

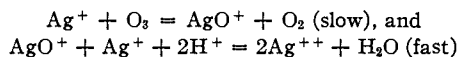
(21.2 *N*, 90% nitric acid, density 1.49) gave, however, the striking result that, as the acid becomes nearly anhydrous, scarcely any argentic silver is produced—after seventy-five hours not even enough to give more than a slight brown color to the solution. The table further shows that the percentage of argentic silver produced at any given acid concentration decreases, though its absolute concentration increases, when the total silver concentration increases. It is seen that a reagent 0.2 formal in $\text{Ag}(\text{NO}_3)_2$ can be prepared in 10 or 15 *N* nitric acid.

Summary

It has been shown that at 0 and 24° ozone slowly oxidizes argentous nitrate in nitric acid solution to a black soluble argentic nitrate; but that the reaction ceases before it has become complete, with the establishment of a steady state. This steady state is shown to arise, not at all from the occurrence of the reverse reaction, but from the reduction of the argentic salt by the water with evolution of oxygen. (For the overall reactions see the equations at the beginning of this paper.) The argentic salt is assumed to consist mainly of bipoisitive silver on the basis of the results of the next following paper.

The initial rate of the silver oxidation at 0° was found to be proportional to the argentous and

ozone concentrations. This is accounted for by the following mechanism which involves the direct production of tripositive silver in the form of an oxygenated ion



The initial rate is found to be only slightly affected by the acid concentration or by the ionic strength.

The rate of the argentic decomposition by water (at 0 and 24°) was found as a first approximation to be proportional to the square of the argentic concentration (except when the latter is high), and inversely proportional to the argentous concentration, and to be greatly diminished by increasing the acid concentration. The mechanism is discussed (see equations 13, 14, and 15); but complications in the reaction make the conclusions somewhat uncertain. The effect of temperature on the reaction rate was found to be very much larger than for most reactions.

The steady state at 0° was roughly determined at various acid concentrations. The percentage of the silver converted into the argentic form varied from 18% in 1.8 *N* nitric acid to about 90% in 12 *N* nitric acid when the total silver was about 0.04 formal, but in 21 *N* (90%) nitric acid scarcely any formation of argentic salt resulted.

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Argentic Salts in Acid Solution. II. The Oxidation State of Argentic Salts

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Purpose of this Investigation

In the preceding article Noyes, Hoard and Pitzer¹ described the general nature and the rates of the reactions by which argentic salt is produced by the ozone oxidation of acid argentous nitrate, and by which it is reduced by water. Before proceeding further with the description of these researches, however, it seemed essential to establish the state of oxidation, whether bipoisitive or tripositive, in which argentic silver exists in nitric acid solution. To a description of experiments made with this purpose this article is devoted.

It is remarkable that this question has re-

(1) Noyes, Hoard and Pitzer, *THIS JOURNAL*, **57**, 1221 (1935).

mained unanswered so long in the case of a readily obtainable oxidation state of a common element—an oxidation state, moreover, that explains the striking catalytic effects of silver salts. Furthermore, the question is of much theoretical interest in relation to the valence states compatible with the periodic system, inasmuch as copper exists mainly unipositive and bipoisitive, and gold unipositive and tripositive.

Previous Investigations

The most definite indications previously existing as to the oxidation state of argentic silver in its solutions were furnished by the composition of the solid phases that separate in preparing or

in diluting the solutions or upon adding other substances to them, but these indications seem contradictory.

First, it may be mentioned that Barbieri² found upon mixing an aqueous solution of potassium peroxysulfate with a solution of silver nitrate in pyridine (Py) that orange-colored crystals separated of the composition $\text{AgS}_2\text{O}_8 \cdot 4\text{Py}$, and that this compound forms isomorphous crystals with the previously prepared cupric compound $\text{Cu}_2\text{S}_2\text{O}_8 \cdot 4\text{Py}$, both these facts showing the existence of a complex salt containing bivalent silver, as is more clearly expressed by the formula $[\text{Py}_4\text{Ag}]^{++}\text{S}_2\text{O}_8^-$. There have since been prepared³ complex salts of bivalent silver with various other bases (having coordination numbers four or six) and with other anions than peroxysulfate.

On the other hand, several investigators (Higson,⁴ Yost,⁵ Barbieri,⁶ Jirsa⁷) have shown that argentic solutions give precipitates with water of solid basic salts that contain tripositive silver. This is proved by the fact that they yield on drying compounds such as $\text{AgONO}_3 \cdot \text{Ag}_2\text{O}_3 \cdot 4\text{AgO}$ or $(\text{AgO})_2\text{SO}_4 \cdot 6\text{AgO}$. Such salts are precipitated even from a solution prepared by dissolving pure silver oxide in nitric acid, and even though this dissolving is attended by evolution of oxygen (Jirsa). The results obtained by Yost⁵ and by Carman⁸ are especially significant, in that they show that all the silver is originally precipitated in the tripositive state. Thus the compounds prepared by them from neutral potassium peroxysulfate and silver nitrate contain per atomic weight of silver 2.0 oxidation equivalents (measured with reference to reduction to argentous salt). Carman showed, however, that if the precipitates stood one to three hours before filtering, the ratio drops to a fairly constant value of 1.51–1.53, corresponding to about equal quantities of bivalent and tripositive silver.

