CXV. THE CREATINE-CREATININE EQUILI-BRIUM. THE APPARENT DISSOCIATION CONSTANTS OF CREATINE AND CREATININE.

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THE facile conversion of creatine into creatinine under the influence of strong acids and the partial reversal of the reaction in neutral and in alkaline solutions have long been familiar. Yet only recently has there become available any quantitative data upon the equilibrium conditions. Hahn and Barkan [1920] were the first to report any systematic kinetic studies. They determined the equilibrium constant in solutions of sodium hydroxide of varying concentration, observed that the component velocities increased with increasing [OH-] and showed that the order of the reaction creatinine \rightarrow creatine, under these conditions, was that of a reversible monomolecular system. In a molar solution of hydrochloric acid, on the other hand, the reverse reaction went to completion and followed the course of a simple monomolecular change. Hahn and Meyer [1923] later reported a few observations which indicated that the velocity of this reaction in buffered solutions increased rapidly from $p_{\rm H}$ 6 to 4. A more elaborate study of this system has been made by Edgar and his associates. Edgar and Wakefield [1923] determined the monomolecular velocity constants (k_2) of the dehydration of creatine in hydrochloric acid solutions of varying concentration and at various temperatures. They succeeded in relating k_2 to the temperature by means of the Arrhenius equation and, further, concluded that k_2 was, probably, proportional to the hydrogen ion activity. Finally, Edgar and Shiver [1925] have made an extensive series of determinations of the equilibrium constant (K) at 50° in buffered solutions of $p_{\rm H}$ values 1 to 6. Hahn and Barkan had suggested that their observations could be interpreted upon the assumption that the molecular species whose concentrations determined the equilibrium were the undissociated molecules of creatine and of creatinine. Confirming this, Edgar and Shiver obtained fairly satisfactory agreement between the observed values of K and those calculated from the dissociation constants of the two bases and the value of K in unbuffered solution (*i.e.* where the two reactants were not significantly dissociated).

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Consideration of the results summarised above will indicate that, although the hypothesis of Hahn and Barkan has been useful in co-ordinating the equilibrium data, it fails to comprehend the relations between [H+] and the velocities of the reactions. If the equilibrium be determined by the ratio of the concentrations of the undissociated molecules of creatine and creatinine, the velocities of the two contributing reactions should be governed by the same factors. Thus, if k_b be the dissociation constant of either reactant and k the velocity constant for its decomposition into the other, then k should vary with $\frac{[OH^-]}{k_b + [OH^-]}$. That is to say, the velocity should be inversely proportional to $[H^+]$ on the acid side of the buffer range of k_b and should be independent of [H+] on the alkaline side. The available experimental evidence, however, indicates that the velocities are proportional to [H+] in solutions of strong acids and are inversely proportional in strongly alkaline solution. The reactions under discussion are of such direct biological interest that it was decided to undertake a series of kinetic studies in buffered solutions between $p_{\rm H}$ 1 and 10 as an attempt to elucidate these discrepancies.

It was necessary, in the first place, that there should be available dependable values for the dissociation constants of creatine and creatinine. Since the values in the literature differ rather seriously, a redetermination of these constants was undertaken.

Dissociation constants of creatine and creatinine.

The method employed was that of electrometric titration of dilute solutions of the two bases with standard hydrochloric acid in the presence of the hydrogen electrode. The routine technique of this laboratory has already been described [Cannan and Knight, 1927]. The reference electrode was a saturated calomel cell which was standardised against 0.05 M acid potassium phthalate [Clark, 1922]. Two palladinised gold-plated platinum electrodes were employed as duplicate hydrogen electrodes. No difficulty was encountered in attaining stable potentials in any of the solutions titrated and the two electrodes agreed within 0.3 mv. at all significant points on the titration curves.

The creatine was prepared from a good commercial sample by repeated recrystallisation from water. After drying to constant weight over calcium chloride, a typical preparation gave

Nitrogen (Kjeldahl)	28.19~%	Water 12.18 %
Theory for $C_4H_9O_2N_3.H_2O$	28.19	12.08

A saturated solution gave no reaction for creatinine upon applying Weyl's test.

Creatinine was prepared from the creatine by treating the latter with hydrochloric acid gas and subsequent liberation of the base by aqueous ammonia. The product was recrystallised from acetone [Edgar and Hinegardner, 1923]. Nitrogen and water determinations were quantitative for anhydrous creatinine. Folin's colorimetric method for the determination of

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creatinine (using creatinine picrate as standard) gave results in agreement with the nitrogen values, but this method is, admittedly, not sufficiently accurate to detect traces of impurity in creatinine.

