AMMONIA EXCRETION AND RENAL ENZYMATIC ADAPTATION IN HUMAN SUBJECTS, AS DISCLOSED BY ADMINISTRATION OF PRECURSOR AMINO ACIDS\textsuperscript{1,2}

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Urinary ammonia is produced in renal tubular cells by the deamination of glutamine and asparagine and the deamination of a variety of amino acids (1–5). Many ammonia-producing enzymes have been identified in mammalian kidney and both their presence and activity appear to vary widely with species (5–14). In the dog, 60 per cent of the urinary ammonia has been attributed to the deamination of serum glutamine, the remaining 40 per cent apparently originating from amino acid oxidase activity (1). In the human kidney, glutaminase activity is only 0.1 to 0.01 that present in dogs (15). Renal \textit{l}-amino acid oxidase activity is low or absent in most species save the rat (8) and renal \textit{d}-amino acid oxidase varies widely in different species, the activity in the cat being 20 times that in the guinea pig (14).

Recently it has been shown in rats that strong acid loads increase renal ammonia production consequent to adaptation of glutaminase (16–19), glycine oxidase and \textit{l}-amino acid oxidase (16). Increased renal ammonia production and excretion also follows the administration of precursor amino acids; in dogs the administration of a \textit{dl}-alanine, \textit{l}-leucine, glycine, glutamine and \textit{dl}-aspartic acid results in a prompt increase in urinary ammonia excretion (1–4).

It is not known precisely which ammonia-producing enzymes are present in the human kidney and which of these undergo adaptive change during the administration of strong acid loads. The purposes of this study were threefold: to determine which amino acids are precursors of urinary ammonia, thereby furnishing evidence for the presence of various ammonia-producing enzyme systems in the human kidney, to ascertain which of these enzymes show adaptive increase in activity and finally to help define the relationship between ammonia production and ammonia transport by observing the changes in urinary acid excretion during the administration of precursor amino acids.

PROCEDURE

Five normal male subjects, housed on the metabolic ward and maintained on a diet of constant composition, were given \textit{NH}_{4}\textit{Cl} loads continuously for from one to five months. One hundred twenty-five studies, each lasting 8 to 10 hours, were performed. To identify the amino acid precursors of urinary ammonia, the subjects were kept on 15 Gm. of ammonium chloride until ammonia excretion was constant. Single amino acids (230 to 400 mM) were then given orally over a 60 to 90 minute period and changes in urinary ammonia, pH and titratable acid (TA) were followed hourly for six to eight hours. These values then were compared to those obtained during a control period of equal duration, performed during the same hours of the day in order to obviate changes related to diurnal variation.

To determine which renal enzyme systems adapt to strong acid loads, subjects were studied under three circumstances: standard diet (SD) plus 5 Gm. of \textit{NH}_{4}\textit{Cl} per day; SD plus 10 Gm. \textit{NH}_{4}\textit{Cl} per day; and finally, SD plus 15 Gm. \textit{NH}_{4}\textit{Cl} per day. Each level of \textit{NH}_{4}\textit{Cl} administration was continued from 10 to at least 30 days. When a steady state was attained, as evidenced by a constant urinary ammonia excretion during a fixed six to eight hour period each day (9:00 a.m. to 5:00 p.m.), equimolar amounts (300 mM) of different amino acids were given on different days.\textsuperscript{8} Experiments were performed at no less than three day intervals to allow adequate time for return to a steady state. The amino acids, dissolved in 500 to 1,000 ml. of water, were administered orally over a 60 to 90 minute period. Urine specimens were collected hourly under mineral oil in bottles containing phenyl mercuric nitrate as a preservative.

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\textsuperscript{2} Presented at the meeting of the American Federation for Clinical Research in Atlantic City, May, 1955.

