structure is viewed along the *b* axis. This projection illustrates very clearly that the external shape of a crystal – in this case the prominence of $\{10\overline{1}\}$ – is a function of the packing of the molecules in the crystal lattice.

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Effective Ionic Radii in Oxides and Fluorides*

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An empirical set of 'effective' ionic radii has been compiled by the use of techniques similar to those of Goldschmidt. The actual radius values were derived from: (1) 1000 experimental interatomic distances and (2) an approximately linear relationship between ionic volume and unit cell volume of over 60 isotypic series of oxides and fluorides. These radii take into account electronic spin state and the coordination of both cations and anions and have been found to reproduce closely the experimental interatomic distances in most oxides and fluorides. The choice of absolute values for anion radii and certain deviations from additivity are important and must be taken into consideration.

Introduction

The traditional sets of ionic radii of Goldschmidt, Barth, Lunde & Zachariasen (1926), Pauling (1927), Zachariasen (1931), and Ahrens (1952) have been used with considerable success by chemists, physicists, mineralogists, and crystallographers alike. Ionic radii have been important to the crystal chemist because structure types and cation coordination numbers are determined principally by cation/anion radius ratios. The ability to substitute one cation for another in a particular structure is largely dependent upon matching ionic radii; thus, good values of ionic radii have been important to the chemist synthesizing inorganic compounds. The crystallographer has found radii helpful in comparing experimental interatomic distances with those calculated for other structures.* These interatomic distances in turn can help the chemist to infer oxidation states.

There are, however, several deficiencies in these traditional sets of radii: (1) they do not reproduce interatomic distances to the accuracy now possible in modern structural analyses; (2) although several papers (Pauling, 1927; Zachariasen, 1931) present techniques for calculating the effect of coordination number on *interatomic distances*, no provision is made for determining

^{*} Contribution No. 1454.

^{*} Goldschmidt's & Pauling's radii and correction factors for coordination number $(CN) \neq VI$ were included in Volume II of Internationale Tabellen zur Bestimmung von Kristallstrukturen (1935). Interatomic distances in numerous inorganic compounds were published in Volume III of International Tables for X-Ray Crystallography (1962) and in Tables of Interatomic Distances and Configuration in Molecules and Ions (1958, 1965).

values of the *radii* in other than octahedral coordination; (3) the effect of variations in electronic spin state on radii has not been included.

This paper presents an extensive table of empirical radii that can be used to closely reproduce average observed cation-anion distances in most oxide and fluoride crystal structures. These radii, which take into consideration both anion and cation coordination and electronic spin state, were derived from over 1000 average interatomic distances. Certain radii were made consistent with regular plots of effective ionic radii (r) or volume (r^3) vs.(1) unit cell volumes for over 60 isotypic series of compounds, (2) electron configuration, (3) coordination number (CN), and/or (4) valence.

Background of ionic radii

Crystal chemists have tried for many years to treat atoms and ions as hard spheres and to calculate atomic and ionic radii that would closely reproduce interatomic distances. Approximate additivity of atomic and ionic radii was noted by the earliest investigators (Goldschmidt *et al.*, 1926; Pauling, 1927; Zachariasen, 1931). However, since the additivity of a given set of radii was not found applicable to all bond types, separate tables of radii were derived for ionic, covalent, and metallic bonding. The history of radii determinations for ionic compounds, treated in detail by Pauling (1927), Ahrens (1952), Slater (1964), and Tosi (1964), will be reviewed here only briefly to point out similarities and differences in the methods.

The first attempt at such determinations was by Bragg (1920) who derived a set of atomic radii which reproduced interatomic distances in both ionic and metallic crystals to an average deviation of about 0.06 Å. Bragg & West (1927) later revised these radii so as to be more in accord with Wasastjerna's (1923) conclusion that $r(O^{2-}) = 1.32$ Å.* Bragg & West arrived at a value of $r(O^{2-}) = 1.35$ Å from close-packed oxygen structures. Goldschmidt et al. (1926) derived a set of radii to be used only in ionic crystals. Goldschmidt's radii were obtained from average interatomic distances using Wasastjerna's radii of $r(O^{2-}) = 1.32$ Å and $r(F^{-}) = 1.33$ Å. Because structural data were scarce, Goldschmidt's table was necessarily limited. However he was able to extend the able to cover radii for which there were no experimental interatomic distances by interpolation or extrapolation of unit-cell dimensions of isotypic series e.g. In^{3+} , Y^{3+} , and Tl^{3+} from the C rare earth oxide structure type. If his radii are based on $r(O^{2-}) = 1.40$ Å, the value generally used by later investigators, they agree well with more recent sets of radii. Goldschmidt was the first to note that radii varied with coordination number and provided a few examples of this variation.