The fact that the rate of oxidation of argentous salt by peroxysulfate, as well as by ozone,

is proportional to the first power, not to the square, of the argentous concentration, indicating that the mechanism involves primarily the production of tripositive silver, has also doubtless aided in creating a presumption that this is the form of silver that predominates in the solution; but in reality this fact would be fully accounted for by the existence of only a very small proportion of tripositive silver.

The Oxidation State Determined by a Solubility Method

Principle of the Method.—The principle of this method of determining the oxidation state of the silver is illustrated by the following description of the preliminary experiments which were made by K. S. Pitzer. Oxygen containing 5–7% ozone was passed for twelve hours through a gas bottle (without a porous plate) containing about 200 ml. of 10 or 16 *N* nitric acid and a great excess (25 g.) of crystalline silver nitrate at 0°. Known volume samples of the (approximately) steady-state mixture were then drawn off and analyzed: (1) for oxidizing power, correcting the result for the dissolved ozone, and (2) for total silver (univalent plus bivalent plus tripositive). Then, by an experiment in which (to agitate the liquid) oxygen instead of ozonized oxygen was passed for forty hours, the solubility of argentous nitrate was determined in a solution of the same acid content at the same temperature, and this (univalent) silver was subtracted from the total silver found in the steady-state mixture.

This method was fairly precise because the solubility of silver nitrate in strong nitric acid was found to be unexpectedly small (0.12–0.36 *f* at 0°). It involves the assumption that this solubility is not appreciably affected by the argentic silver, which, however, was present at only 0.22 to 0.25 formal.

The Results of the Preliminary Experiments.—These are given in Table I. It will be seen that in the more accurate, second experiment the

TABLE I

OXIDATION STATE OF ARGENTIC SILVER DETERMINED BY A SOLUBILITY METHOD: PRELIMINARY RESULTS AT 0°

Expt.	HNO_3 Volume- formal	AgNO_3 Solubility	Reaction mixture (Ag^{I}) + (Ag^{III})		Oxid. Equiv.	Ratio of oxid. equiv. to (Ag^{II}) + (Ag^{III})
1	15.2	0.119	0.366	0.247	0.217	0.89
2	10.0	.357	.583	.226	.218	.96
			.583	.226	.222	.98
			.588	.231	.222	.96

(2) Barbieri, *Atti accad. Lincei*, [5] **21**, 560–563 (1912); *Gazz. chim. ital.*, **42**, 7 (1912).

(3) Hieber and Mühlbauer, *Ber.*, **61**, 2149 (1928); Morgan and Burstall, *J. Chem. Soc.*, **133**, 2594 (1930); Barbieri, *Ber.*, **60**, 2424–2427 (1927), and *Atti accad. Lincei*, [6] **15**, 44–47 (1932).

(4) Higson, *J. Chem. Soc.*, **119**, 2048 (1921).

(5) Yost, *THIS JOURNAL*, **48**, 152–164 (1926).

(6) Barbieri, *Atti accad. Lincei*, [6] **13**, 882–887 (1931).

(7) Jirsa, *Z. anorg. Chem.*, **148**, 130–140 (1925); **158**, 33–60, 61–66 (1926).

(8) Carman, *Trans. Faraday Soc.*, **30**, 566–577 (1934).

number of oxidation equivalents was 97% of the number of formula weights of the total argentic salt, indicating that this salt was substantially all in the bipoisitive state.

Procedure for the Final Determination.—A final experiment was then made with great care under more favorable conditions. In this experiment we were fortunate in having the detailed coöperation of Professor Isaburo Wada, Director of the Wada Research Laboratory, of Tokio, who was then visiting this Institute. In developing the analytical methods and in making the analyses we were ably assisted also by Messrs. C. D. Coryell and Alexander Kossiakoff. In this direction we are also indebted to Professor E. H. Swift for important suggestions.

The principle involved in determining the state of oxidation of the silver was the same as before. The temperature at which the solubility of silver nitrate was determined and at which the steady-state mixture was prepared was $-5.8 \pm 0.2^\circ$, instead of 0° , concentrated nitric acid (15.2 *N* at 20°) being used, as in one of the previous experiments. These conditions made possible much greater experimental accuracy, for at this lower temperature the ratio of argentic to argentous silver is much larger, and the spontaneous decomposition of the argentic silver is much slower. Moreover, precise analytical methods whose reliability had been well tested were used, and the quantities of the samples were all determined by weight, not by volume.

The Procedure in Detail.—The solutions were prepared in glass-stoppered gas absorption bottles with porous sintered glass plates, of the kind described in the preceding article,¹ to which reference may be made also for the details of the ozonizer, thermostat, and other features of the apparatus. In one gas bottle were placed 7 g. of fine silver nitrate crystals and 150 ml. of a nitric acid solution 15.55 *N* (by volume) at 0° (or 15.19 *N* at 20°), and in another bottle were placed 20 g. of silver nitrate crystals and 200 ml. of the same nitric acid; and these bottles were kept in a large alcohol-water bath at -6.15° for twenty-four hours, and then at $-5.8 \pm 0.2^\circ$ for forty-eight hours more. During this whole period there was passed steadily a slow current of oxygen through the first bottle (to stir its contents), and of ozonized oxygen through the second bottle (to produce the steady-state mixture). Before entering the latter bottle (so as to avoid evaporation of the oxidized solution and to determine the solubility of the ozone) the ozonized oxygen was passed through a bottle containing only nitric acid of the same strength.

