
JOURNAL OF THE AMERICAN CHEMICAL SOCIETY

VOLUME 55

FEBRUARY, 1933

NUMBER 2

Third Report of the Committee on Atomic Weights of the International Union of Chemistry

BY G. P. BAXTER (*Chairman*), MME. M. CURIE, O. HÖNIGSCHMID,
P. LE BEAU AND R. J. MEYER

The following report of the Committee covers the twelve month period, September 30, 1931 to September 30, 1932.¹

Only two changes in the table of atomic weights are recommended, from 126.932 to 126.92 in the case of iodine and from 138.90 to 138.92 in the case of lanthanum.

Physico-chemical Methods

Carbon.—Moles and Salazar² have determined the normal density of carbon monoxide prepared by four methods: (1) reaction of sodium formate with phosphorus pentoxide, (2) dehydration of formic acid with phosphoric acid, (3) dehydration of formic acid with concentrated sulfuric acid, (4) reaction of potassium ferrocyanide with concentrated sulfuric acid. In each case the gas was subjected to chemical purification and then was fractionally distilled.

THE DENSITY OF CARBON MONOXIDE

Preliminary Series

Method	Globe N-2 608.87 ml.	Globe G 987.52 ml.	Average
1	1.25058	1.25006	1.25032
2	1.25076	1.24984	1.25030
	1.25013	1.24986	1.24999
	1.25130	1.25018	1.25074
	1.25105	1.25024	1.25064
	1.25073	1.25027	1.25050
	(1.25124)	(1.25081)	(1.25102)
	1.25095	1.25090	1.25092
	1.25019	1.24945	1.24982
Average	1.25073	1.25011	1.25042

(1) Authors of papers bearing on the subject are requested to send copies to each of the five members of the Committee at the earliest possible moment: Prof. G. P. Baxter, Coolidge Laboratory, Harvard University, Cambridge, Mass., U. S. A.; Mme. Prof. M. Curie, Institut du Radium, 1 Rue Pierre Curie, Paris Ve, France; Prof. O. Hönigschmid, Sofienstrasse 9/2, Munich, Germany; Prof. P. Le Beau, Faculté de Pharmacie, 4 Avenue de l'Observatoire, Paris VIe, France; Prof. R. J. Meyer, Landshuter Str. 11-12, Berlin W 30, Germany.

(2) Moles and Salazar, *Anales soc. españ. fis. quim.*, 30, 182 (1932).

THE DENSITY OF CARBON MONOXIDE (Concluded)

Method	Final Series		
	Globe N 992.04 ml.	Globe G 987.52 ml.	Average
2	1.24953	1.24980	1.24966
	1.25000	1.25053	1.25026
	1.24941	1.24992	1.24966
	1.24953	1.25046	1.24999
Average	1.24962	1.25018	1.24989
3	1.24920	1.24959	1.24939
	1.24972	1.25039	1.25005
	1.25179	1.25067	1.25123
	1.25003	1.25109	1.25056
	1.25061	1.24997	1.25029
	1.24930	1.25019	1.24974
Average	1.25029	1.24973	1.25001
	1.25013	1.25023	1.25018
	1.25013	1.25023	1.25018
4	1.24966	1.25056	1.25011
	1.24968	1.25105	1.25036
Average	1.24967	1.25081	1.25023
Average of all	1.24990	1.25030	1.25010

This result is in exact agreement with that previously found by Pirc and Moles. With the values 22.414 and 1.00050 for R and $1 + \lambda$, respectively, the atomic weight of carbon is calculated to be 12.006. Because of the small coefficient of deviations from Boyle's law, carbon monoxide is as well suited for precise molecular weight determination as the permanent gases, so that this result may be accepted as a valuable addition to the evidence which has been accumulating recently that the atomic weight of carbon is somewhat higher than 12.00.

