The nitrogen-sparing effect of methionine sulfoxide and some other sulphur-containing amino acids

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1. The nitrogen-sparing effect of methionine, methionine sulfoxide, homocystine, cystine and choline was studied in rats by determining daily N excretions during 5 d after changing from a high-protein diet to a protein-free diet. L-glutamic acid was used as a negative control.

2. L-, D- and DL-methionine were equally active in sparing N. L-methionine sulfoxide, DL-homocystine and L-cystine were as active as L-methionine. D-methionine sulfoxide was slightly less active than L-methionine sulfoxide. Choline hydrogen tartrate was not different from the negative control.

3. It is concluded that in short-term experiments cystine is the key substance in the N-sparing effect. The question of whether methionine sulfoxide is biologically available as a methionine source for rats and other experimental animals is a matter of some controversy (Gjøen & Njaa, 1977). In the present study we have tested whether the sulfoxide has a nitrogen-sparing effect (Dreyer, 1970) similar to that reported for methionine in experiments with rats, dogs, pigs and chicks (Nielsen et al. 1939; Miller, 1944; Allison et al. 1947; Brush et al. 1947; Muramatsu & Okumura, 1980).

The N-sparing effect is observed when the experimental animals are given methionine in small amounts in an otherwise protein-free diet. The urinary N excretion is then reduced as compared to that seen without the methionine addition. The effect is most pronounced early after changing from a protein-containing to a protein-free diet (Miller, 1944; Dreyer, 1970; Lubaszewska et al. 1973).

We have studied the urinary N excretion during 5 d after changing rats from a stock diet containing approximately 240 g protein/kg to a protein-free diet to which the amino acids under study were added. The amino acids tested were L-, D- and DL-methionine, L- and D-methionine sulfoxide, L-cystine, DL-homocystine and L-glutamic acid; choline hydrogen tartrate was also tested.

METHODS

Analytical procedures

Urine was collected as described by Njaa (1963). N in the daily urine samples was determined essentially as described by Crooke & Simpson (1971). In Expt 6 urea was determined as described by Potts (1967) omitting the extraction procedure. The results were analysed by analyses of variance and comparisons between treatments were done after partitioning the treatment sum of squares into single degrees of freedom as described by Snedecor (1956).

Experimental procedures

Six experiments were performed. Each experiment comprised three or four experimental groups of five or six rats, all experiments, except Expt 1, were run with litter mate controls. The rats used were males which had been given the stock diet for at least 2 weeks before they were given the experimental diets. The latter consisted of a protein-free diet to which were added the amino acids tested. The composition of the protein-free diet was (g/kg diet): partially-dextrinized potato starch 680, sucrose 200, arachis oil 60, mineral mixture 40,
vitamin mixture 10, cellulose powder 10. The dietary levels of minerals and vitamins were as described by Gjøen & Njaa (1977). The amount of N added to the diet from the amino acids was 300 mg/kg diet which is equivalent to 3.2 g methionine/kg diet. Each rat was given 10 g food daily and a record was kept of food intake. The urine was collected daily from each rat and analysed separately for N. The experiments lasted for 5 d and urine was collected from the first day.

The design of the experiments is shown in Table 1. L-glutamic acid was used as a negative control, and L-methionine as a positive control in all experiments except in Expt 2 where the positive control was omitted.

Expt 1. L-methionine, L-methionine sulphoxide and L-methionine sulphoxide (225 mg N/kg diet) together with L-cystine (75 mg N/kg diet) were compared for their N-sparing effect.

Expt 2. L-methionine sulphoxide (300 mg N/kg diet), and L-methionine sulphoxide (150 mg N/kg diet) together with L-glutamic acid (150 mg N/kg diet) were compared.

Expt 3. L-, D- and DL-methionine were compared.

Expt 4. L-methionine, L-methionine sulphoxide and D-methionine sulphoxide were compared.

Expt 5. L-methionine, DL-homocystine, and DL-homocystine together with choline were compared. The choline added could theoretically methylate 33% of the homocystine, bearing in mind that only one of its methyl groups is transferable. Thus, in the group receiving choline the level of added N was 400 mg/kg diet.

Expt 6. L-methionine, L-cystine and choline were compared at equivalent N levels.

RESULTS
The results obtained are summarized in Table 2. They refer to the entire 5 d period. Similar tables for the separate days could have been presented, but as the results were rather clear-cut the daily excretions are only occasionally referred to.

In all experiments in which L-methionine and L-glutamic acid were compared the urinary N excretion was significantly less with the former. In most instances the difference was already highly significant on the second day of the experiment; in Expt 4, in which the individual variations were great, the difference was highly significant from day 3.
Table 2. Urinary nitrogen (U, mg/rat per d), mean daily weight loss (ΔW, g/rat) and mean daily food intake (Fd, g/rat per d)

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<th>Expt no.</th>
<th>Mean wt at start of experiment (g)</th>
<th>L-M</th>
<th>D-M</th>
<th>DL-M</th>
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<th>L-S + L-C</th>
<th>L-S + Glu</th>
<th>D-S</th>
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L-M, L-methionine; D-M, D-methionine; DL-M, DL-methionine; L-S, L-methionine sulphoxide; D-S, D-methionine sulphoxide; L-C, L-cystine; DL-HC, DL-homocystine; Ch, choline hydrogen tartrate; Glu, L-glutamic acid.

a, b, c. Values with different superscript letters were significantly different (P < 0.01 or 0.001).
Fig. 1. Expt 6. Relationship between the daily excretions of total urinary nitrogen (TN) and urea-N (UN), and between values calculated from Lubaszewska et al. (1973). (●), L-methionine; (○), L-cystine; (▲), choline hydrogen tartrate; (△), L-glutamic acid. UN = 0·901 TN - 10·61, r = 0·992, n = 20. (■), Lubaszewska et al. (1973). UN = 0·739 TN - 9·68, r = 0·893, n = 59. (The circled value was not used in the calculation.)

