The Solubility of Pd(NH₃)₂Cl₂ in Electrolyte Solutions¹

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Solubilities have been determined at 25° for *trans*-Pd(NH_a)₂Cl₂ in solutions of NH₄NO₃, NH₄Cl, and NaCl, in which media base hydrolysis is repressed. The data are interpreted in terms of the heterogeneous equilibrium involving the uncharged solute molecule **and** the homogeneous equilibrium for the first step of aquation, **The** concentration of neutral solute in the saturated solution is given as 9.5×10^{-4} *M* and the aquation equilibrium constant **as** 3.9 **X** 10-8.

The rates and equilibria of aquation (acid hydrolysis) of a number of complexes of Pt(1I) containing only C1-, H_2O , and NH₃ as ligands have been investigated by Martin and his coworkers.² The equilibrium constants range over a factor of nearly 1000, whereas the rate constants for the forward step vary by a factor of less than five, 3 indicating that nearly the entire kinetic effect is to be found in the reverse reaction. Banerjea and Tripathi4 have suggested that the mechanism of the substitution reactions of Pd(I1) differs from that for $Pt(II)$; and this difference is such that it may be expected that, considering the aquation of a series of complexes with various ligands, the effect on the trend in reaction rate within the series caused by replacing Pt by Pd will differ for the forward and the reverse reactions. In such case, the aquation equilibrium constants will vary through the series in qualitatively different ways for the complexes of the two metals.

One example may be seen from the following equilibrium constants for aquation^{2,5}: PtCl₄⁻², 30×10^{-3} ; PtCl₃(H₂O)⁻, 1.0 \times 10⁻³; PdCl₄⁻², 3×10^{-3} ; PdCl₃(H₂O)⁻, 4 $\times 10^{-3}$. An even more profound difference seems to be shown by the report by Banerjea and Tripathi⁴ that the aquation of trans- $Pd(NH_3)_2Cl_2$ proceeds to completion, while² for the *trans*-Pt complex $K = 8 \times 10^{-5}$

(3) For further discussion of this point see F. Basolo and R. *0.* **Pearson, "Mechanisms of Inorganic Reactions," John Wiley and Sons, Inc., New York, N. Y., 1958, pp. 152ff.**

(4) D. Banerjea and K. K. Tripathi, *J.* **Inorg.** *N~cl. Chcm., 7,* **78 (1954).**

(6) H. A. Droll, B. P. Block, and W. C. Fernelius, *J. Phys. Chcm., 61,* **1000 (1957).**

and for the *cis-Pt, K'* (at ionic strength 0.3) = 3.3 \times 10⁻³. On the other hand, the interpretation of Banerjea and Tripathi in **this** connection is based on the assumption that the observed conductivity of the solution is due to the aquation process, whereas it had been recognized earlier⁶ that $Pd(NH_3)_2Cl_2$ undergoes a decomposition when placed in contact with water. It therefore seems likely that it was this latter process, rather than aquation, which was observed by Banerjea and Tripathi.

The lack of consistency in the nineteenth century data⁷ on the solubility of $Pd(NH_3)_2Cl_2$ also can be attributed to this decomposition. It is the purpose of the present study to determine the solubility under conditions in which the decomposition is prevented and to utilize these data to investigate the aquation equilibria.

The Nature of the Decomposition Process.- Qualitative evidence was obtained for the decomposition by the very noticeable darkening of the solid when put into contact with water, with the formation of a brownish colloidal substance. In solutions unsaturated with respect to Pd- $(NH₃)₂Cl₂$ a reddish brown sol formed, and this ultimately coagulated into a brown gel. The presence of the colloidal material rendered phase separation very difficult and led invariably to erratic results in attempts to measure solubility, In the unsaturated solution, decomposition was substantially complete, whereas in the more concentrated systems it was always possible **to** recover some yellow crystals of starting material, even after prolonged boiling.

It was observed, however, that in the presence

⁽¹⁾ This research was supported in part by the Office of Naval Research.

^{(2) (}a) D. S. Martin, Jr., and R. J. Adams, in S. Kirschner, *ed.,* **"Advances in the Chemistry of the Coordination Compounds,'' The Macmillan Company, New York, N. Y., 1961, p. 679 (includes** earlier references); (b) T. S. Elleman, J. W. Reishus, and D. S. **Martin, Jr.,** *J. Am. Chem. Sod., 81,* **10 (1959); (c) C. I. Sanders and D. S. Martin, Jr.,** *ibid.,* **83, 807 (1961); (d) J. W. Reishus and D. S. Martin, Jr., ibid., 83, 2457 (1961).**

⁽⁶⁾ (a) H. D. K. Drew, F. W. Pinkard. G. H. Preston, and W. Wardlaw, *1. Chem. Sod.,* **1895 (1932); (b) A. A. Grinberg, V. M. Shulman, and S. 1. Khorunzhenkov,** *Isv. Ins;. Isuchcniyu Plofiny,* **12, 119 (1936).**

⁽⁷⁾ See, e.&, H. MPllef, *Liebigs* **Ann.,** *88,* **341 (1853).**

of a sufficient concentration of $NH₄^+$, Cl⁻⁻, or H⁺ the darkening did not take place. The highest electrolyte concentrations at which decomposition was observed to occur were: NH₄Cl, 0.0010 M; NaCl, 0.015 *M*; NH₄NO₃, 0.005 *M*; HCl, HNO₃, HC104, less than **0.005 M.** The color of either partly or completely decomposed $Pd(NH_3)_2Cl_2$ was restored by the addition of NH4C1 or HC104. Aqueous NaNO₃ had no effect in preventing decomposition and NaOH at millimolar concentrations promoted it.