\textsuperscript{8} On the days of the amino acid loading and control experiments, \textit{NH}_{4}\textit{Cl} was not administered until after the study was completed. The subjects therefore did not receive \textit{NH}_{4}\textit{Cl} for 14 hours prior to each experiment.
**TABLE I**

*Mammalian renal enzyme systems capable of ammonia production*

<table>
<thead>
<tr>
<th>ENZYME</th>
<th>SUBSTRATE</th>
<th>PRODUCTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Glutaminase*(5,12,15)*</td>
<td>Glutamine</td>
<td>Glutamic acid + ammonia</td>
</tr>
<tr>
<td>2. Asparaginase*</td>
<td>Asparagine</td>
<td>Aspartic acid + ammonia</td>
</tr>
<tr>
<td>3. D-Amino acid oxidase*</td>
<td>Most D-amino acids</td>
<td>Corresponding α-keto acid + ammonia</td>
</tr>
<tr>
<td>4. L-amino acid oxidase*</td>
<td>Most L-amino acids</td>
<td>Corresponding α-keto acid + ammonia</td>
</tr>
<tr>
<td>5. Glycine oxidase*</td>
<td>Glycine</td>
<td>Glyoxylic acid + ammonia</td>
</tr>
<tr>
<td>6. Glutamic acid dehydrogenase</td>
<td>L-glutamic acid</td>
<td>α-ketoglutaric acid + ammonia</td>
</tr>
<tr>
<td>7. Transaminase*</td>
<td>a.L-aspartic + α-ketoglutaric acid</td>
<td>Oxaloacetic acid + glutamic acid</td>
</tr>
<tr>
<td></td>
<td>b.L-alanine + α-ketoglutaric acid</td>
<td>Pyruvic acid + glutamic acid</td>
</tr>
<tr>
<td>8. Proline oxidase*</td>
<td>Proline</td>
<td>L-glutamic acid</td>
</tr>
</tbody>
</table>

* Neither the transaminases nor proline oxidase directly produce ammonia but if coupled with glutamic dehydrogenase, a net production of ammonia occurs. The superscript indicates a reference number.

Immediately after voiding, ammonia, TA and pH were determined. In 42 experiments titratable acid minus bicarbonate (TA–HCO₃⁻) was measured. The values obtained on experimental days, when amino acid was given, at each level of NH₄Cl administration, were compared to the values obtained on control days at the same level of NH₄Cl administration. The net change (experiment minus control) in ammonia excretion, attributable to the administration of each amino acid at each level of NH₄Cl administration, was thus obtained. The net change in urinary ammonia [or ammonia plus (TA–HCO₃⁻)] after a specific amino acid, while the subject was receiving 5

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**Fig. 1. Effect of Administration of Glycine on Urinary Ammonia Excretion, pH and Titratable Acid**
Gm. NH₃Cl per day, was then compared to the net change that occurred when the subject received 10 and then 15 Gm. NH₃Cl per day.

**Methods.** Urinary ammonia was determined in duplicate by a modification of the microdiffusion method of Conway (20). Urine pH was measured with a Beckman pH meter using external glass electrodes and corrected to 37.5° in accord with the findings of Wesson for human urine (21). Urine was titrated to pH 7.4 with 0.1 N NaOH to determine titratable acid. TA-HCO₃ was measured by the method of Dawson, Dempsey, Bartter, Leaf and Albright (22). Serum CO₂ content was determined manometrically (23). Serum chloride was measured by the method described by Peters and Van Slyke (23); serum sodium and potassium concentrations were determined on the flame photometer (24).

### RESULTS

1. **Identification of ammonia-producing enzyme systems in human kidney by administration of precursor amino acids**

The enzymes producing ammonia from amino acids which have been identified *in vitro* in mammalian kidney are listed in Table I. A typical experiment showing an augmented ammonia excretion after administration of glycine to a subject stabilized on 15 Gm. NH₃Cl daily is presented diagramatically in Figure 1. The prompt rise in urinary ammonia excretion lasting four to six hours suggests that glycine oxidase is present in human kidney. The results after the administration of 12 different amino acids suggest that all

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**TABLE I**

Protocols of typical amino acid loading and control studies.*

<table>
<thead>
<tr>
<th>PREPARATION</th>
<th>AMINO ACID</th>
<th>CONTROL</th>
<th>EXPERIMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Standard diet + 5 gm. NH₄Cl day</td>
<td>Glycine</td>
<td>27.9</td>
<td>0-60</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Standard diet + 10 gm. NH₄Cl q.d.</td>
<td>Glycine</td>
<td>24.0</td>
<td>0-60</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Standard diet + 15 gm. NH₄Cl q.d.</td>
<td>Glycine</td>
<td>20.7</td>
<td>0-60</td>
</tr>
</tbody>
</table>