Nitrogen.—Batuecas³ has continued the investigation on nitrous oxide, previously reported, by determination of the density at pressures below one atmosphere. The gas was prepared (1) by the reaction of hydroxylamine hydrochloride and sodium nitrite, (2) by the decomposition of ammonium nitrate. Chemical purification was followed by condensation and fractional distillation. The values given in the following table are

THE DENSITY OF NITROUS OXIDE

$P = 506.67$ mm.

Method	Globe G 1007.55 ml.	Globe N-3 772.58 ml.	Average
1	1.9749	1.9741	1.9745
1	1.9750	1.9744	1.9747
1	1.9743	1.9746	1.9745
Average	1.9747	1.9744	1.9746
2	1.9744	1.9744	1.9744
2	1.9738	1.9745	1.9741
2	1.9745	1.9758	1.9751
Average	1.9742	1.9749	1.9745

(3) Batuecas, *J. chim. phys.*, **28**, 572 (1931); *Anales soc. españ. fis. quim.*, **29**, 538 (1931).

THE DENSITY OF NITROUS OXIDE (*Concluded*)

Method	Globe G 1007.55 ml.	Globe N-3 772.58 ml.	Average
$P = 380 \text{ mm.}$			
1	1.9723	1.9721	1.9722
1		1.9719	1.9719
1		1.9706	1.9706
1	1.9724	1.9724	1.9724
Average	1.9724	1.9718	1.9718
2	1.9718	1.9719	1.9718
2	1.9726	1.9730	1.9728
2	1.9722	1.9730	1.9726
Average	1.9722	1.9726	1.9724
$P = 253.33 \text{ mm.}$			
1	1.9695	1.9701	1.9698
1	1.9685	1.9700	1.9692
1		1.9677	1.9677
1	1.9691	1.9691	1.9691
Average	1.9690	1.9692	1.9690
2	1.9701	1.9698	1.9700
2	1.9691	1.9701	1.9696
2		1.9700	1.9700
Average	1.9696	1.9700	1.9699

referred to one atmosphere without correction for the deviation from Boyle's law.

The limiting density is calculated on the assumption that a linear relation holds between pressure and density referred to one atmosphere.

Pressure, atmospheres	Density	Limiting density	$1 + \lambda$
1	1.9804		
$\frac{2}{3}$	1.9746	1.9630	1.00886
$\frac{1}{2}$	1.9722	1.9640	1.00835
$\frac{1}{4}$	1.9694	1.9639	1.00840
	Average	1.9636	1.00854

The calculation of $1 + \lambda$ by linear extrapolation of values for PV found from the expression $L_1/L_0 \frac{P_1}{P}$ gave the following result

Pressure, atmospheres	PV	$1 + \lambda$
1	1.00000	
$\frac{2}{3}$	1.00294	1.00882
$\frac{1}{2}$	1.00416	1.00832
$\frac{1}{3}$	1.00559	1.00839
	Average	1.00851

With the values 1.9804, 22.414 and 1.0085 for the density at one atmosphere, R and $1 + \lambda$, respectively, the molecular weight of nitrous oxide is found to be 44.014 and the atomic weight of nitrogen 14.007.

In an article devoted chiefly to criticism of the report of this Committee for 1931, Moles⁴ applies corrections to the above results (1) for diminution in volume of the globes when filled at low pressures and (2) for adsorption as found by Crespi in work as yet unpublished. The corrected figures are as follows

Pressure atmospheres	Density	Limiting density	$1 + \lambda$	N ₂ O
1	1.98034			
$\frac{2}{3}$	1.97459	1.96309	1.00880	44.001
$\frac{1}{2}$	1.97223	1.96412	1.00829	44.023
$\frac{1}{3}$	1.96946	1.96402	1.00833	44.022
Average		1.96364	1.00845	44.016