The partial pressure of the ozone was found to be 0.0625 atm. and its solubility at -5.8° in the nitric acid (15.2 *N* at 20°) was found to be 0.00321 mole per liter of solution. These quantities were determined by driving the gas,

by means of a current of carbon dioxide, out of a gas bottle of known volume, filled first with the ozonized oxygen, and second containing known volumes of this gas and of its saturated solution in 15.2 *N* nitric acid. The expelled gas was passed in each case into a 0.5 *f* potassium iodide solution saturated with sodium hydrocarbonate; then the mixture was acidified, and the liberated iodine titrated with thiosulfate.

After three days the ozone was displaced from the steady-state mixture by passing a rapid current of oxygen through it for about five minutes, and duplicate samples of the pure silver nitrate solution and of the steady-state mixture were removed from the respective bottles through their side-arms (so that the crystals of undissolved silver nitrate would be retained by the porous plate near the bottom of the bottles).

The samples of unozonized silver nitrate were run into empty weighed flasks, and those of the steady-state mixture into weighed flasks containing 1.8 g. of thalious nitrate, placed in the cold bath. After coming to room temperature these flasks and their contents were weighed, thus giving the weights of the samples, which were later analyzed as will now be described.

Methods of Analysis.—The solubility of the silver nitrate in the pure nitric acid solution was determined by diluting this solution, nearly neutralizing it with ammonium hydroxide, and adding a 2% excess of sodium chloride solution. The precipitated silver chloride was collected in a weighed porous plate filter funnel, dried at 200° , and weighed. It was proved that even the very large quantity of ammonium nitrate present did not dissolve a significant quantity of silver chloride.

The determination of the oxidizing power of the steady-state mixture involved much preliminary study, owing to the presence in the mixture of concentrated nitric acid and owing to the rapid decomposition of argentic silver when the solution is diluted or allowed to warm up. It was necessary to find some reducing substance not oxidized by concentrated nitric acid, but oxidized instantaneously, even at -5.8° , by argentic silver—a substance, furthermore, whose oxidation product could afterward be analyzed for with accuracy. After many experiments thalious nitrate was found to be well suited for this purpose. It is not oxidized by cold nitric acid, but reduces argentic silver at -5.8° within a few seconds, as was shown by the flash-like disappearance of the black color as soon as the cold sample of the steady-state mixture was run onto the solid thalious nitrate and the mixture was shaken. It was shown, moreover, by an experiment in a small eudiometer that this reaction is attended by no oxygen evolution, and therefore by no spontaneous or catalytic decomposition of the argentic silver.

After a long study of possible thallium determinations (made mainly by Mr. C. D. Coryell) it was found that the quantity of thallic salt produced could be determined by neutralizing the nitric acid in the cooled sample with 15 *N* ammonium hydroxide, filtering off and washing the brown thallic hydroxide, dissolving this in cold 3 *N* sulfuric acid, and analyzing the solution for its thallic content. This last analysis was made by using the method described by Kolthoff⁹ and by Zintl and Rienäcker¹⁰ for titrating thal-

(9) Kolthoff, *Rec. trav. chim.*, **41**, 189 (1922).

(10) Zintl and Rienäcker, *Z. anorg. allgem. Chem.*, **153**, 275 (1926).

TABLE II

Solution	Sample withdrawn, g.	Determinations of total silver			Determinations of oxidation-equivalents		
		% of sample analyzed	AgCl obtained, mg.	At. wt. Ag per kg. solution	% of sample analyzed	KBrO ₃ used, ml.	Oxid. eq. per kg. solution
Satd. AgNO ₃ solution	36.78	100	316.3	0.0600
	74.01	100	632.2	.0597
Steady-state mixture	38.36	51.05	851.5	.3033	26.31	45.96	0.2334
	38.36	48.81	812.4	.3027	38.54	67.22	.2331
	38.36	34.98	61.13	.2335
	38.45	51.66	859.1	.3017	35.32	61.78	.2331
	38.45	48.19	803.9	.3014	30.05	52.51	.2329
	38.45	34.19	59.75	.2329
				Mean 0.3023			Mean 0.2332

lous thallium with standard bromate solution with the aid of methyl orange as an oxidation indicator. This is possible since under proper conditions this indicator is decolorized only after an excess of bromate over that required to oxidize the thallos salt has been added.

In detail this thallic determination was carried out as follows. The thallic sulfate solution (about 140 ml.) obtained by dissolving the thallic hydroxide from each of the two samples of the steady-state mixture was divided by weighing into three nearly equal portions. To each of these portions were added 11 g. of solid potassium chloride and one drop of 0.2% methyl orange solution, and, after heating the solution to 50–60°, bromate solution (standardized against pure thallos chloride) was run in from a buret until the indicator was decolorized, thus estimating any thallos thallium (in fact about 2%) that was carried down in the thallic hydroxide precipitate. Then the solution was heated nearly to boiling, and sulfur dioxide gas was passed through it for ten minutes, after which the solution was boiled for ten to fifteen minutes to expel the sulfur dioxide, and was titrated at 50–60° with the bromate solution, adding one drop of methyl orange solution as the end-point was approached. The amount of bromate required to decolorize the indicator and the amount added before reducing the thallic salt with sulfur dioxide were subtracted, thus giving the oxidizing power of the thallic sulfate solution and hence of the original argentic solution. Test analyses, made with weighed samples of pure thallos chloride oxidized to the thallic form, had previously shown this analytical procedure to be remarkably precise.

