumption of a HP meal enriched with MSG alone or MSG + IMP would lead to an earlier satiety and thereby reduce energy intake at a subsequent second course. Energy intake, however, was increased (on average) by 0.644 MJ following the HP condition with added MSG as compared with no added MSG. Given that we also observed a significant negative association between energy intake at the buffet meal and the taste detection threshold of individuals for the taste of MSG + 0.25 % IMP-5, our findings may raise some concern for users of MSG, particularly those who are sensitive to its taste. On the other hand, the observed association was weak. Moreover, it is interesting that energy intake following the HP + MSG + IMP condition was not significantly different from the HP no-additives condition. The present study was sufficiently powered (80 % power, α = 0.05) to detect differences between the three HP fed conditions to the magnitude of 0.65 (SEM 0.96) MJ. If the 256 (SEM 938) MJ difference between the HP + MSG + IMP condition and the HP no-additives condition were statistically significant, it would have detected it despite the large within-subject variation in responses. Therefore, further research, in human subjects, is necessary to confirm the effects of MSG on energy intake, and perhaps using MSG in combination with IMP-5 can prevent MSG-induced over-consumption if it is a real phenomenon in humans. Ideally, a larger population with equal number of tasters at both ends of the MSG taste sensitivity range would clarify these results.

At the second-course buffet, we anticipated that subjects might reduce their overall energy intake by continuing to eat foods that exuded the similar ‘meaty’ and/or ‘savoury’ taste as provided in the first course (i.e. buffet included ham, hard cheese and unsalted peanuts) until they reached ‘sensory-specific satiety’. The buffet foods that provided those sensory properties were all higher in protein than the other foods that had a higher carbohydrate content and which provided more of a ‘sweet’ and/or ‘creamy/fatty’ taste (i.e. sweet cookies, rice cream, cream-topped custard, potato crisps). We did also include, however, some higher-protein items with a ‘sweet’ and/or ‘creamy/fatty’ taste (i.e. natural yoghurt, fruit yoghurt and a protein snack bar). Of course, what might have happened was that the subjects had reached ‘sensory-specific satiety’ at the end of the first course because it was a relatively large energy load (i.e. 2.2 MJ on average). Therefore, at the next course, they may have eaten to satisfy their desire for another taste sensation. In addition, the 30 min break between the first and second courses may have been too long and thereby any desire to continue eating foods with similar sensory properties was lost. Furthermore, our buffet design may have limited our ability to see any effect because we closely matched the overall fat content of the six foods that provided the ‘meaty’/’savoury’ taste to those that provided the ‘sweet’ or ‘creamy/fatty’ taste. It is also possible that some individuals ate a bit of everything, and because the food items were similar in their fat content, the increase in the percentage energy derived from fat that we observed was the cause of the increased energy intake between the HP conditions with and without MSG. The addition of MSG to the diet of rats (but not when ingested in the absence of energy) has also been shown to increase the thermic effect of feeding and promote a rapid switch in substrate utilisation (30) over the first 30 min after the test meal. Although we did not measure these indices, we believe that they would have been underlying mechanisms of an early satiety and reduced energy intake, had we found that to occur in the present study.

The sham-fed HP condition with added MSG and IMP-5 was included in the study design to discriminate the pre-ingestive/cephalic effects of the HP meal with MSG (alone or in combination with IMP-5) (i.e. those evoked by the thought, sight, smell and taste only) from the post-ingestive effects. The results revealed that thinking, seeing, smelling and tasting the meal containing MSG + IMP-5 had no significantly greater impact on post-ingestive and post-absorptive appetite and metabolism than did drinking an equal volume of water (i.e. mechanical effect).

We conclude that the addition of MSG to a HP meal does not influence perceptions of satiety but it may increase energy intake at a second course. Cephalic responses observed after the sham-fed HP + MSG + IMP-5 condition were of similar magnitude as the control. This provides evidence that just tasting food is not enough to influence satiety and energy intake. GLP-1 was not involved in cephalic or post-prandial responses observed in the present study. Further research, in human subjects, is warranted to clarify the effect of MSG on energy intake and feeding behaviour. A study in a larger population with equal numbers of ‘highly sensitive’ and ‘not sensitive at all’ tasters of MSG may provide some insight into mechanisms that lead some individuals (but not all) to over-consume energy when food is enriched with MSG.

Acknowledgements
We would like to thank Loek Wouters for analysing the 3H-labelled urine samples for determination of body composition, and Jos Stegen and Wendy Sluijsman for analysing...
the blood samples for glucose and insulin. N. D. L.-M. designed the experiment, collected the data, analysed the data and wrote the manuscript. A. J. P. G. S. read the manuscript and contributed to the discussion. M. S. W.-P. helped to design the experiment, analyse the data and write the manuscript, and supervised the project. The present study was supported by project funding (RTD line 5, Obesity and Food Technology) from The European DiOGenes program 2002–2006 and Top Institute Food & Nutrition, The Netherlands.

None of the authors had any financial or personal interest in any company or organisation sponsoring the research.

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