Whence $N = 14.008$

Fluorine.—Moles⁵ discusses critically earlier determinations of the atomic weight of fluorine with the final conclusion that the atomic weight of this element is very close to integral, 19.000, and suggests that the higher value recently found by Patterson, Whytlaw-Gray and Cawood, through the density of methyl fluoride, is due to the effect of impurity of methane in the methyl fluoride employed by the latter. Patterson, Whytlaw-Gray and Cawood⁶ have prepared methyl fluoride by their own (Collie's) method (pyrolysis of tetramethylammonium fluoride) and that of Moles and Batuecas (action of potassium methyl sulfate on potassium fluoride) and found that the two samples had identical critical constants within the error of the experiments. A direct determination of the compressibility of methyl fluoride yielded the values 1.0114 at 0° and 1.0087 at 21°. The latter is essentially identical with the value calculated from their experiments with the micro-displacement-balance but the former is much lower than that found by Moles and Batuecas from gas density measurements, 1.018. Batuecas⁷ discusses the calculation of Patterson, Whytlaw-Gray and Cawood's density determinations with the micro-balance and points out that since the methyl fluoride employed by Moles and Batuecas possessed a vapor pressure obeying the law $\log p = -(A/T) + B$ it must have been essentially free from impurity. The Committee feel that the preponderance of evidence is in favor of the lower value for fluorine and see no reason at present to alter the value in current use.

Potassium.—Heller and Wagner⁸ were unable to detect by analysis of potassium chloride from plant sources any concentration of the higher isotope of potassium. This confirms their previous finding.⁹

Selenium.—Hönigschmid¹⁰ has synthesized silver selenide from pure silver and pure selenium. The product was brought to constant weight in

(4) Moles, *Anales soc. españ. fis. quim.*, **30**, 460 (1932).

(5) Moles, *J. chim. phys.*, **29**, 53 (1932); *Nature*, **128**, 966 (1931).

(6) Patterson, Whytlaw-Gray and Cawood, *ibid.*, **129**, 245 (1932); *J. Chem. Soc.*, 2180 (1932).

(7) Batuecas, *J. chim. phys.*, **29**, 269 (1932).

(8) Heller and Wagner, *Z. anorg. allgem. Chem.*, **206**, 152 (1932).

(9) Heller and Wagner, *ibid.*, **200**, 105 (1931).

(10) Hönigschmid, *Naturwiss.*, **20**, 659 (1932).

vacuum. Eleven experiments yielded the ratio $2\text{Ag} : \text{Ag}_2\text{Se} = 0.732081$ and the atomic weight of selenium 78.962. This value agrees with Aston's value and is considerably lower than the current one. Until further details are available no change is recommended in the table.

Iodine.—Hönigschmid and Striebel¹¹ have continued their experiments on the conversion of silver iodide to silver chloride, with precipitated silver iodide, instead of material synthesized from the elements used previously.

Iodine was three times distilled from iodide solutions prepared from similar iodine and was sublimed in an oxygen current over red hot platinum. From this iodine hydriodic acid was prepared by direct synthesis with hydrogen and after solution in water this acid was distilled.

Sample A of silver iodide was precipitated by adding an excess of 0.15 *N* hydriodic acid solution to an acid solution of the purest silver (0.15 *N*). Sample B was precipitated in ammoniacal solution and then an excess of nitric acid was added. Sample C was prepared by adding 0.2 *N* ammoniacal silver nitrate to an excess of 0.2 *N* ammoniacal ammonium iodide. Sample D resulted from adding an excess of 0.16 *N* silver nitrate to 0.16 *N* hydriodic acid.

After washing and drying the silver iodide was fused in an air current containing free iodine and then in pure air in a weighed quartz tube, and weighed. Then the silver iodide was converted to silver chloride by heating in a chlorine current at gradually increasing temperatures up to fusion. Proof was secured that the iodide did not retain iodine and that the conversion was complete. Vacuum weights are given.