The total silver present in the same samples of the steady-state mixture was determined by acidifying the filtrate from the thallic hydroxide precipitate, adding sodium chloride, and weighing the precipitated silver chloride, as in the case of the pure silver nitrate solution.

Experimental Results and Final Value of Oxidation State.—The experimental data for all the determinations so made are given in Table II. The headings are all self-explanatory, except that the bromate solution used was 0.05124 *N* (in an oxidation sense).

The mean values derived from all the data of Table II, and the final values of the oxidation state, are shown in Table III. In order to be able

to refer the mean values of the quantities of silver and of the oxidation equivalents to 1 kg. of acid, as well as to 1 kg. of solution, the latter values have been corrected by subtracting from the weight of the solution that of the solid silver nitrate calculated to have been dissolved, and (in the case of the steady-state mixture) also that of the oxygen calculated to have been absorbed in producing the argentic silver. The nitric acid used was 15.2 *N* at 20°.

TABLE III

FINAL MEAN VALUES OF THE SOLUBILITY OF SILVER NITRATE AND OF THE COMPOSITION OF THE STEADY-STATE MIXTURE AT -5.8°

Solution	Atomic weights of silver		Oxidation equivalents		Ratio of oxid. eq. to argentic salt
	In kg. of solution of 15.2 <i>N</i> HNO ₃	Per kg. HNO ₃	In kg. of solution of 15.2 <i>N</i> HNO ₃	Per kg. HNO ₃	
Steady-state mixture	0.3023	0.3193	0.2332	0.2463	...
Satd. AgNO ₃ solution	.0598	.0604	.0	.0	...
Difference	.2425	.2589	.2332	.2463	0.951

The difference 0.2589 between the number of equivalents of silver in the steady-state mixture and in the pure silver nitrate solution (referred to the same weight of nitric acid) evidently corresponds, at least approximately, to the argentic silver in the mixture, and the ratio 0.2463/0.2589 of the oxidation equivalents to this difference, which equals 0.951, shows how much the oxidation number of the argentic silver exceeds that of argentous silver. This corresponds to an oxidation number of 1.951 for the silver in the argentic salt. The silver is therefore *bi-positive* in the argentic compound existing in the steady-state mixture produced at -5.8° in nitric acid 15.2 *N* at 20°.

The difference between 1.95 and 2.00 may be reasonably attributed to the spontaneous decomposition of the compound during the time required for removal of the ozone from the mixture and for the addition of the sample to the thallos

nitrate. And there is doubtless some inaccuracy in the assumption that the amounts of argentous nitrate present as such in the pure solution and in the steady-state mixture are identical.

The Oxidation State Determined by Displacement of the Steady State

EXPERIMENTS BY CLARENCE L. DUNN

Principle of the Method.—A second principle employed in these researches for determining the oxidation state of argentous silver involves displacement of the steady state. Namely, since argentous solutions have been prepared under conditions where the ratio of oxidation equivalents to total silver is already not very far from unity (which would correspond to complete conversion to the bivalent state), one may expect to determine, upon making the conditions progressively much more favorable, whether this ratio approaches unity asymptotically, or whether it goes through this value without any noticeable discontinuity. For example, starting with the fact that with 0.025 *f* silver nitrate in 12 *N* nitric acid 0.93 oxidation equivalents of argentous silver are produced at 0° for each formula weight of (total) silver present when the solution is brought to a steady state with ozone at 0.06 atm., one might determine upon increasing the ozone partial pressure, say to 0.4 atm., whether the equivalents of argentous salt increased only from 0.93 to nearly 1.00, or whether they increased up to values much above unity. This evidently involved on the experimental side making ozone at these higher partial pressures, saturating acid silver nitrate solution with it, and analyzing the mixture.

Preparation of Ozone Gas at High Concentration and Oxidation of the Silver Solution.—The concentrated ozone gas needed for these experiments was made from tank oxygen which was purified by passing it through an all-glass train of tubes containing an electrically heated platinum wire, soda-lime, strong sulfuric acid and phosphorus pentoxide, and through a very fine sintered glass filter (to remove particles of dust, rubber, etc.). This gas after emerging from the ozonizer, when it contained about 6% of ozone by volume, was fractionally condensed in a 15-ml. cylindrical bulb sealed to the ozonizer. This bulb had a sealed-in entrance tube delivering 5 cm. above the bottom of the bulb and had a side-arm near the top out of which the gas issued. The bulb was placed within a Dewar tube containing liquid air, which could be brought to any desired level by raising or lowering the Dewar tube. During the liquefaction the tube was kept so low as to condense ozone containing very little oxygen, the composition of the condensate being judged by the depth of its blue