* TV refers to total urine volume.
2. Adaptation of ammonia-producing enzymes in the human kidney during the chronic administration of ammonium chloride loads

The protocols from a typical series of experiments performed to determine whether enzyme adaptation occurred are shown in Table II. Net ammonia excretion (experiment minus control) progressively increased from 4.2 to 21.6 to 41 mEq. following the administration of 400 mM of glycine to a subject chronically maintained on 5, then 10 and finally 15 Gm. of NH₄Cl per day. Data from three subjects showing the increasing magnitude of net ammonia excretion following the administration of 400 mM of glycine at different levels of NH₄Cl intake are shown in Figure 2.

Typical examples of the results of experiments in two other subjects performed in a similar manner to determine whether or not adaptation of glutaminase, asparaginase, d-amino acid oxidase and l-amino acid oxidase occurred are shown in Figures 3 and 4.

The mean increase in net ammonia excretion for the five subjects following the administration of this group of precursor amino acids (glycine,
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FIG. 4. EFFECT OF DIFFERENT AMINO ACIDS AND AMINO ACID AMIDES ON NET URINARY AMMONIA EXCRETION FOLLOWING STEPWISE INCREASE IN THE MAGNITUDE OF THE CHRONIC AMMONIUM CHLORIDE LOAD.

The numbers in the columns refer to the net increase in urinary ammonia after the substrate load. Asparagine, glutamine, l-alanine, d-alanine and l-leucine) increased significantly (p < 0.01) from 4.8 to 12.2 to 20.6 mEq. as the chronic maintenance dose of NH₄Cl was increased from 5 to 10 to 15 Gm. a day. By contrast, the administration of glutamate, aspartate or proline resulted in only a small relatively fixed net increment in ammonia excretion (2 to 7 mEq.) despite increasing magnitude of the ammonium chloride loads (Figure 5). To be

FIG. 5. EFFECT OF GLUTAMATE, ASPARTATE AND PROLINE ON NET URINARY AMMONIA EXCRETION FOLLOWING STEPWISE INCREASE IN THE MAGNITUDE OF THE CHRONIC AMMONIUM CHLORIDE LOAD.

The numbers in the columns refer to the net increase in urinary ammonia after the substrate load.
certain that a small but significant increase did not occur, an amino acid solution containing 600 mM of a mixture of amino acids composed for the most part of glutamic acid, proline and aspartic acid was infused at two different levels of ammonium chloride administration. Total ammonia excretion was greater, but again there was no progressive net increase in ammonia excretion with increasing magnitude of the ammonium chloride load (Figure 5, last two columns).

3. The effect of increased ammonia production following amino acid administration on total urinary acid excretion

The invariable and significant augmentation of total acid excretion \([\text{NH}_4^+ + (\text{TA} - \text{HCO}_3^-)]\) after amino acid loads is indicated by the four typical protocols listed in Table III.

4. Relationship between ammonia production and transport

The administration of precursor amino acid to subjects maintained on a chronic strong acid load invariably causes a rise in urine pH as \(\text{NH}_4^+\) excretion increases. Typical changes in urine pH and ammonia excretion which occurred in one subject are shown in Figure 6. Mean ammonia excretion for the five subjects studied increased significantly \((p < 0.001)\) from 6.1 to 10.1 mEq. per hour concomitant with a significant rise \((p = < 0.001)\) in mean urine pH from 5.60 to 6.12. Similar increases in urine pH following amino acid loads have been found in dogs by some (25) but not by others (2).

When subjects continuously received \(\text{NH}_4\text{Cl}\) it was noted that, after a steady state was attained at each level of \(\text{NH}_4\text{Cl}\) administration, urinary ammonia increased as the magnitude of the chronic
acid load increased but urinary pH, instead of falling, rose progressively (Figure 7). In order to determine whether or not, during chronic acid loads, urinary ammonia excretion is independent of changes in urine pH, acute alkalosis was produced by the infusion of sodium bicarbonate during a time when ammonia production was being stimulated toward maximal levels. One subject (M. A. S.) was maintained on 15 Gm. of NH₄Cl a day for about 30 days. At this time 600 mM of glycine was administered, resulting in an increase in ammonia excretion from 8 to 15.6 mEq. per hour. The identical experiment was repeated but in addition during the second study 250 mM of NaHCO₃ was infused producing an acute alkalosis, serum CO₂ rising from 22 to 35 mEq. per L. and urine pH from 5.59 to 7.63. The expected increase in ammonia excretion incident to the anticipated increase in ammonia production was not forthcoming. Instead urinary ammonia excretion fell progressively from 8.2 to 2.4 mEq. per hour. These experiments are compared in Figure 8.