The following experiment served to identify the ultimate products of decomposition : In **500** cc. **of** H_2O was placed 0.587 mmole of $Pd(NH_3)_2Cl_2$ (less than saturation) and the mixture was heated at 100° for *4* **hr.** until coagulation of the preeipitate was complete. The solid, after having been dried at 105' for 24 hr., weighed **52.0** mg.; after standing in the air, the mass increased to **54.3** mg. These data are consistent with the formula PdO. xH_2O , with x ranging from 0.6 to 1.⁸ A portion of the solid was dissolved in $6 \, M \, HNO₃$ and roughly analyzed for Cl, *via* AgCl; Cl found: 4% , corresponding to $\frac{1}{7}$ Cl atom per Pd atom.

The colorless filtrate contained at most a trace of Pd, and no excess H^+ or NH₃. NH₄⁺ and Cl⁻⁻ were identified qualitatively; a $\frac{1}{10}$ aliquot, on evaporation, yielded 4.6 mg. of residue, corresponding to **2.2** moles of NH4C1 per mole of original $Pd(NH_3)_2Cl_2.$

These observations suggest that the decomposition of the compound results from base hydrolysis in which formation of the hydroxo complex is followed by olation and eliminatioa of NHs. The prevention of this hydrolysis by H^+ or Cl^- thus is due to interference with the first step. The effect of $NH₄$ ⁺ in preventing the reaction, however, is too large to be explained on the basis of its acidity. From this fact, and from the observation that in none of the experiments (save only those in which H^+ or OH^- was added) did the pH of the solution differ appreciably from that of the distilled water used (*ca.* 6), it is apparent that approximately one NH_4^+ , resulting from NH_3 elimination, accompanies each Cl^- formed in the initial step of the base hydrolysis. (On the other hand, it cannot be asserted whether the hydrolysis proceeds as far as hydrous PdO in the systems saturated with respect to the complex.)

Experimental

For each determination, the sample of $Pd(NH_3)_2Cl_2$ was agitated in contact with the appropriate salt solution in a constant-temperature bath maintained at 25.00". Aliquot samples were removed from time to time until the analysis of an initially unsaturated system agreed with that of an initially supersaturated system (prepared by previously warming to *ca*. 40°, at which temperature botn solubility and rate of solution are much greater than at *25").* Twenty-four to thirty-six hours was required for equilibrium.

Analysis was performed on samples pipetted through cotton woo!. Pd was determined gravimetrically as the salt of dimethylglyoxime, following substantially the procedure of Treadwell and Hall." It was found that a more crystalline precipitate could be obtained for those samples initially low in salt concentration by the addition of ammonium acetate to about 0.01 *M.*

 $trans-Pd(NH₈)₂Cl₂$ was prepared following the procedure given by Cohen and Davidson.¹⁰ A sample which had reached equilibrium with an $NH₄NO₃$ solution analyzed 51.0% Pd; another, from an I\rH4cI solution, *50.8%* Pd. CaIcd.: **50.4%** Pd. Several of the equilibrated solids were tested for the cis-isomer. following the procedure of Grinberg and Shulman, $¹¹$ with negative results in each case.</sup>

ReSultS

In Table I are given the solubilities¹² of Pd- $(NH₃)₂Cl₂$ in solutions of $NH₄NO₃$, $NH₄Cl$, and NaCl at 25° . Data for systems in which there was noticeable decomposition have not been included.

TABLE I

SOLUBILITIES OF $Pd(NH_3)_2Cl_2$		
Salt concn.,	Soly.,	
М	М	Salt
0.00939	0.00300	NH4NO3
.03052	.00316	NH4NO:
.0939	.00334	NH4NO3
.00180	.00227	NH.Cl
.00199	.00226	NH.Cl
.00300	.00200	NH Cl
.00600	.00165	NH Cl
.0100	.00141	NH.Cl
.0150	.00122	NH.Cl
.0150	.00129	NH4C1
.0300	.00113	NH ₄ Cl
.0601	.00105	NH.Cl
. 1002	.00102	NH.Cl
.0377	.00106	NaC1
.0754	.00103	NaCl
.1005	.00100	NaCl

⁽⁰⁾ F. P. Treadwell and W. T. Hall, "Analytical Chemistry," (10) A. J. Cohen **and** N. **Davidson,** *J. Am. Chem.* Soc., **73, 1955** Vol. **11,** John **Wiley and Sons, Inc., New York,** N. **Y., 1942, p. 140. (1051).**