It will be convenient, throughout the paper, to conduct the discussion in terms of hydrogen ions rather than of hydroxyl ions and, consequently, all constants will be treated as though they were acid constants. That is to say, the kation of a base will be regarded as an acid which dissociates a hydrogen ion [Bronsted, 1923]. The constants so derived (k') are related to the familiar k_b values by the equation $p_{k'} = p_{kw} - p_{kb}$.

Table I. Uncorrected apparent dissociation constant of creatinine.

	Molar			
Authors	conc.	Temp.	$p_{k'}$	k.,
Wood [1903]	0.1	40.2	2.97	3.57×10^{-11}
McNally [1926]		40 ·0	4.42	1.01 × 10−9
Cannan and Shore	0.1	30·0	4 ·77	$0.98 imes 10^{-9}$
>>	0.02	30.0	4.72	0·15 × 10−9
,,	0.1	25.0	4 ·78	$0.76 imes 10^{-9}$
McNally [1926]		25.0	4.71	0·70 × 10−9
Eadie and Hunter [1926]	0.1	20.0	4.87	0·64 × 10−9
Hahn and Barkan [1920]	0.04	17.0	4.44	0.19×10^{-9}
Cannan and Shore	0.02	15.0	4.91	$0.47 imes 10^{-9}$

Wood, and Hahn and Barkan calculated k_b from the degree of hydrolysis of solutions of the hydrochloride; Eadie and Hunter employed the electrometric titration; McNally's results are the mean of results from the conductance, hydrogen ion concentration and distribution of the hydrochloride.

In Table I are assembled several determinations of $p_{k'}$ for creatinine together with values calculated from the k_b values recorded in the literature. The important effect of temperature upon the constant is evident and renders difficult the comparison of the results of different observers. But it would seem that, apart from the two earliest determinations which were made with methods open to considerable experimental errors, the various values are in substantial agreement. If is unnecessary, therefore, to report our experimental data in any greater detail. For purposes of the analysis of the kinetic studies which follow, the value for the dissociation constant of creatinine at 30° will be taken to be $k' = 1.90 \times 10^{-5}$, *i.e.* $p_{k'} = 4.72$.

The case of creatine is less satisfactory. The various determinations are summarised in Table II. In Table III is given the analysis of a typical titration curve to indicate the degree of concordance of the data. The calculations have been made with the aid of the Henderson-Hasselbalch equation. The values of [H+], used in calculating the "corrected equivalents of acid," are obtained from the observed $p_{\rm H}$ after correction for the activity of the hydrogen ion by the equation $\log \dot{\tau}_{\rm H} = 0.20 \sqrt{\Sigma i v^z}$ [Simms, 1926]. $\tau_{\rm H}$ is the activity coefficient ratio for the hydrogen ion, $\Sigma i v^z$ is the sum of all the ion concentrations each multiplied by the z power of its valency. The value of z was assumed to be unity. The constants have not been corrected for activity.

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Table II. Uncorrected apparent dissociation constants of creatine.

Authors	Molar conc.	Temp.	k'_1	$p_{k'_1}$
Wood [1903]	0.1	40.2	2.1×10^{-3}	2.68
Cannan and Shore	0.1	30 •0	$2 \cdot 4 \times 10^{-3}$	2.62
**	0.02	30.0	$2 \cdot 4 \times 10^{-3}$	2.62
	0.1	$25 \cdot 0$	$2 \cdot 2 \times 10^{-3}$	2.66
Eadie and Hunter [1926]	0.02	20.0	0.9×10^{-3}	3.05
Hahn and Barkan [1920]	0.04	17.0	1.4×10^{-3}	2.85
Cannan and Shore	0.02	17.0	$2 \cdot 45 imes 10^{-3}$	2.61

Table III. Titration of 50 cc. 0.02 M creatine with 0.1 M hydrochloric acid.

Titre	<i>р</i> н	[H ⁺] corrected	$\begin{array}{c} \text{Corrected} \\ \text{equiv. acid} \\ = \underbrace{[\text{HCl}] - [\text{H}^+]}_{[\text{creatine}]} \end{array}$	$\log \frac{a}{1-a}$	<i>Pk</i> [•] 1
0.00	5.77		0.00		
0.20	4.41	$39.0 imes 10^{-6}$	0.0181	-1.74	2.67
0.52	3.95	$11 \cdot 2 imes 10^{-6}$	0.0452	1.33	2.62
1.02	3.64	$23 \cdot 4 \times 10^{-6}$	0.0901	1.01	2.63
2.01	3·3 0	$51\cdot3 imes 10^{-6}$	0.1744	0.68	2.62
3.02	3.08	$85 \cdot 1 \times 10^{-6}$	0.2568	0.47	2.63
4 ·00	2.92	123.0×10^{-6}	0.3337	0.30	2.62
4·98	2.79	166.0×10^{-6}	0.4064	0.16	2.63
6.00	2.67	224.0×10^{-6}	0.4749	-0.04	2.63
8.02	2.48	347.0×10^{-6}	0.6010	+0.18	2.64
10.00	2.34	501.0×10^{-6}	0.7000	0.36	2.70
13.03	2.15	794.0×10^{-6}	0.8016	0.61	2.76