**DISCUSSION**

The administration of an amino acid during the chronic maintenance of NH₄Cl loads of increasing magnitude results in three types of response: 1) l-Lysine and presumably similar diaminomonicarboxylic acids (2) elicit no increase in urinary ammonia excretion; 2) glutamate, aspartate and proline result in a small, constant increment in ammonia excretion; 3) glycine, glutamine, asparagine, l-alanine, l-leucine and d-alanine each progressively augment ammonia excretion in stepwise fashion paralleling the increase in ammonium chloride load. These data indicate that certain amino acids augment ammonia excretion, and also in the light of certain theoretical considerations help to identify the character of the renal enzymatic adaptation.

These experiments, designed to determine in vivo adaptation of the renal ammonia-producing enzymes, were based on the premise that at any given substrate concentration more end-product will be formed per unit time if more enzyme is present. If the chronic administration of strong acid loads results in an increase in renal tubular enzyme activity,⁸ then the renal production of ammonia following the same quantity of amino acid substrate before and after enzyme adaptation should differ quantitatively. After enzyme adaptation, the identical substrate load should result in an increased ammonia production ascribable to the increased enzyme activity. This hypothesis is schematically represented in Figure 9.

Before enzyme adaptation, as substrate concentration is increased from S₁, the plasma concentration before the administration of the fixed single amino acid load, to S₂, the plasma concentration after the substrate load, ammonia production per unit time by the nonadapted enzyme E¹ increases from P₁ to P². The net increase in ammonia pro-

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⁸ The increased enzyme activity that attends the adaptive process may be the result of either increased enzyme concentration, increased enzyme activator or decreased enzyme inhibitor.
The renal load is increased as identical amounts two E1 from the liver delivered enzymatic activity, reflecting the magnitude of the adaptive process. Since the amino acids were administered orally, traversing the liver prior to entry into the systemic circulation, the plasma concentration is, in part, dependent upon the fate of the amino acid in the liver. It is possible that as the magnitude of the NH4Cl load was progressively increased, the larger doses may have increasingly loaded the hepatic urea-synthesizing mechanism, thereby permitting larger amounts of amino acid to escape to the systemic circulation. Were this true, then the increasing ammonia excretion following identical oral substrate loads administered at high levels of NH4Cl intake, could represent increasing substrate reaching the kidney rather than enzyme adaptation; this seems unlikely for two reasons. One, the subjects did not

![Graph showing the relationship between ammonia excretion and urinary pH during chronic NH4Cl loads.](attachment:image)

**Fig. 7. Relationship Between Ammonia Excretion and Urinary pH During Chronic NH4Cl Loads**

As the chronic acid load was increased in stepwise fashion mean urinary ammonia excretion increased from 3.4 to 6.2 to 8.6 mEq per hour. Urinary pH also increased progressively from 5.18 to 5.39 and reached 5.96 as the NH4Cl load was increased from 5 to 10 to 15 Gm. per day.

production ascribable to the effect of the substrate load on enzyme E1 is therefore equal to (P4 minus P3). If the activity of the ammonia producing enzyme increases from E1 to E2 after the administration of a strong acid load, then the administration of the same quantity of substrate should result in a greater net increase in ammonia production (P4 minus P3) compared to the unadapted enzyme (E1). Finally, since the magnitude of the net increase in ammonia production after a substrate load is dependent on the enzymatic activity, the ratio (P4 minus P3)/(P3 minus P1), should reflect the magnitude of change in enzyme activity from E1 to E2.