THE ATOMIC WEIGHT OF IODINE

Sample	Wt. of AgI	Wt. of AgCl	Ratio AgI: AgCl	At. wt., I
A	14.41889	8.80228	1.638085	126.918
A	10.61180	6.47828	1.638058	126.914
A	13.61047	8.30885	1.638070	126.916
B	17.91554	10.93678	1.638100	126.920
A	14.11519	8.61692	1.638078	126.917
A	14.03900	8.57050	1.638061	126.915
A	13.39032	8.17448	1.638064	126.915
B	11.47497	7.00511	1.638085	126.918
B	13.49506	8.23839	1.638070	126.916
B	14.36421	8.76879	1.638106	126.921
C	10.72744	6.54879	1.638080	126.917
C	8.42456	5.14298	1.638069	126.916
D	14.91865	9.10741	1.638078	126.917
D	12.57197	7.67526	1.638080	126.918
D	8.28549	5.05809	1.638067	126.916
C	13.95958	8.52190	1.638083	126.918
C	9.21692	5.62671	1.638066	126.916
Average			1.638076	126.917

(11) Hönigschmid and Striebel, *Z. anorg. allgem. Chem.*, **208**, 53 (1932).

In view of the concordance of this result and the earlier one of Hönigschmid and Striebel (see report for 1931) the atomic weight of iodine is changed in the table from 126.932 to 126.92.

Tellurium.—Hönigschmid¹² has prepared and analyzed tellurium tetrabromide. From the two ratios $\text{TeBr}_4 : 4\text{Ag}$ and $\text{TeBr}_4 : 4\text{AgBr}$ the atomic weight of tellurium is found to be 127.587.

Krypton and Xenon.—Allen and Moore¹³ have separated krypton and xenon from liquid air residues by means of fractional distillation. The average density of the purest krypton fractions, found with a globe of 22 ml. capacity, was 3.733 and that of the purest xenon 5.887. Using values calculated by Watson in 1910 for the compressibilities of these gases the atomic weights of krypton and xenon are found to be 83.6 and 131.4 with an uncertainty of 0.1 unit in each case. These results, which agree with the recent results of Aston, Watson, and Whytlaw-Gray, Patterson and Cawood (see report for 1931) escaped notice by the Committee in preparing the report for 1931.

Lanthanum.—Baxter and Behrens¹⁴ have determined the atomic weight of lanthanum by analysis of lanthanum bromide. Lanthanum ammonium nitrate which had already been brought to a high state of purity in an earlier investigation was subjected to thirty series of fractional crystallizations. After conversion to bromide this salt was carefully dehydrated in a current of dry nitrogen and hydrogen bromide in such a way that melting of the hydrate was avoided, and the salt was finally fused in an atmosphere of pure hydrogen bromide. After being weighed the salt was dissolved and compared with silver in the conventional way and finally the silver bromide was collected and weighed. The following table gives weights corrected to vacuum. The fractions of lanthanum are numbered in the order of increasing solubility of the double nitrate, No. 371 representing the head fraction of the last series of crystallizations.

THE ATOMIC WEIGHT OF LANTHANUM							
Fraction	Wt. of LaBr ₃	Wt. of Ag	Ratio LaBr ₃ :3Ag	At. wt. of La	Wt. of AgBr	Ratio LaBr ₃ :3AgBr	At. wt. of La
376	4.01090	3.42801	1.170037	138.923	5.96743	0.672132	138.923
379	5.19186	4.43727	1.170057	138.929	7.72475	.672107	138.909
372	6.57727	5.62133	1.170056	138.929	9.78548	.672146	138.931
371	6.38414	5.45612	1.170088	138.939	9.49843	.672126	138.920
372	6.91830	5.91287	1.170061	138.931	10.29285	.672146	138.931
371	6.19359	5.29339	1.170041	138.924			
	Average		1.170057	138.929		.672131	138.923

The average value of the two methods, 138.926, is slightly higher than the value previously found by Baxter, Tani and Chapin, corrected for a new determination of the density of lanthanum chloride, 138.916. The

(12) Hönigschmid, *Naturwiss.*, **20**, 659 (1932).

