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Metabolic Activity of Protein Amide Groups

Recently an enzymically catalyzed, Ca\(^{++}\)-activated incorporation of aliphatic amines, such as cadaverine or ethanolamine, into a protein fraction of liver, kidney, and brain was described (1). To the list of amines acting as substrates for incorporation may now be added putrescine, spermine, and histamine which were tested as C\(^{14}\)-labeled compounds. The incorporation of spermine and putrescine suggests an explanation for the inhibition of cadaverine incorporation by these amines reported previously. In addition it was shown that ammonia acted as an inhibitor of cadaverine incorporation (1). Therefore, the incorporation of N\(^{15}\)-ammonia into the protein fraction active in amine fixation was tested. A purified protein fraction (1) of guinea pig liver (105 mg.) was incubated with N\(^{15}\)H\(_2\)NO\(_3\) (80 μM; 60 atom % excess N\(^{15}\)) in a total volume of 8 ml. of 0.04 M barbital buffer, pH 8.2, in the absence (A) and presence (B) of 160 μM Ca\(^{++}\) at 37°C. for 30 min. The incubation mixtures were dialyzed against a large volume of 0.01 M phos-

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phate buffer, pH 7.6, for 14 hr. (two changes). The protein was precipitated with HClO₄, washed (2), hydrolyzed in 6 N HCl, the hydrolyzate made alkaline, NH₃ collected and analyzed for N¹⁵ concentration. A: 0.0; B: 2.1 atom % excess N¹⁵. It is assumed that the ammonia obtained corresponds mainly to the amide groups of the protein. The results suggest that amines as well as ammonia are incorporated by replacement of amide groups of the protein, a mechanism already proposed for the Ca++-activated incorporation of lysine into protein (3). The extent of the above-found incorporation of N¹⁴H₃ into protein would correspond to a net replacement of 3% of the amide groups.

During the course of this study it was observed that the protein fraction active in amine incorporation was inactivated very easily, particularly when incubated in the presence of Ca++. It could be shown that during this inactivation considerably greater amounts of ammonia were liberated in the presence than in the absence of Ca++ (Table I). The Ca++-activated liberation of ammonia exceeded considerably the estimated amount of amide groups replaced by C¹⁴-labeled amines or N¹⁵ ammonia. The ammonia liberation was activated by Ca++ but not by Mg++, Sr++, or Ba++, nor was it shown by another protein fraction from liver inactive in amine incorporation. At present the fraction of the ammonia derived from sources other than the amide groups of protein has not been established (5). The possibility is now under investigation that the liberation of ammonia in the presence of Ca++ is not solely hydrolytic but may be an exchange of the amide group with the ε-amino group of protein-bound lysine with liberation of ammonia. This possibility is supported by a study of Ca++-inactivated protein from which, after exhaustive deamination with nitrous acid, small amounts of intact lysine could be isolated.

In summary, these experiments point to the occurrence in liver and other organs of a protein fraction with metabolically labile amide groups. Whether

| TABLE I |
| Ammonia Liberation by Protein Fraction of Liver |
| µM NH₃ liberated per 100 mg. protein |
| -Ca++ | +Ca++ (80 µM) |
|---|---|---|
| 1 | P | 3 | 8.6 |
| 2 | P + cadaverine (16 µM) | 2.8 | 8.4 |
| 3 | P + cadaverine (16 µM)* | 1.1 |
| 4 | P + Sr (Ba or Mg) (80 µM) | 0.7-3.1 |
| 5 | P' | 0.4 |

P partially purified fraction (obtained at 18 to 26% ammonium sulfate saturation of 100,000 x g supernatant fluid of guinea pig liver homogenates in 0.25 M sucrose solution) which incorporated 800 cts./min./infinite thickness of C¹⁴-cadaverine (1). This corresponds to 2.3 µM cadaverine incorporation per 100 mg. protein. Total volume of 4 ml. of 0.04 M barbital buffer containing 100 mg. P, incubated at 37°C. for 60 min.

* Experiment 3: reaction stopped at 0 time. P' protein fraction inactive in cadaverine incorporation obtained from liver homogenate at 25 to 45% ammonium sulfate saturation. Reactions stopped by addition of HClO₄; NH₃ determined according to (4).
Substrate and enzyme protein are two different entities cannot be stated at present. The activity of the amide groups in amine incorporation may be of considerable biological significance as pointed out previously (1), not only as concerns the fixation and metabolism of biologically important amines but also because of the possibility of the formation of modified proteins. The Ca⁺⁺-activated liberation of ammonia from proteins supports the recent claim that some unassigned tissue ammonia originates in the amide groups of proteins (6). The biological significance of a change of properties of tissue proteins by loss of the amide group cannot be assessed at present. A possible involvement of the ε-amino group of protein-bound lysine in the mechanism of amide-ammonia liberation is of particular interest since an analysis of collagen (7) appears to indicate the natural occurrence of lysine substituted in this amino group.

References


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