The results obtained with L-methionine sulphoxide showed no significant difference from those obtained with methionine (Expt 1). This was also the situation when 250 mg methionine sulphoxide-S/g was replaced by L-cystine-S. In Expt 4 there was a tendency for L-methionine sulphoxide to be less effective than L-methionine, and for D-methionine sulphoxide to be less effective than L-methionine sulphoxide. The differences were not significant when taken over the entire 5 d period. However, for the last 2 d the difference between D-methionine sulphoxide on the one hand and L-methionine sulphoxide and L-methionine on the other was significant (P < 0·05) whereas the difference between the latter two was not. In Expt 2, L-methionine sulphoxide was tested at two levels. At the level equivalent to 3·2 g methionine/kg diet the difference between the urinary N excretions on the L-glutamic acid diet and that on the sulphoxide diet was 32 mg N/rat per d. When the sulphoxide level was halved the difference was also approximately halved to 17 mg N/rat per d. In Expt 5 the effect of DL-homocystine was not significantly different from that of L-methionine and the addition of choline to the homocystine was without any additional effect.

In Expt 6 the effect of L-cystine was not significantly different from that of L-methionine, and the N excretion on the diet with added choline was not significantly different from that found on the diet with added L-glutamic acid. The urea-N content of the urine samples paralleled closely the total N content. The close linear correlation between the two variables is shown in Fig. 1. The mean weight losses and the mean food intakes showed significant differences in Expts 4–6, but not in Expts 1–3 (Table 2).
Our main concern in these experiments was whether the N-sparing effect of L-methionine was retained by L-methionine sulphoxide. This could be answered in the affirmative (Expts 1 and 4). There were, however, indications that D-methionine sulphoxide was less effective (Expt 4). The effect of L-methionine sulphoxide was dose related (Expt 2). As L-cystine and DL-homocysteine were also as effective as L-methionine (Expts 2, 5 and 6) the N-sparing effect seemed not to be due to the furnishing of labile methyl groups. Mudd (1980) pointed out that methylneogenesis is so controlled that labile methyl groups are made available to meet the needs for methionine. Moreover, choline showed no effect (Expts 5 and 6). The fact that L-, D- and DL-methionine as well as DL-homocystine were equally effective indicates that the configuration around the α-carbon was of little importance (Expts 3, 5 and 6). The indication that D-methionine sulphoxide was less effective than L-methionine sulphoxide could probably be interpreted to show some importance of the configuration around the sulphur atom. In an earlier communication it was shown in N-balance experiments that D-methionine sulphoxide was less effective as a substitute for L-methionine than was L-methionine sulphoxide (Njaa, 1962).

The finding that L-cystine had the same N-sparing effect as L-methionine would indicate that cystine is the key substance in this connexion. Cystine is outside the reversible pathways of methione metabolism (Mudd, 1980). Assuming that a system exists for the reduction of methionine sulphoxide to methionine, cystine can be formed from all the substances found to show a N-sparing effect but it cannot serve as a source for any of them. It may be relevant in this connexion that Finkelstein et al. (1980) suggested that trans-sulphuration, rather than methionine conservation, is a major metabolic response in hepatic regeneration. Also, the results with cystine obtained with dogs (Nielsen et al. 1939; Miller, 1944) and with rats (Brush et al. 1947) indicate that methionine itself is not necessary to obtain the N-sparing effect.

During protein depletion the urine produced contains progressively less urea-N, both in absolute amounts and relative to total N (Dreyer, 1970; Lubaszewska et al. 1973). Our results confirm this. In Fig. 1 it is shown that we found a linear correlation between urea-N and total urinary N with a positive regression coefficient and a negative constant term. Thus, the regression equation is consistent with an absolute and a relative decrease in urea-N as total N excretion decreases. Fig. 1 also includes the regression line between urea-N and total urinary N calculated from the results of Lubaszewska et al. (1973). For the purpose of comparison with our results the observations were calculated as mg N/rat per d assuming a body-weight of the rats of 280 g. The constant terms in the two equations were very similar. They may be interpreted to represent a relatively constant part of the urinary N excretion which is independent of the total loss. Together with a minimum obligatory loss of urea-N it would constitute the constant endogenous N loss. The fact that the regression coefficients were different may probably be a consequence of different experimental techniques. Lubaszewska et al. (1973) fasted their rats for 12 h before the start of the experiments whereas we took our rats directly from the stock diet to the protein-free diet. They also collected urine for five 3-d periods whereas we analysed the daily urines. Similar differences in technique are also obvious between our experiments and those of Dreyer (1970).

In our experiments we did not always find reduced body-weight losses in the groups in which the N-sparing effect was observed. This is in contrast to the findings of Yoshida & Moritoki (1974) and to later publications from the same group. However, our feeding periods were so short that the measurement of weight loss would be subject to great error. In Expts 4–6, in which significant differences were observed, the differences were in the expected direction and were accompanied by parallel differences between the food intakes.

The N-sparing effect of methionine has tentatively been explained by assuming that it is needed for the production of an essential metabolite, and that the body raids its own
tissue if this amino acid is not supplied by the diet (Hoover et al. 1949). The present study was aimed mainly at establishing whether methionine sulfoxide retained the N-sparing effect of methionine. The experiments indicate that it does. Moreover, as it seems probable that cystine is the key substance in short-term experiments, reduction of methionine sulfoxide to methionine is indicated.

REFERENCES

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