(13) Allen and Moore, *THIS JOURNAL*, **53**, 2512 (1931).

(14) Baxter and Behrens, *ibid.*, **54**, 591 (1932).

average, 138.92, is probably nearer the truth than that recently given in the International table and the atomic weight of lanthanum is therefore changed in the table from 138.90 to 138.92.

Thallium.—Briscoe, Kikuchi and Peel¹⁵ have redetermined the atomic weight of thallium by comparison of the chloride with silver. The processes of purification consisted in repeated crystallization of thallous sulfate and chloride, prepared from (A) English thallium of unknown origin and (B) German thallium obtained from Westphalian pyrites. The novel feature of the investigation lies in the method of titrating the chloride with silver. A few milligrams (2-3) excess of silver was added in every case to the thallous chloride solution and after standing and settling the supernatant solution (6-7 liters) was decanted, evaporated to small bulk and the silver content estimated by titration with $n/1000$ thiocyanate. Vacuum weights are given in the following table.

THE ATOMIC WEIGHT OF THALLIUM

	Wt. of TlCl	Wt. of Ag	TlCl/Ag	At. wt., Tl
English	2.41969	1.08861	2.22273	204.33
	4.97940	2.24057	2.22234	204.29
	4.90541	2.20686	2.22280	204.34
	4.90351	2.20608	2.22272	204.33
	4.90391	2.20615	2.22283	204.34
	6.27962	2.82533	2.22261	204.32
				204.33
German	5.05837	2.27554	2.22293	204.35
	5.25891	2.36607	2.22264	204.32
	9.53621	4.28987	2.22296	204.35
	10.01985	4.50788	2.22274	204.33
	7.25159	3.26254	2.22268	204.33
				204.34
English A	10.04965	4.52210	2.22234	204.29
	11.55090	5.19723	2.22251	204.31
	10.60755	4.77211	2.22282	204.34
	9.91726	4.46146	2.22288	204.35
	9.53722	4.29023	2.22300	204.36
	11.88154	5.34523	2.22283	204.34
				204.33
English B	9.49327	4.27091	2.22277	204.34
	9.91925	4.46205	2.22302	204.36
	10.43230	4.69329	2.22281	204.34
	10.06308	4.52668	2.22305	204.36
	9.07037	4.08074	2.22272	204.33
				204.35
				204.34
Average of all				204.34

Exception may be taken to the analytical method, which was adopted because of alleged uncertainty in the conventional nephelometric method of comparison. The large bulk of supernatant liquid and washings must

(15) Briscoe, Kikuchi and Peel, *Proc. Roy. Soc., (London)* **A133**, 440 (1931).

have contained many milligrams of dissolved silver chloride, since the excess of silver used was very small. The ultimate fate of this silver chloride is difficult to guess but it is hard to believe that the titration of the excess of silver was not seriously affected by it especially in view of the well-known disturbance of the thiocyanate titration of silver in the presence of silver chloride. In fact it is hard to understand why the excess of silver in the individual experiments was so small as actually found. The effect of this difficulty would be, however, to raise rather than lower the atomic weight of thallium so that it is surprising that the atomic weight found is lower rather than higher than that recently found by Hönigschmid, 204.39.

One experiment in which the nephelometric method was used showed erratic behavior, the final result for the atomic weight of thallium being 204.43.

The ratio of thallium to thalious nitrate also was determined. Buttons of fused electrolytic thallium were polished with chamois, fused in hydrogen and bottled in nitrogen for weighing. Solution in nitric acid followed by repeated evaporation with oxalic acid and fusion of the residue failed to give a product of constant weight, so that the authors consider the method (Crookes) unreliable. The results of two experiments are given in which the final weight of thalious nitrate is employed.

Tl	TlNO ₃	At. wt., Tl
12.08412	15.74910	204.42
12.33736	16.07846	204.46

The results of both methods seem of doubtful value.