Pauling (1927) calculated a set of radii based on an inverse variation with effective nuclear charge in isoelectronic series of the alkali ions. The radii of the alkali ions were determined from interatomic distances in alkali halides assuming $r(O^{2-}) = 1.40$ Å and $r(F^{-}) =$ 1.36 Å. Pauling prepared two tables of radii: (1) a univalent set from which radii of ions of any charge could be calculated and (2) a table of ionic octahedral crystal radii (ionic radii for multivalent ions). A correction factor was provided to allow calculation of interatomic distances in crystals where the CN of the cation differed from six. However, the coordination of the anion was neglected. Pauling (1928) noted that additivity in the alkali halides alone was poor and introduced extra terms in the expression for interatomic distance to correct for cation-cation and anion-anion repulsion. With these corrections, calculated and observed distances agreed to within about 0.001 Å.

Subsequently, Zachariasen (1931) derived a set of radii from the best available interatomic distances, assuming $r(K^+) = 1.33$ Å and $r(Cl^-) = 1.81$ Å. He calculated univalent radii using Pauling's (1927) method and provided equations with which interatomic distances could be calculated if valence and CN of the cation were known. Zachariasen's radii differed considerably from Pauling's but not from Goldschmidt's corrected radii using $r(O^{2-}) = 1.40$ Å. Although Zachariasen's radii reproduced interatomic distances quite accurately. they seem to have been used less than some of the other sets of radii. Ahrens' or Pauling's radii were probably preferred because of the simplicity of extracting radii directly from a table rather than of calculation from univalent radii with corrections for valence and CN. However, even Zachariasen's table did not allow direct comparison of cation radii when the cation CN was different from 6, nor did it differentiate between high and low spin ions.

The next extensive list of radii was prepared by Ahrens (1952) who revised Pauling's radii by the use of slightly different values for the alkali ions. His method utilized the regularities found by plotting radii vs. ionization potentials. The regular curves obtained by Ahrens were used to estimate the radii of ions for which no values previously existed. Ahrens' table was thus the most complete table for many years and consequently was widely used. However, Ahrens considered only octahedral radii and made no provision for variation of radius with electronic spin state.

More recently, two further sets of radii were introduced. Slater (1964) extended the idea of Bragg (1920) and published a table of atomic radii in which $r(^{VI}O^{2-})$ = 0.60 Å, which reproduced interatomic distances in 1200 metallic, covalent, and ionic solids to a deviation of ± 0.12 Å. These radii have not proved to be particularly useful in oxide and fluoride crystal chemistry. Fumi & Tosi (1964) derived a set of ionic radii for the alkali halides based on (1) a correlation between Born repulsive parameters and the ratio r_c/r_a in alkali halides and (2) the minimum in the LiF and NaCl electron

^{*} Throughout this paper a Roman numeral at the upper left of the element will be used to designate CN. Thus, ^xBa²⁺ refers to the ten-coordinated barium ion. If no symbol for CN is present, six coordination is implied.

density maps. Their most significant conclusion was that the ratio r_c/r_a is much larger than that found in all the other traditional sets of radii. Fumi & Tosi's cation and anion crystal radii are larger and smaller respectively by ~ 0.2 Å than the traditional radii. These radii will be considered in more detail in the discussion.

The advent of accurate structure analysis by the use of Guinier *d*-value data, intensities collected with counter diffractometers, and least-squares refinements of crystal structures has provided a wealth of new accurate cell dimensions and interatomic distances. These data have been used in our derivation of an empirical set of 'effective' ionic radii by an extension of Goldschmidt's (1926) method, with the further assumption that the radii of the anions also vary with CN. This derivation uses approximately 1000 interatomic distances in oxides and fluorides, assumes $r(^{VI}O^{2-}) = 1.40$ Å and r(VIF) = 1.33Å, and takes into consideration both coordination number and electronic spin state of firstrow transition metal ions. Low-spin tetrahedrally coordinated ions and high-spin second-row transitionmetal ions are not included because of their infrequent occurrence. The term 'effective' ionic radii is used to emphasize the fact that these radii are empirical and include effects of covalence in specific metal-oxygen or metal-fluorine bonds. We feel these radii more closely reproduce interatomic distances in solids than the previous sets of radii and will be more useful to those who wish to compare radii rather than interatomic distances.