It is probable that the last two calculations suffer by reason of the uncertainty of the correction for hydrogen ion activity.

It will be seen from Table II that differences exist between the determinations of different observers which cannot be attributed to differences of temperature or of concentration. In particular, it is difficult to explain the conflicting results of Eadie and Hunter and of ourselves since the same method was employed and was prosecuted with the same degree of precision. No plausible source of error in the titrimetric method peculiar to creatine suggests itself. The possibility of a significant amount of conversion of creatine into creatinine during the course of a titration seems to be excluded by the velocity measurements recorded in the second part of this paper. Provisionally we will take the value for k' at 30° as $2\cdot40 \times 10^{-3}$, *i.e.* $p_{k'} = 2\cdot62$.

A question of some interest to the chemical behaviour of creatine arises from a consideration of its electrolytic dissociation. The conventional formula for creatine contains both a carboxyl and an amino-group. Creatine might be expected to behave, therefore, as an ampholyte. Only basic properties are, however, evident in its chemical behaviour and only one dissociation constant is detected by titration. This is, therefore, described as a basic constant. Hahn and Fasold [1925] have, however, found that the solubility of creatine in solutions of sodium hydroxide is greater than in water and they conclude that some dissociation of creatine as an acid occurs in solutions of great hydroxyl concentration. From their observations they calculate a value of 14.28 for p_{k_a} . Now the allocation of the first dissociation constant of

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creatine to the amino-group and the assignment of only negligible acid properties to the carboxyl is difficult to justify upon the grounds of organic chemical experience. Yet it is usual to describe creatine as a base. A more plausible interpretation of the acid-base behaviour of this substance would seem to follow the application to it of Bjerrum's [1923] treatment of the aminoacids. The first constant $(k_1' = k_w/k_b)$ then becomes the acidic constant and the second—in this case, inaccessible—constant (k_2') is the association constant of the basic group. With this assignment of constants creatine becomes an acid comparable in strength with other carboxylic acids. At the same time a new difficulty is created for it is required that the basic dissociation shall be as great as that of the alkali hydroxides. It would be difficult to concede this to a simple amino-group and it is of interest, therefore, that creatine does not behave as a primary amine either towards nitrous acid or towards formaldehyde. In this connection the strong basic properties and anomalous behaviour of guanidine itself will be recalled. It is significant that several of the structural formulae which have been proposed to explain the anomalous reaction with nitrous acid contain a nitrogenous group which might be expected to dissociate strongly as a base [Hunter, 1928, p. 99].

The above considerations in no way prejudice the application of the dissociation constants to the co-ordination of kinetic data. It is a matter of no immediate moment whether the velocity of dehydration of creatine is determined by the concentration of undissociated creatine or of "zwitterion" —the mathematical relation to k_1 remains unmodified.

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Solutions of creatine (0.0106 M) and of creatinine (0.00354 M) were prepared in a series of the 0.05 M buffers recommended by Clark [1922]. The mixtures were covered with 10 cc. of toluene and stored in stoppered bottles in an air-bath maintained at $30^{\circ} \pm 1^{\circ}$. At intervals appropriate to each experiment, a sample was removed and the concentration of creatinine present was determined by the method of Folin. The standard solutions for this method were prepared from a purified specimen of creatinine picrate. The $p_{\rm H}$ values of the various reaction mixtures were determined at the beginning, and again at the conclusion, of each experiment by means of the hydrogen electrode. At the end of each experiment determination was also made of the total creatine + creatinine. In agreement with other investigators it was found that some conversion occurred of these two substances into products which reacted neither as creatine nor as creatinine. The extent of the loss during the period of experiment varied from 0.5 to 5 % according to the $p_{\rm H}$ of the solution. In view of the temperature at which the solutions were maintained and of the precarious antiseptic properties of toluene over long periods, the occurrence of bacterial decomposition may be suspected. This source of error cannot be absolutely excluded but the results are so concordant amongst themselves and fit in so well with the equilibrium data of Edgar and Shiver

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