Johnson¹⁶ claims that the equal opalescence method of titrating halogen compounds with silver is insensitive and gives erroneous results, and that the silver halide is probably contaminated with adsorption products. A new method of finding the end-point is proposed, by comparing the test solution with standard solutions of the reactants containing known amounts of silver and halide.

Wild¹⁷ has compared the values for $1 + \lambda$ as found (1) by extrapolation of high pressure measurements of PV , (2) from low pressure measurements of PV , and (3) from gas density determinations. In computing the high pressure values the expression $PV = A + Bp + Cp^2 + Dp^4$ was employed. In case of the low pressure and density determinations the relation of PV to P is assumed to be linear.

Observer	HIGH PRESSURE					
	H ₂	N ₂	O ₂	He	Ne	A
Onnes	0.99942	1.00041	1.00096	0.99949	0.99959	
Holborn and Otto	.99938	1.00046	1.00098	.99947	.99952	1.00098
Verschoyle	.99937	1.00049				
Average	.99939	1.00045	1.00097	.99948	.99956	1.00098

(16) Johnson, *J. Phys. Chem.*, **35**, 540, 830, 2237, 2581 (1931); **36**, 1942 (1932).

(17) Wild, *Phil. Mag.*, **12**, 41 (1931).

Observer	LOW PRESSURE					
	H ₂	N ₂	O ₂	He	Ne	A
Chappuis	0.99942	1.00043				
Gray and Burt			1.00097			
Geneva Lab.	0.99935	1.00043	1.00086			
Heuse and Otto	.99942	1.00048	1.00097	0.99948	0.99953	1.00094
Average	.99940	1.00045	1.00093	.99948	.99953	1.00094
DENSITY						
Baxter and Starkweather		1.00040	1.00093		0.99941	1.00107
Final mean	0.99939	1.00044	1.00094	0.99948	.99951	1.00099

Atomic Weights from Isotopes

A comprehensive report of the German Chemical Society by Hahn¹⁸ covers the year 1931.

The question of the conversion factor from the physical to the chemical scale still seems to be unsettled. Birge and Menzel¹⁹ prefer the Mecke and Childs value, 1.00022, to that of Babcock and Naudé, 1.00012. Aston²⁰ has been successful in photographing the lines of O¹⁶O¹⁷ and O¹⁶O¹⁸ of sufficient intensity for comparison with O₂¹⁶. From the relative intensities 1 : 4 : 1072 the relative abundance of O₁₇, O₁₈ and O₁₆ is 0.24 : 1 : 536, a result which seems to support the ratio of O¹⁸ to O¹⁶ found by Mecke and Childs, 1 : 630. Since the accuracy of the mass spectrograph is admittedly no greater than 1/10,000 and the uncertainty of the conversion factor obviously is as large, while the measurement of isotopic ratios has inherent uncertainties, the degree of precision of the determination of atomic weights from mass spectrographic data still seems to be inferior to that of the best chemical methods, although the striking concordance of the two methods in most cases is reassuring.

Recent results obtained by Aston²¹ with the mass spectroscope are given in the following table. The factor used for conversion from the physical to the chemical basis is 1.000125.

Hydrogen.—Urey, Brickwedde and Murphy,²² by spectroscopic examination of the least volatile fractions remaining from the evaporation of large quantities of liquid H₂, have detected the lines corresponding to the principal series of H². Examination of the spectrum of ordinary hydrogen indicated the proportion to be approximately 1 in 4000. Bainbridge²³ finds the mass of H² to be 2.01353 assuming He = 4.00216 and H¹ = 1.00778. The atomic weight of hydrogen is therefore slightly higher than the value for H¹ found by Aston.

(18) Hahn, *Ber.*, **65A**, 1 (1932).

(19) Birge and Menzel, *Phys. Rev.*, **37**, 1669 (1931).

(20) Aston, *Nature*, **130**, 21 (1932).

