

Thus there exists an imposing body of kinetic evidence which requires that the mechanism of hydrolysis of "specific" substrates by α -chymotrypsin follow eq. 1.

(10) M. L. Bender, G. E. Clement, F. J. Kézdy and B. Zerner, *J. Am. Chem. Soc.*, **85**, 358 (1963).

(11) Department of Chemistry, Harvard University, Cambridge 38, Massachusetts.

(12) Alfred P. Sloan Foundation Research Fellow.

DEPARTMENT OF CHEMISTRY
NORTHWESTERN UNIVERSITY
EVANSTON, ILLINOIS

BURI ZERNER¹¹
MYRON L. BENDER¹²

RECEIVED NOVEMBER 17, 1962

THE APPARENT MOLAR VOLUME OF SODIUM IN AMMONIA AT -45°

Sir:

We have previously² reported measurements of the apparent molar volume, V , of sodium in ammonia at 0° ; V , defined as (volume of solution - volume of solvent)/g. atoms Na, is nearly independent of concentration, being 56.3 ml. mole⁻¹ at 0.35 M and about 2 ml. mole⁻¹ greater at 0.009 M . The change is but little outside of experimental error. Evers and Filbert³ have reported similar measurements at -45° ; they found V to decrease slowly from 62 ml. mole⁻¹ at 0.2 M to 60 at 0.050 M and steeply to 53 at 0.040 M , and then to rise steeply from 53 at 0.023 M to 59 at 0.018 M , 64 at 0.012 M and 65 at 0.0036 M . They have stated their accuracy to be better than 3%; however, it would be expected that the probable errors of such measurements in dilute solution would be roughly inversely proportional to the concentration.

The dissimilarity of these two reports has led us to perform measurements at -45° also; the procedure was essentially as previously described. A stirred methyl alcohol bath was used, cooled by a system described elsewhere⁴ controlled by a Hallikainen controller and electrical heater and monitored with a platinum thermometer, Mueller bridge, and recorder. The dilatometer bulbs were sealed together, rather than being assembled with waxed joints. Results are given in Table I; over-all uncertainties are estimated separately for each run. Within experimental accuracy, V is constant at 60.7. Agreement with the data of Evers and Filbert at the higher concentrations is satisfactory, but there is no evidence for a minimum or any considerable change at lower concentrations.

TABLE I
APPARENT MOLAR VOLUME OF Na IN NH₃ AT -45°

| C (mole liter ⁻¹) | V (ml. mole ⁻¹) | C (mole liter ⁻¹) | V (ml. mole ⁻¹) |
|----------------------------------|--------------------------------|----------------------------------|--------------------------------|
| 0.1640 | 60.7 \pm 0.2 | | |
| .0940 | 60.7 \pm 0.2 | 0.0276 | 61.5 \pm 1.0 |
| .0493 | 59.8 \pm 1.0 | .0269 | 61.2 \pm 0.8 |
| .0428 | 59.7 \pm 1.0 | .0234 | 59.9 \pm 0.8 |
| | 59.7 \pm 0.5 | .0161 | 60.4 \pm 1.2 |
| .0388 | 60.0 \pm 0.7 | .0114 | 59.4 \pm 1.2 |
| .0317 | 61.4 \pm 1.2 | .0058 | 59.9 \pm 3.0 |

It may be noted that V decreases from -45 to 0° ; for aqueous solutions of electrolytes at ordinary temperatures, V increases with increasing temperature.

It is also of interest to note that both the volume and absorption spectra⁵ of sodium-ammonia solutions are essentially constant through a concentration range where the magnetic, electrochemical, and thermochemical properties change drastically. Becker, Lind-

(1) This work was performed under the auspices of the U. S. Atomic Energy Commission.

(2) S. R. Gunn and L. G. Green, *J. Chem. Phys.*, **36**, 363 (1962).

(3) E. C. Evers and A. M. Filbert, *J. Am. Chem. Soc.*, **83**, 3337 (1961).