color. After about 2 ml. of liquid had condensed, requiring about three hours, the liquid was distilled at a moderate rate for a period of about three hours into a 12 *N* nitric acid solution exactly 0.025 *f* in silver nitrate, contained in a gas bottle in a thermostat at 0°. This solution had been previously treated with 6% ozone until the steady state corresponding thereto had been practically reached. The gas emerging from the reaction bottle contained at the end of one hour about 40% of ozone, and at the end of two and one-half hours about 65% of ozone by volume. A concentration of 12 *N* nitric acid was used, since this had been found to be approximately that at which the largest fraction of argentous salt was produced with 6 to 8% ozone. The small silver concentration of 0.025 formal was used, since the oxidation is larger the more dilute the silver, and since on the other hand this is the smallest concentration that would give sufficient analytic accuracy.

Analysis of the Reaction Mixture.—Before taking a sample for determining the oxidizing power the dissolved ozone was removed from the gas bottle by subjecting it for one minute to the reduced pressure of a water pump and then drawing a rapid stream of oxygen through it for half a minute more. A blank experiment with 12 *N* nitric acid saturated with ozone showed that the ozone remaining after this treatment was negligible. Then there was withdrawn from the gas bottle with a cooled pipet 50 ml. of the reaction mixture, and this was delivered into a two-fold excess of 0.05 *f* thallic nitrate solution at 0°. The mixture was analyzed for thallic thallium as was described in the earlier part of this article, except that hydrochloric acid was used for dissolving the ammonium hydroxide precipitate of thallic hydroxide, instead of sulfuric acid; approximately 0.05 *N* potassium bromate was used as before for titrating the thallium after reducing it by sulfur dioxide.

Experimental Results and Conclusion as to the Oxidation State.—A series of experiments was first made to determine the steady state resulting with 9 or 6% ozone at 0° in 12 *N* nitric acid solutions 0.025 *f* in silver nitrate. Such solutions were treated with ozonized oxygen of this strength for several days till no further change in their oxidizing power took place. The ratio of oxidation equivalents to total silver equivalents had at the steady state in different experiments the following percentage values: with 9% ozone, 97.4, 96.5, 98.4%; with 6% ozone, 93.0% (84.0% when 15 *N* nitric acid was used).

Then eight other experiments were made with 12 *N* nitric acid 0.025 *f* in silver nitrate at 0°, by carrying the oxidation at first only to 34 to 83% with 6% ozone, and afterward making it as complete as possible by treatment with 40 to 65% ozone for three hours in the way described above. The percentage ratios of oxidizing power to total silver obtained with the 6% and the 40–65% ozone are those shown in Table IV in the first and second rows, respectively.

TABLE IV

Ozone used, %	Percentage of oxidation equivalents to total silver equivalents							
6	83.2	81.0	80.4	80.0	56.5	49.0	48.0	34.1
40-65	100.0	100.8	98.1	99.0	99.1	103.0	90.0	98.6

It will be seen, irrespective of the oxidation (83-34%) attained with the 6% ozone, that the more concentrated ozone raised the percentage of oxidation equivalents to equivalents of total silver (except in one case) only to 98.1 to 103.0%—never to any larger value such as would correspond to any considerable proportion of tripositive silver. (Incomplete removal of ozone probably accounts for the one slightly high value of 103%.) The conclusion seems, therefore, justified that under the conditions here involved (12 *N* nitric acid, 0.025 *f* silver nitrate, 0°) even fairly concentrated ozone converts little, if any, of the silver into a state more highly oxidized than the bivalent state.

The Oxidation State of Argentic Silver Determined from the Magnetic Susceptibility of its Solutions

EXPERIMENTS BY CLARENCE L. DUNN

Theoretical Principle of the Method.—A third method of studying the state of oxidation of argentic silver was suggested by Professor Linus Pauling. This rests upon the fact that the magnetic moment of the silver ion in its different valence states and the corresponding change in the susceptibility of the solution may be expected to afford a direct indication of the oxidation state of the silver.

The theoretical principle involved in the method is the simple one, valid for certain types of atoms, that the magnetic moment of an atom or ion is determined primarily by the spin of its unpaired electrons, and therefore by the number of such electrons, being 1.73 Bohr magnetons for one such electron, 2.83 Bohr magnetons for two, and so forth.¹¹ The univalent silver ion has a complete sub-shell of 10 outer (4d) electrons and has no spin magnetic moment, but only a diamagnetic one. The bivalent silver ion with only nine of these outer electrons has an odd and therefore an unpaired electron, and has a spin moment of 1.73 magnetons. The trivalent silver ion with eight outer (4d) electrons probably has only paired electrons, and is probably therefore dia-

(11) Stoner, *Phil. Mag.*, **8**, 250 (1929); Pauling, *THIS JOURNAL*, **58**, 1391 (1931).

magnetic; for this has been shown by Janes¹² to be true of bivalent palladium, which has the same electronic configuration. It is, however, possible, though less likely, that the tripositive silver ion, like the structurally similar bivalent nickel ion (with eight 3d electrons), has two unpaired electrons, in which case it would be paramagnetic, but would have a spin moment (2.83 magnetons) different from that of the bivalent silver ion.