(21) Aston, *Proc. Roy. Soc. (London)*, **A134**, 571 (1932); *Nature*, **128**, 221 (1931); **128**, 725 (1931); **129**, 649 (1932).

(22) Urey, Brickwedde and Murphy, *Phys. Rev.*, **39**, 164 (1932).

(23) Bainbridge, *ibid.*, **41**, 115 (1932).

	Isotopic weights and percentages				Packing fraction	At wt.
	6	7				
Lithium	91.7	8.3			$\begin{cases} 20 \times 10^{-4} \\ 17 \times 10^{-4} \end{cases}$	6.928
Scandium	45				-7×10^{-4}	44.96
	100				assumed	
Rubidium	85	87			-8.2×10^{-4}	85.43
	75	25			assumed	
Strontium	86	87	88		-8.2×10^{-4}	87.64
	10	6.6	83.3		assumed	
Cesium	133				-5×10^{-4}	132.91
	100					
Barium	135	136	137	138	-6.1×10^{-4}	137.43
	5.9	8.9	11.1	74.2		
Thallium	203	205			1.8×10^{-4}	204.41
	70.6	29.4				
Lead	203	204	205	206	0.1×10^{-4}	207.19
(common)	(0.04)	1.50	(0.03)	27.75		
Lead				20.20	209	
(Katanga)				207	(0.85)	
Lead				206		206.067
(Wilberforce)				93.3	0.1×10^{-4}	
Lead				6.7		
(Norwegian)				207		206.20
Lead (Thorite)				206	0.1×10^{-4}	
				85.9		
				8.3		
				207		207.895
				206	0.1×10^{-4}	
				4.6		
				1.3		
Uranium	238					
	100					

ATOMIC WEIGHTS

1933

	Symbol	Atomic Number	Atomic Weight		Symbol	Atomic Number	Atomic Weight
Aluminum	Al	13	26.97	Molybdenum	Mo	42	96.0
Antimony	Sb	51	121.76	Neodymium	Nd	60	144.27
Argon	A	18	39.944	Neon	Ne	10	20.183
Arsenic	As	33	74.93	Nickel	Ni	28	58.69
Barium	Ba	56	137.36	Nitrogen	N	7	14.008
Beryllium	Be	4	9.02	Osmium	Os	76	190.8
Bismuth	Bi	83	209.00	Oxygen	O	8	16.0000
Boron	B	5	10.82	Palladium	Pd	46	106.7
Bromine	Br	35	79.916	Phosphorus	P	15	31.02
Cadmium	Cd	48	112.41	Platinum	Pt	78	195.23
Calcium	Ca	20	40.08	Potassium	K	19	39.10
Carbon	C	6	12.00	Praseodymium	Pr	59	140.92
Cerium	Ce	58	140.13	Radium	Ra	88	225.97
Cesium	Cs	55	132.81	Radon	Rn	86	222
Chlorine	Cl	17	35.457	Rhenium	Re	75	186.31
Chromium	Cr	24	52.01	Rhodium	Rh	45	102.91
Cobalt	Co	27	58.94	Rubidium	Rb	37	85.44
Columbium	Cb	41	93.3	Ruthenium	Ru	44	101.7
Copper	Cu	29	63.57	Samarium	Sm	62	150.43
Dysprosium	Dy	66	162.46	Scandium	Sc	21	45.10
Erbium	Er	68	167.64	Selenium	Se	34	79.2
Europium	Eu	63	152.0	Silicon	Si	14	28.06
Fluorine	F	9	19.00	Silver	Ag	47	107.880
Gadolinium	Gd	64	157.3	Sodium	Na	11	22.997
Gallium	Ga	31	69.72	Strontium	Sr	38	87.63
Germanium	Ge	32	72.60	Sulfur	S	16	32.06
Gold	Au	79	197.2	Tantalum	Ta	73	181.4
Hafnium	Hf	72	178.6	Tellurium	Te	52	127.5
Helium	He	2	4.002	Terbium	Tb	65	159.2
Holmium	Ho	67	163.5	Thallium	Tl	81	204.39
Hydrogen	H	1	1.0078	Thorium	Th	90	232.12
Indium	In	49	114.8	Thulium	Tm	69	169.4
Iodine	I	53	126.92	Tin	Sn	50	118.70
Iridium	Ir	77	193.1	Titanium	Ti	22	47.90
Iron	Fe	26	55.84	Tungsten	W	74	184.0
Krypton	Kr	36	83.7	Uranium	U	92	238.14
Lanthanum	La	57	138.92	Vanadium	V	23	50.95
Lead	Pb	82	207.22	Xenon	Xe	54	131.3
Lithium	Li	3	6.940	Ytterbium	Yb	70	173.5
Lutecium	Lu	71	175.0	Yttrium	Y	39	88.92
Magnesium	Mg	12	24.32	Zinc	Zn	30	65.38
Manganese	Mn	25	54.93	Zirconium	Zr	40	91.22
Mercury	Hg	80	200.61				