In applying this hypothesis to in vivo experiments two assumptions were made:

1) The quantity of amino acid substrate delivered to the tubular enzymes per unit time after an identical substrate load does not progressively increase as the magnitude of the ammonium chloride load is increased from 5 to 15 Gm. per day. The renal hemodynamic changes that accompany the administration of ammonium chloride loads indicate a decrease in glomerular filtration rate and renal plasma flow (27). A decrease in the delivery of substrate with increasing NH4Cl loads would not interfere with the interpretations since it would tend to minimize, not maximize, the magnitude of the adaptive process. Since the amino acids were administered orally, traversing the liver prior to entry into the systemic circulation, the plasma concentration is, in part, dependent upon the fate of the amino acid in the liver. It is possible that as the magnitude of the NH4Cl load was progressively increased, the larger doses may have increasingly loaded the hepatic urea-synthesizing mechanism, thereby permitting larger amounts of amino acid to escape to the systemic circulation. Were this true, then the increasing ammonia excretion following identical oral substrate loads administered at high levels of NH4Cl intake, could represent increasing substrate reaching the kidney rather than enzyme adaptation; this seems unlikely for two reasons. One, the subjects did not
receive any NH₄Cl for 14 hours prior to control or amino acid loading studies (see Procedure). Moreover when 600 mM of asparagine was administered to a subject maintained on 5 Gm. of NH₄Cl per day, net ammonia excretion was 15 mEq. By contrast when maintained on 15 Gm. of NH₄Cl, the same subject's net ammonia excretion after 600 mM of glycine was 14 mEq. When maintained on 10 and 15 Gm. of NH₄Cl an intake of only 400 mM of glycine resulted in a net ammonia excretion of 22 and 41 mEq., respectively.

2) It must also be assumed in these experiments that ammonia excretion quantitatively reflects the magnitude of renal ammonia production. Ammonia produced within the tubular cells has three different fates, i.e., excretion into tubular urine; diffusion into renal venous blood; and intracellular utilization. Despite these alternative pathways, the present analysis is not adversely affected since the bulk of the ammonia formed in acidotic animals is excreted (1, 25) and, therefore, although not equal to, reflects quantitatively ammonia production. In support of this assumption are other studies from this laboratory in rats given increas-

![Image](attachment:image.png)

**FIG. 8. EFFECT OF ACUTE ALKALOSIS ON URINARY AMMONIUM EXCRETION AFTER AMINO ACID ADMINISTRATION**

See text for details.

![Image](attachment:image.png)

**FIG. 9. EFFECT OF ENZYME ADAPTATION ON THE MAGNITUDE OF AMMONIA PRODUCTION FOLLOWING A FIXED SUBSTRATE LOAD**

The conditions before enzyme adaptation (Eₑ) are shown on the left and after enzyme adaptation (Eₑ') on the right. Substrate concentration in arbitrary units is plotted along the abscissa and the rate of reaction, in arbitrary units of end-product formed per unit time, along the ordinate. The curved line in each panel describes the effect of increasing substrate concentration on the rate of reaction. The steeper curve on the right is the result of the increased activity of the adapted enzyme (26). See text for details.
ing ammonium chloride loads. In those experiments, ammonia excretion increased in direct proportion to ammonia production, as reflected by the in vitro determination of renal glutaminase activity (19).

Finally, in the present study at all levels of ammonium chloride intake, urinary pH was in the acid range. Moreover, mean urinary pH increased as the magnitude of the chronic ammonium chloride load was increased (Figure 7). With such changes in urinary pH, a progressively increasing net ammonia excretion in response to a fixed substrate load cannot be ascribed to improved conditions for transport.

In the light of these considerations, the data indicate that glycine oxidase, glutaminase, asparaginase, L-amino acid oxidase and D-amino acid oxidase show adaptive increases in the human kidney following the chronic administration of strong acids.

Although the data do not permit the measurement of the magnitude of the adaptive response of each of these five enzyme systems in absolute terms, the relative increase in enzyme activity as the size of the NH₄Cl load is increased can be estimated. Since the magnitude of the net increase in ammonia production after a substrate load is dependent on the enzyme activity, then the ratio, net increase in ammonia production after substrate during maintenance on 15 Gm. NH₄Cl per day/net increase in ammonia production after identical substrate load during maintenance on 10 Gm. of NH₄Cl per day, should reflect the magnitude of increase in enzyme activity as the chronic load of NH₄Cl was increased from 10 to 15 Gm. per day. These changes for each of the enzymes which undergo adaptive increase are compared in Table IV. The ratio of change for this group of enzymes varied from 1.5 to 1.9, indicating a one and a half to twofold increase in enzyme activity as the chronic NH₄Cl load was increased from 10 to 15 Gm. per day. This similarity in response suggests a symmetrical adaptation of several enzymes.