While in an ion having unpaired electrons the spin of such electrons usually gives to the ion its main magnetic moment, yet the orbital motion of such electrons may produce a large additional moment. Much of this orbital moment, however, is usually found (except in the rare-earth group) to be quenched by the surrounding molecules. This quenching of the orbital moment of a bivalent silver ion may be expected to occur to about the same extent as does that of the similarly constituted cupric ion, for which in solution several investigators¹³ have found a total moment of about 1.92 magnetons. Moreover, Sugden¹⁴ has found that this moment lies between 1.74 and 2.16 magnetons for various solid salts of bivalent silver with pyridine. On the other hand, the trivalent silver ion, if it is not diamagnetic, would probably have a total moment like that of the bivalent nickel ion, which has been found by Cabrera and Duperier¹⁵ to have a total moment of 3.19 magnetons in the solution of its nitrate.

Experimental Method of Determining Susceptibilities.—The effect of oxidizing the silver upon its magnetic moment was derived by comparing the susceptibilities of appropriate solutions. These susceptibilities were measured by Gouy's method in the form described by Shaffer and Taylor,¹⁶ with an apparatus set up and used in this Laboratory by Dr. E. B. Wilson, Jr., and Dr. G. W. Wheland. The process consists in weighing the solution in a vertical glass tube between the pole-pieces of an electromagnet both before and after the magnet is activated. Then the two weighings are repeated under identical conditions with a substance of known susceptibility (pure water). The ratio of this known value to the change in weight observed gives the constant of the apparatus, by which other observed weight differences can be multiplied to give susceptibilities.

Experimental Procedure in Detail.—In detail the experiments were made in the following way. Solutions of a

(12) Janes, *ibid.*, **57**, 472 (1935).

(13) Liebknecht and Wills (1900), Jacobsohn (1916), Cabrera and Moles (1914-1917), Cherbuliez (1917). For a curve showing their results, see Blrch, *J. phys. radium*, **9**, 138 (1928).

(14) Sugden, *J. Chem. Soc.*, 161-170 (1932).

(15) Cabrera and Duperier, by Barnett, "International Critical Tables," 1929, Vol. VI, p. 352.

(16) Shaffer and Taylor, *THIS JOURNAL*, **48**, 844 (1926).

definite amount of silver nitrate in 10 *N* nitric acid were made, and the larger part of each was treated with ozone at 0° for twenty-four hours. The rest of the unoxidized solutions was reserved for comparative susceptibility measurements.

Samples of the oxidized and unoxidized solutions were placed in a tube of about 7 mm. bore fitted with a ground-glass top and of the design described by Shaffer and Taylor, and were weighed in and out of the magnetic field. The same tube was then charged with conductivity water, and the weighing repeated, this serving as a standard of known susceptibility.

The oxidized solution, before it was weighed, had been brought to 22° and the ozone had been removed by a current of air. The concentration of the oxidized silver in the solution was determined by removing samples for analysis just before and just after making the weighing with the field on, and taking the mean of the two results, this being done to eliminate the effect of the rather rapid decomposition of the oxidized silver. Each sample was run into a weighed flask containing standard thalious nitrate solution, the flask was reweighed to give the weight of the sample, and the amount of thalious salt unoxidized was determined with bromate by the method described above.

As a check the oxidized solution was allowed to decompose spontaneously in the weighing tube, and measurements were made on the resulting solution. The susceptibility of this solution was found to be identical with that of the original unoxidized solution.

Accuracy in the magnetic measurements required that the concentration of oxidized silver be as high as possible, yet the available ranges of concentrations of acid and of total silver were very limited, owing to the fact that the percentage of the silver oxidized by 6% ozone is large only when the acid is concentrated, and then the silver nitrate is only moderately soluble. A number of measurements were made with each of two silver solutions, 10 *N* in nitric acid, in which the percentage of oxidized silver was varied in the different experiments by allowing decomposition to take place or by retreating the solution with ozone at 0°.

The Experimental Data and the Calculated Magnetic Moments.—The data of the measure-

ments at 22° are given in the first seven columns of Table V, and the change in susceptibility upon oxidation and the magnetic moments of the oxidized silver atom, calculated as described below, are given in the last three columns. The symbols have the following significance.

c_1 and c_2 are the equivalents of argentic silver per gram of sample found in the oxidized solution just before and just after it was weighed in the magnetic field; and the mean of these values is the concentration to which the observed change in weight is considered to correspond. As given in the table these quantities are multiplied by 10^6 .

Δw_{UN} is the increase in weight in milligrams which the unoxidized sample showed when the magnetic field was on over that when it was off; Δw_{OX} is the corresponding increase in weight which the oxidized sample (with the same total silver concentration) showed under the same conditions; and Δw_{H_2O} is the increase in weight which pure water showed under these conditions.

$\chi_{OX} - \chi_{UN}$ is the change in susceptibility per cubic centimeter (in cgs. units) produced by oxidizing the sample. This is calculated by the expression $-7.49 \times 10^7 (\Delta w_{OX} - \Delta w_{UN}) / \Delta w_{H_2O}$, the numerical factor being the volume susceptibility of water against air at 22°. μ is the increase in magnetic moment, expressed in Bohr magnetons, attending the conversion of one atom of argentous silver into argentic silver, calculated from the data in the way described below, first assuming the silver to be bipoisitive, and second, assuming it to be tripositive.