Lithium.—Van Wijk and van Koevinge²⁴ and Nakamura²⁵ from a study of band spectra find a low value for the isotopic ratio of Li^7 to Li^6 , and the corresponding atomic weight 6.89.

Boron.—Elliott²⁶ by a study of the band spectrum of boron monoxide finds the isotopic relationship 3.63 and the atomic weight 10.794.

Lithium, Sodium, Potassium, Cesium.—Bainbridge,²⁷ using a Dempster mass spectrograph, finds the ratio of the abundance of the lithium isotopes to be independent of the temperature of the source of the ions. The abundance ratio was determined in three experiments to be 10.75, 11.28 and 11.51, average 11.18, while Aston finds the ratio 11.

Sodium was found to contain less than $1/3000$ of Na^{21} or Na^{25} and less than $1/800$ of Na^{22} , if any, while potassium was found to be free from more than $1/1500$ of K^{43} , $1/600$ of K^{42} and $1/300$ of K^{40} referred to K^{39} .

Bainbridge²⁸ supports Aston's conclusion that the cesium mass spectrum is simple. The discrepancy between the atomic weights determined by chemical and physical methods is far larger than the apparent error of either.

Zinc.—The mass spectrum of zinc²⁹ obtained with a new form of discharge tube which avoids the presence of zinc hydrides gave no evidence of the isotopes Zn^{66} and Zn^{69} reported by Aston. The atomic weight recalculated from Aston's ratios, assuming the relation of the metallic line to the corresponding hydride to be that of the lines 64 and 65 in Aston's spectra, is found to be 65.33, with the conversion factor 1.00022.

Tellurium.—Additional isotopes of tellurium³⁰ of mass numbers 122, 123, 124 (and 127?) were discovered, the mass numbers and percentages of the complete spectrum being as follows

122	123	124	125	126	127	128	130
2.9	1.6	4.5	6	19.0	?	32.8	33.1

With Aston's packing fraction -5×10^{-4} and the conversion factor 1.00022 the atomic weight of tellurium is calculated to be 127.58, a value in much closer agreement with the chemical value than Aston's, 128.03.

RECEIVED DECEMBER 6, 1932
PUBLISHED FEBRUARY 9, 1933

(24) Van Wijk and van Koevinge, *Proc. Roy. Soc. (London)*, **A132**, 98 (1931).

(25) Nakamura, *Nature*, **128**, 759 (1931).

(26) Elliott, *Z. Physik*, **67**, 75 (1931).

(27) Bainbridge, *J. Franklin Inst.*, **212**, 317 (1931).

(28) Bainbridge, *Phys. Rev.*, **36**, 1668 (1930).

(29) Bainbridge, *ibid.*, **39**, 847 (1932).

(30) Bainbridge, *ibid.*, **39**, 1021 (1932).