(4) S. R. Gunn, *Rev. Sci. Instruments*, **33**, 880 (1962).

(5) M. Gold and W. L. Jolly, *Inorg. Chem.*, **1**, 818 (1962).

quist, and Alder⁶ proposed the equilibria $(1/2)M_2 = M = M^+ + e^-$ to explain the conductance and magnetic properties. Gold, Jolly and Pitzer⁷ proposed that the intermediate non-conducting paramagnetic species M is an ordinary ion-pair and that M_2 is a quadrupolar ionic assembly. However, dissociation of salt ion-pairs in ammonia produces a decrease of 20 to 30 ml. mole⁻¹ in V .

(6) E. Becker, R. H. Lindquist and B. J. Alder, *J. Chem. Phys.*, **25**, 971 (1956).

(7) M. Gold, W. L. Jolly and K. S. Pitzer, *J. Am. Chem. Soc.*, **84**, 2264 (1962).

UNIVERSITY OF CALIFORNIA
LAWRENCE RADIATION LABORATORY
LIVERMORE, CALIFORNIA

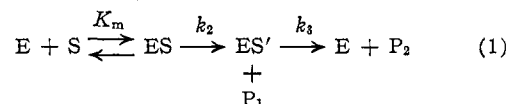
STUART R. GUNN
LEROY G. GREEN

RECEIVED NOVEMBER 9, 1962

SIGMOID AND BELL-SHAPED pH-RATE PROFILES IN α -CHYMOTRYPSIN-CATALYZED HYDROLYSES. A MECHANISTIC CORRELATION¹

Sir:

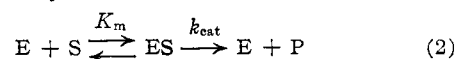
The thesis enunciated earlier, that all chymotrypsin-catalyzed hydrolyses follow eq. 1, has been substantiated by spectrophotometric and kinetic evidence.^{2,3} On experimental and theoretical grounds the two steps of acylation and deacylation (k_2 and k_3) were postulated



to be mechanistically equivalent, the enzymatic components of their transition states being identical.³ On this basis, the same pH dependence might be expected in all chymotrypsin steps and reactions. However, some chymotrypsin-catalyzed reactions show bell shaped pH-rate profiles while others show sigmoid pH-rate profiles.⁴

The pH dependencies of the rate constants of the two individual steps (k_2 and k_3) (eq. 1 and Table I) indicate that pH-rate profiles of *deacylation* obey a *sigmoid* curve, while the pH-rate profile of the only *acylation* step observed over a sufficient pH range follows a *bell-shaped* curve.

The pH dependencies of the *catalytic rate constants* (turnover) (eq. 2 and Table II) of this research and literature data may be correlated (on the basis of Table



I) with the pH dependence of a rate-controlling acylation or deacylation step. In the hydrolyses of acetyl-L-tryptophanamide and acetyl-L-tryptophan ethyl ester, acylation and deacylation, respectively, were shown to be rate-controlling.² The respective bell-shaped and sigmoid pH dependencies of these reactions are in complete accord with such a designation. The steric and electronic similarity of acetyl-L-tryptophanamide to the other amides and hydroxamides in Table II suggests that they should exhibit acylation rate-controlling steps and therefore bell-shaped pH-rate profiles, as found experimentally (the other amide "bells" do not, however, reflect k_{cat}). The similarity of the three ethyl ester substrates in Table II suggests that they should exhibit rate-controlling deacylation steps and sigmoid pH-rate profiles, as found experimentally. All these results are self-consistent and lead one to

(1) This research was supported by Grant H-5726 of the National Institutes of Health. Paper XIX in the series, The Mechanism of Action of Proteolytic Enzymes.

(2) B. Zerner and M. L. Bender, *J. Am. Chem. Soc.*, **85**, 356 (1963).

(3) M. L. Bender, *ibid.*, **84**, 2582 (1962).

(4) These observations apply to reactions in which K_m (or $K_{m(app)}$) has been separated from the appropriate rate constant.