The similarity both in the magnitude of the adaptive increases of these five enzyme systems and in the magnitude of the net ammonia excretion after substrate loads suggests that when substrate is furnished in abundance by the administration of appropriate amino acids, these enzyme systems apparently can contribute equally to urinary ammonia. However, under normal circumstances there is a considerable variation in the concentration of precursor amino acids in plasma. In plasma, glutamine constitutes 20 to 25 per cent of the free α-amino nitrogen (15, 28) and L-alanine and glycine, another 24 per cent (29); D-alanine is apparently absent and asparagine is present in trivial concentrations (28). The contribu-

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\begin{array}{|c|c|c|c|c|}
\hline
\text{ENZYME} & \text{GLYCINE OXIDASE} & \text{GLUTAMINASE} & \text{ASPARAGINASE} & \text{L-AMINO ACID OXIDASE} & \text{D-AMINO ACID OXIDASE} \\
\hline
(p^* - p) & 1.6 & 1.9 & 1.9 & 1.9 & 1.5 \\
\hline
\end{array}
\]

* Refers to mean values for the five subjects.
† Equals net increase in NH₄ excretion after substrate load during chronic maintenance on 15 Gm. NH₄Cl per day.
‡ Equals net increase in NH₄ excretion after substrate loads during chronic maintenance on 10 Gm. NH₄Cl per day.

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* The changes in enzyme activity when the chronic NH₄Cl load was increased from 5 Gm. per day to higher levels were not used because the net increase in ammonia excretion after substrate at this level was negligible or small compared to the control values. On the other hand, the net increases in ammonia excretion after substrate when the subjects were maintained on 10 and 15 Gm. per day were larger and therefore were considered to reflect more accurately enzymatic alterations.
tion to urinary ammonia of the enzyme systems which are capable of adaptation may therefore vary widely because of substrate limitation.

The biological role of these enzymes, capable of adaptation, is not adequately defined. Significant activity of l-amino acid oxidase has been demonstrated in vitro in only the rat kidney (8). Despite its exceedingly low turnover number of 6, it nevertheless appears to be active in vivo in the human kidney and evidence from these studies and those in the rat (16) indicate that adaptation occurs. The disparity between the in vivo and in vitro evidence of its activity may exist because the ideal conditions for demonstrating in vitro activity have not been elucidated.

More enigmatic is the d-amino acid oxidase, an ubiquitous enzyme, found in all mammalian kidneys tested and having a high (1,440) turnover number (11). The presence of a highly active enzyme without apparent substrate beclouds its physiologic role. Nonetheless, the administration of d-alanine to human subjects resulted in augmentation in ammonia excretion and the evidence presented indicating adaptive change during chronic acid loading bespeaks a physiologic role. Amino acid racemases, capable of interconverting l and d forms, have been found in microorganisms (11). Such a racemase may be present in mammalian kidney and undergo adaptive change, thereby shunting, as acid load requires, a portion of the l-amino acid pool into d-amino acid substrate for ammonia production.

In contrast to the apparent physiologic importance in the renal regulation of acid-base balance of those enzyme systems showing adaptive changes is the role of glutamic dehydrogenase, aspartic transaminase and proline oxidase for which no evidence of adaptation was found. Lotspeich and Pitts failed to find any augmentation of ammonia excretion after the infusion of glutamic acid in acidotic dogs (2). Neither proline oxidase nor aspartic transaminase directly increase ammonia production (Table I). Both, however, augment the synthesis of glutamic acid which may then be deaminated by glutamic dehydrogenase, thereby increasing ammonia production. The failure of proline or aspartic acid administration to augment ammonia excretion progressively may, therefore, indicate that either proline oxidase and aspartic transaminase do not adapt or/and that glutamic dehydrogenase does not adapt.