Experiments 1-6 were made with a solution containing 0.252 formula weight and experiments 7-12 with one containing 0.257 formula weight,

TABLE V
SUSCEPTIBILITY MEASUREMENTS AND THE MAGNETIC MOMENT OF ARGENTIC ION

Expt.	Equivalents of $Ag_{OX} \times 10^6$ per g.			Δw_{UN} in mg.	Δw_{OX} in mg.	Δw_{H_2O} in mg.	$\chi_{OX} - \chi_{UN}$ $\times 10^7$	μ in magnetons	
	$10^6 c_1$	$10^6 c_2$	Mean					For Ag^{II}	For Ag^{III}
1	61.2	56.6	58.9	-20.9	-16.7	-22.5	1.30	1.97	2.78
2	91.0	83.5	87.2	20.6	14.8	22.5	1.93	1.97	2.78
3	85.8	76.7	81.2	19.5	14.2	21.3	1.87	2.01	2.84
4	116.0	108.1	112.0	20.5	12.9	22.4	2.54	1.99	2.82
5	92.1	84.6	88.4	20.4	14.4	22.3	2.02	2.00	2.83
6	65.1	60.1	62.5	20.7	16.5	22.6	1.39	1.97	2.79
7	32.5	31.4	31.9	20.4	18.3	22.3	0.705	1.96	2.78
8	63.0	62.0	62.5	20.4	16.2	22.3	1.41	1.98	2.81
9	67.1	60.5	63.7	20.0	15.8	21.8	1.44	1.99	2.81
10	105.9	98.5	102.2	19.5	13.0	21.3	2.28	1.97	2.79
11	110.3	97.3	104.8	19.5	12.8	21.3	2.36	1.98	2.80
12	88.5	73.5	81.0	19.8	14.3	21.7	1.90	2.02	2.86
								Mean 1.98	2.81

of total silver per kilogram of solution. The densities at 22° of the two (unoxidized) solutions were 1.343 and 1.350, respectively.

Calculation of the Magnetons per Atom of Argentic Silver.—The calculation of the magnetic moment from the data was made as follows.

(1) The magnetic changes in weight are first converted into change in susceptibility per cubic centimeter (in cgs. units) in the way described above. Thus is calculated the quantity $\chi_{OX} - \chi_{UN}$, recorded in the table.

(2) This change in susceptibility per unit volume is then converted into susceptibility per equivalent of argentic salt produced. This is done by multiplying it by $1/cd$, since this is the volume of one equivalent when d represents the density and c (equal to the mean of c_1 and c_2 in the table) represents the equivalents of argentic salt per gram of the given sample. The expression $(\chi_{OX} - \chi_{UN})/cd$ so obtained is evidently also that for the susceptibility per atomic weight of oxidized silver produced in case this be bivalent, and twice this quantity is this susceptibility in case the silver be trivalent.

(3) Now an expression for the atomic susceptibility in terms of the magnetic moment of the atom is given by the Langevin formula.¹⁷ Namely, the change in susceptibility (in cgs. units) per atomic weight of an element each of whose atoms increases its magnetic moment from 0 to μ cgs. units is by this formula equal to $N_A \mu^2 / 3RT$, where N_A is Avogadro's number. This expression is therefore equated with that for each of the two atomic susceptibilities obtained in (2), these equations are solved for μ , and the values are multiplied by $10^{21}/9.23$ to convert them from cgs. units into Bohr magnetons. There are thus obtained the expressions:

For bivalent silver

$$\mu = \frac{10^{21}}{9.23 N_A} \sqrt{(\chi_{OX} - \chi_{UN}) 3RT/cd}$$

For trivalent silver

$$\mu = \sqrt{2} \times \text{this quantity}$$

The Magnetic Moment of the Argentic Ion.—

From the next to last column of Table V it is seen that the mean value found for the magnetic moment of the argentic ion is 1.98 magnetons if the argentic silver be provisionally assumed bivalent. This is not greatly different from the value 1.92 for the bivalent copper ion, and it

(17) See Van Vleck, "Electric and Magnetic Susceptibilities," Oxford Press, 1932. p. 226.

lies between the values 1.74 and 2.16 found for the solid salts of bivalent silver with pyridine. Moreover, it is only 0.25 magneton larger than the value (1.73 magnetons) predicted theoretically for the magnetic moment arising from the spin of an unpaired electron, such as the bivalent silver and copper ions possess, this excess of 0.25 unit being no greater than may well arise from the contribution of the unquenched orbital moment of the electron. This all points to the conclusion that the argentic silver is in fact largely in the bivalent state.