Methods of determining 'effective' ionic radii

Assumptions

It has already been shown that interatomic distances (and therefore 'effective' ionic radii) depend on CN (Goldschmidt, 1926), electronic spin state (van Santen & van Wieringen, 1952; Blasse, 1965), covalency (Pauling, 1927), and the presence of strong anion-anion or cation-cation repulsive forces (Pauling, 1928). Changes due to CN variation are of the order of 0.0–0.3 Å; those due to change in magnetic spin state are ~0.1 Å. The changes due to covalency effects are unknown but are probably reasonably constant for combinations of cations with anions whose electronegativities are approximately the same. Repulsive forces cause changes in the order of 0.0–0.1 Å.

In deriving the set of 'effective' ionic radii listed in Table 1(a) the following assumptions were made:

(1) The principle of additivity of both cation and anion radii to accurately reproduce interatomic distances is valid if one takes into consideration CN, electronic spin, covalency, and repulsive forces. (Repulsive forces are discussed further in another section.)

(2) The effect of covalency in shortening M-X bonds is comparable in all M-F or M-O bonds (Phillips & Williams, 1965).

(3) With the exceptions noted in assumption 1, radii are independent of structure type.

(4) The radii of both *cation* and *anion* vary with coordination number. (The variation in anion radii with CN is discussed in detail in another section.)

(5) Although individual cation-anion distances vary in a polyhedron, the average cation-anion distance over all similar polyhedra in one structure is constant.

(6) With a constant anion, the volume of the unia cells of isostructural series is proportional to the volume of the cations.

Assumption 5 is not basically different from 6 in that it says an ion can be deformed in any fashion but the total volume of the polyhedron will not change.* The test of assumptions 1-6 is the consistency of radii derived from different structures.

The principle of obtaining radii from relative unit volumes for a particular structure type was originated by Grimm & Wolff (1926) and Goldschmidt *et al.* (1926). Goldschmidt estimated the radii of numerous rare earth ions by comparing cell volumes of rare earth oxides with those of Sc_2O_3 and Y_2O_3 for which interatomic distances were known. A similar procedure was used by others to obtain sets of radii applicable to certain structures: Templeton & Dauben (1954) – rare earth fluorides, chlorides, oxychlorides, and oxides; Geller (1957) – oxide perovskites; Geller & Mitchell (1959) – garnets; Roth & Schneider (1960) – rare earth oxides; Sasvári (1960) – rocksalt, rutile, fluorite, and, antifluorite compounds; Knox (1961) – fluoride perovskites; and Espinosa (1962) – garnets.

Procedure

As a first approximation, anion radii (O^{2-} and F^{-}) with CN=III, IV, and VI were determined by subtracting Ahrens' sixfold radii from interatomic distances in simple oxide structures: III – rutile; IV – corundum and C rare earth; VI – rocksalt. In a second step, cation radii for various CN's were determined. These radii were in turn used to obtain anion radii with CN=II, III, IV, and VIII. The results are listed in Table 2. The values in parentheses represent anion radii obtained at a later stage by repeating this procedure with appropriate refined cation radii, *e.g.* in quartz $r(^{II}O^{2-}) = R_{av}(Si-O) - r(^{IV}Si^{4+}) = 1.607 - 0.26 =$ 1.347 Å. The consistency of these values can be taken as a measure of their validity.

^{*} The approximate linearity of radii-volume plots has been shown by Bertaut & Forrat (1957) for $A_3B_5O_{12}$ garnets, by Whinfrey, Eckart & Tauber (1960), and Brisse & Knop (1968) for $A_2B_2O_7$ pyrochlores, by Gattow (1963, 1964) for A_2BX_4 compounds, by Pannetier & Courtine (1966) for ABO_4 , by Giglio (1963) for ABX_6 , by Sasvári (1960) for AX_2 and by Roth & Schneider (1960) for A_2O_3 . Sasvári also corrected certain radii of Ahrens to comply with this regularity and derived a set of eight-coordinated radii from the fluorite structures. Although Giglio, Gattow, and Pannetier & Courtine plotted r vs. V, we feel that it is more correct to plot $r^3 vs$. V or r vs. A (cubic). The excellent linearity in simple oxide structures (rocksalt, rutile, corundum and C rare-earth) supports this view.