⁽⁵⁾ See 0. Glemser and *G.* **Peuschel,** *2. alzorg. allgem. Chem.,* **281, 44 (1955).**

⁽¹¹⁾ A. A. Grinberg and V. M. **Shulman,** *Dokl.* **Akad.** *Nauk SSSR,* **215 (1933).**

⁽¹²⁾ Throughout this paper concentrations are given in units Brackets of **moles of indicated solute per liter of solution** *(MI.* **are used around the formulas** *to* **indicate such concentration.**

Discussion

The solubility data can be explained by considering the equilibria

$$
Pd(NH_8)_2Cl_2(s) = Pd(NH_8)_2Cl_2(aq) \qquad (1)
$$

 $Pd(NH_3)_2Cl_2(aq) + H_2O =$ $Pd(NH_3)_2Cl(H_2O)^+ + Cl^-$ (2)

of which reaction 2 is the first step of aquation. In the development that follows, the iolubility product

$$
K_{\rm s} = [\text{Pd}(\text{NH}_3)_2\text{Cl}(\text{H}_2\text{O})^+][\text{Cl}^-]K_{\gamma}
$$

where K_{γ} represents the product of the activity coefficients of the two ions. *x* is the concentration of neutral $Pd(NH_3)_2Cl_2$ in solution, s is the molar solubility, and c the concentration of chloride ion added.12 It will be assumed that *x* is constant in all the saturated solutions and that K_{γ} can be represented by the Giintelberg approximation. **la**

It then follows that

$$
s = x + (K_s/K_\gamma)[\text{Cl}^-]^{-1}
$$
 (3)

A first approximation for *x* was found from a plot based on eq. 3 for large values of c , where $[Cl^-] \cong$ c. This value of *x* then was used to construct Fig. 1, in which s is plotted against $(K\gamma [Cl^-])^{-1}$ for the NH₄Cl and NH₄NO₃ points. The intercept agrees with the first approximation of $x = 9.5 \times 10^{-4} M$ and the slope gives $K_s = 3.7 \times 10^{-6}$ (molar units); or the equilibrium constant for reaction $2 = K_s/x$ $= 3.9 \times 10^{-3}$.

The points for the NaCl solutions, if included in Fig. 1, would show a barely perceptible negative

Fig. 1.-Plot of eq. 3: solid circles, NH₄Cl; open circles, NH₄NO₃; concentrations in moles/l.

deviation, due perhaps to a slight extent of base hydrolysis (since the only source of $NH₄$ + is from

(13) E. Giintelberg, *2. physik.* **Chem., 138, 199 (1926).**

this hydrolysis itself), the chloride from which would tend to lower the solubility. For the NH_{4-} $NO₃$ solutions, however, there should be sufficient Cl^- from the aquation to prevent decomposition. The observed deviations here may be the result of the approximations made with regard to activity coefficients, both of ions and of the neutral molecule. Such effects would be most pronounced in the $NH₄NO₈$ solutions, because of the relatively high ionic strength combined with low chloride ion concentration.

A hypothetical solubility of $Pd(NH_3)_2Cl_2$ in water may be obtained from eq. 3 by setting $c =$ 0 and estimating *K,.* The result, 0.00297 *M,* agrees closely with that found by linear interpolation of the $NH₄NO₃$ data to zero nitrate, 0.00295 *M.* This agreement, plus the lack of positive deviations from the straight line in Fig. 1 in the region of low chloride, indicates that there is no large tendency for further aquation to $Pd(NH₃)₂$ - $(H₂O)₂$ ⁺². On the other hand, a slight extent of this reaction cannot be ruled out: computations performed by selecting *K* for the second step of aquation as 3×10^{-4} and using K_s as 3.5×10^{-6} (only slightly smaller than found from Fig. 1) showed that the solubility data could be reproduced to within 3×10^{-5} *M* for all points in the chloride solutions. On this basis, it is felt that the value given previously for K_s has an uncertainty of perhaps 10% .

If now the equilibrium constant for the aquation of Pd(NH₃)₂Cl₂, 3.9 \times 10⁻³, is compared with those of the several palladium and platinum complexes referred to earlier,^{2,5} it is seen that the large variation in *K* manifested by the Pt compounds does not appear for these particular palladium analogs,14 consistent with a difference in mechanism of the sort discussed previously. It must be recognized, however, that it is unwarranted to make any more than a general comparison of this kind until isomer analysis data are available for the palladium compounds. Thus, while there is no reason to doubt that all the complexes present in a solution of *trans*- $Pt(NH_3)_2Cl_2$ are in the *trans* configuration, the same cannot be assumed, *a priori,* for the Pd complex. There is no necessary connection between the thermodynamic stability of the *trans* form in the crystalline phase¹¹ and that in aqueous solution; and the expected electrostatic stabilization of the *trans* form is not always found, as with $Pt(NH_3)Cl_2(H_2O)^{2b}$ and with

⁽¹⁴⁾ But note the smaller values (ref. 5) for PdClz(Hz0)z and PdCl(Hs0)s +.