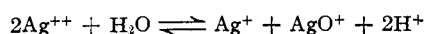
Before drawing this conclusion, however, it is necessary to show that the results are not also consistent with the conclusion that the argentic silver is tripositive. The last column of Table V shows that the measured susceptibilities lead to the magnetic moment 2.81 Bohr magnetons if the argentic silver be provisionally assumed to be tripositive. Now from a theoretical standpoint, as has been already mentioned, there are two possibilities to be considered for trivalent silver: either the eight outer 4d electrons are all paired, or two of them are unpaired. The first case is by far the more probable, because this structure is possessed by bivalent palladium, which is diamagnetic in all of its compounds which have been studied magnetically, including palladous nitrate as crystals and in dilute solution,¹² and because trivalent silver presumably has exactly the same structure as bivalent palladium, in that the number of electrons and types of electron orbits available are just the same in the two atoms. The observed susceptibility for the argentic solutions leads, however, to a large calculated moment (2.81 magnetons) instead of to the value zero, showing that the argentic silver is not present as trivalent silver with the electronic structure that is to be expected for it.

We may however also consider the second, less probable, alternative that the trivalent silver is similar in structure to the bivalent nickel ion, in that it contains two unpaired electrons (4d for silver, 3d for nickel). In this case the contribution of the orbital moment to the susceptibility would be expected to be about the same for the tripositive silver ion as for the bivalent nickel ion. But for the latter ion Cabrera and Duperier¹⁵ have found, by susceptibility measurements of solutions of nickel nitrate, a total moment of 3.19 magnetons, which is 0.38 magnetons larger than that derived for tripositive silver from our measurements.

Molecular State of Argentic Compounds in Acid Solutions

The above-established conclusion that in concentrated nitric acid solutions of the argentic compounds produced by ozone the silver is present almost wholly in the bipoisitive oxidation state is in accord with the above-cited results of Barbieri and others, which showed that organic bases, such as pyridine, precipitate complex salts of bipoisitive silver from argentic solution. This conclusion and these results, however, must be reconciled with the fact also above cited that the solid phases separating from acid solutions contain all the silver in the tripositive state. This reconciliation can be made with the aid of the following hypothesis.

Though argentic compounds in strongly acid solutions contain the silver very largely in the form of the Ag^{++} ion, yet this ion is involved in a rapidly established equilibrium with appreciable quantities of Ag^+ ion and Ag^{+++} ion, which last however probably mainly exists as a hydrolyzed product, such as AgO^+ . Thus, for definiteness it may be assumed, as was done in the preceding article on the reaction rates, that the following reaction quickly attains equilibrium



In that case the mass action expression $(\text{Ag}^{++})^2 = K(\text{Ag}^+)(\text{AgO}^+)(\text{H}^+)^2$ would determine the various concentrations prevailing in the solution.

From this mass action expression it follows that upon diluting an acid argentic solution with water the concentration of AgO^+ would increase until the solution became saturated with some compound produced out of this tripositive silver radical. This tripositive compound is shown to be a basic salt by the fact that it yields upon drying (which results incidentally in partial reduction of the tripositive silver) substances like $\text{AgONO}_2 \cdot \text{Ag}_2\text{O}_3 \cdot 4\text{AgO}$ or $(\text{AgO})_2\text{SO}_4 \cdot 6\text{AgO}$ which contain the acidic constituent of the solution.

The proportion of tripositive silver that as a result of this reaction is present in more dilute (1 to 4 *N*) nitric acid has not been determined by any of the three above-described methods, for these could be used only in presence of concentrated acid (10–15 *N*). In the following article on the argentous–argentic electrode potential, however, evidence is presented (by a fourth method) that the tripositive silver even in 1 to 4 *N* nitric acid is relatively small.

Summary

The argentic salt produced in solution by oxidizing with ozone argentous nitrate dissolved in concentrated nitric acid at 0° has been shown to contain the silver almost entirely in the bipoisitive form. This has been done by three independent methods.

1. The total silver content and the oxidizing power of the steady-state mixture produced in contact with an excess of solid silver nitrate was determined, and from the total silver was subtracted the unipoisitive silver present as determined by measuring the solubility of silver nitrate in the same strength of nitric acid.

2. Total silver and oxidizing power determinations showed that, though 85–90% of the silver is converted (stoichiometrically) into bipoisitive silver by passing 5–8% ozone into a 0.025 *N* solution of silver nitrate in 12 *N* nitric acid, yet the use of even 40–65% ozone results in converting only 98 to 103%.

3. The change in magnetic susceptibility resulting when an argentous solution is partially oxidized was measured, and from this the magnetic moment of the argentic ion was computed. This was found to have a magnitude (1.98 magnetons) that corresponded with that to be expected for the bivalent silver ion (1) from its structural similarity with the bivalent copper ion and (2) from the susceptibilities of solid salts of bipoisitive silver with pyridine previously measured by Sugden. It is shown that the alternative assumption of trivalent silver leads to an experimental value (2.81 magnetons) of the magnetic moment entirely inconsistent with the diamagnetism of the presumably almost identical bivalent palladium ion, and discordant even with the magnetic moment (3.19 magnetons) of the less closely related bivalent nickel ion.

These results are in accord with the well-established fact that organic bases, such as pyridine, precipitate salts of bipoisitive silver from argentic solutions. To account also for the well-known fact that from such solutions water precipitates basic salts of trivalent silver, the hypothesis is proposed that in more dilute acid solutions an equilibrium rapidly establishes itself, such as: $2\text{Ag}^{++} + \text{H}_2\text{O} = \text{Ag}^+ + \text{AgO}^+ + 2\text{H}^+$. This hypothesis has already proved useful in explaining the mechanisms of the silver oxidation and reduction reactions.

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