The inferential identification of the adaptation of a specific enzyme by the administration of its precursor amino acid implicitly assumes that the rise in ammonia excretion under such circumstances is the consequence of the effect of the administered precursor amino acid on the specific enzyme system for which it is substrate. It is conceivable, however, that the administered amino acid might be converted by a transamination reaction to another substrate that would then act upon another enzyme system which then would be responsible for the augmented ammonia excretion. This possibility seems unlikely. All transaminases including glutamine-a-keto acid and aspargine-a-keto acid transaminase are pyridoxal phosphate and pyridoxamine phosphate dependent (13, 28); yet in pyridoxine deficient rats with reduced transaminase activity (30) the renal response to acid loads is normal, ammonia excretion is high and glutaminase adaptation occurs (31). This suggests that the transaminase systems do not contribute significantly to the renal production of ammonia. Moreover, it is noteworthy that even though amino acids can be readily transaminated to glutamate and glutamate to other amino acids, nevertheless the administration of glutamate, aspartate and proline resulted in only a very small increase in ammonia excretion. In addition, the adaptation of glutaminase (16–19), glycine oxidase and amino acid oxidase (16) has been shown in vitro to follow the administration of strong acid loads. For these reasons it seems likely that the changes in ammonia excretion after the administration of a specific amino acid can be ascribed to its effect on the enzyme system for which it is substrate.

The increased urinary ammonia excretion that follows amino acid administration was invariably associated with an increase in total acid excretion [NH₃ plus (TA–HCO₃)] These findings differ from those of Orloff and Berliner (25), who failed to find any consistent increase in total acid excretion following the infusion of dl-alanine into dogs. The discrepancy may be related either to species difference or to the fact that their experiments, conducted during acute acidosis, were different from the present studies which were performed during chronic acidosis.
The increased urinary pH that attends the increased ammonia excretion following substrate loads may be the result of either increased diffusion of ammonia from tubular cell to tubular urine thereby titrating tubular H+ and permitting further H+ for Na+ exchange by maintaining an increased gradient from cell to urine, or it may be the result of an increased exchange of cellular ammonium ion for urinary sodium ion which otherwise would have exchanged for H+ (32).

The relationship between urine pH and urinary ammonia that follows the chronic administration of strong acid loads of increasing magnitude contrasts sharply with the inverse relationship that prevails during acute acid loading (33, 34). This rise in urine pH with increases in chronic NH4Cl intake could result from the enhanced diffusion of ammonia from tubular cell to tubular urine caused by the high concentration gradients produced by accelerated ammonia formation. It is also possible that with increased activity of the renal ammonia-producing enzyme systems, the exchange of NH4+ for Na+ increases to some extent at the expense of H+ for Na+ exchange, resulting thereby in a rise in urine pH. Whatever explanation is valid, one consequence of renal enzymatic adaptation is the excretion of increased amounts of ammonia at any given urine pH (19, 35).

Evidence from the present experiments indicates that the superimposition of an acute alkalosis in a subject chronically maintained on 15 Gm. of NH4Cl per day prevents the usual increase in ammonia excretion after a substrate load and results instead in a sharp decline in ammonia excretion (Figure 8). Data concerning the effects of alkalosis upon ammonia production are conflicting. Van Slyke and associates (1) reported that the diminution in urinary ammonia excretion which attends the changing from hydrochloric acid acidosis to bicarbonate alkalosis is associated with a decreased renal utilization of glutamine (1, 9) and a decreased renal production of ammonia. Other evidence (36) indicates unaltered ammonia production and decreased urinary ammonia excretion associated with an acute alkalosis.

SUMMARY

One hundred twenty-five experiments were performed on five subjects maintained on chronic acid loads of increasing magnitude. Under these conditions precursor amino acids were administered and the renal ammonia producing enzymes in the human kidney inferentially identified. By comparing the magnitude of response in ammonia excretion to a fixed amino acid load at several different levels of ammonium chloride administration, those enzymes which adapt were identified. Evidence was presented indicating that, in the human kidney, glutaminase, asparaginase, glycine oxidase, l-amino acid oxidase and d-amino acid oxidase adapt to chronic acid loads whereas glutamic dehydrogenase, proline oxidase and aspartic transaminase do not.

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