Table 1(a). Effective ionic radii

ION	EÇ	CN '	SP	A-IR	cR	'IR'		ION	EC	CN	SP	A-IR	CR	'IR'
AC+3	6P 6	VI		1.18	••			CS+1	5P 6	XII			2,02	1,88
AG+1	4010	II TV50			.81	.67		CU+1	3D10	II			60	.46
		V			1.26	1.12		CU+2	3D 9	VI		.96	76	43
		ŶI_		1.26	1 29	1,15		007-	~~ ,	V			.79	.65
		VII			1,38	1,24				ΫI		(.72)	87	.73
AG+2	4D 9	- VI VIII		- 89	1,44	1,30		DY+3	4F 9	VI.		,92	1.048	.908 R
AG+3	4D 8	IVSQ	I		.79	.65		ER+3	4F11	VI		- 89	1 021	I.UJ K AA1 R
AL+3	2P 6	ĪV			53	.39	*			ŶĪIJ			1,14	1.00 R
		V V T		- 51	.62	.48		EU+2	4F 7	VI			1,31	1,17
∆M↓ 3	5F 6	VI VI		1-07	1 15	1.01	/ K¥	FUAS	4F 6	VIII		٩A	1,39	1,25
AM+4	5F 5	vī.		92		T 0 0 T	R	LUTU	41 G	VIII			1.21	1.07 R
		VIII			1,09	,95		F -1	2P 6	İI			1,145	1.285
A5+3	45 2 3010	VI TV		(,58)	475	136	- D#			III			1,16	1.30
ne		ŶĪ		.46	64	.50	/ K-			NI TA		1.33	1.19	1.31
AT+7	5010	VI		.62	-	-		F +7	15 2	νī		.08		
	5010 50 B	VI		1,37	94	70		FE+2	30 6	IV	HS		.77	.63
AVY-	~~~~	VI		r .85)	104	• / •				V1		74	.75	.61 E
B +3	1S 2	İİI			,16	.02		FE+3	30 5	ĪΥ	HS		.63	.49 ·≭
		IV		61	26	,12	*		· -	γI	LS		69	55 R
84+2	5P 6	VI VI		1.34	1 50	1 36			<0 2		HS	.64	785	.645 R*
e r		VII		110	1.53	1.39		GA+3	3010	1V 1		1.00	61	47 •
		VIII	,		1 56	1,42		VATE	0010	v.			.69	• • 7/ • -55
		îx			1,61	1.47	-			ΎΙ		.62	760	620 R#
		ŶIJ			1.74	1,52	7	GD+3	4F 7	VI		•97	1,078	938 R
BE+2	15 2	ÎĨĨ			31	17		GE+2	45 2	ÅI T		(.73)	1,20	1,06 R
		IV			41	.27	*	GE+4	3010	IV		• • •	.54	.40 *
BT+3	6S 2	Ϋ́Ι		.35	1 13	90		۱	• 5 0	Ϋ́Ι		.53	680	540 R*
		vı		(.96)	1.16	1.02		П +•	79 U	II			- 04	=,38 = 18
F		VIII			1,25	1,11	R	HF+4	4F14	vī		,78	85	.71 R
BI+5	5D10	VI U		•74	• • •		_	1		VIII			.97	.83
BK+5 BK+4	5F 7	VIII			1,10	,96	R	HG+1	65 1	111			1,11	.97
BR-1	4P 6	ŶĪ		1,96			n.	1046	2010	İv			1.10	- 69 - 96
BR+7	3D10	VI		. 39			1			νī.		1.10	1 16	1 02
C +4	15 2	III		15P	,06	-,08	1	1		VIII		~ 1	1,28	1.14 R
		ŶĨ		.16			I	HQ+3	4110	VI		• 91	1,034	.894 R
CA+2	3P 6	VI		,99	1,14	1.00	I	I -1	5P 6	VI		2.20	1,10	1.02 K
		VII			1,21	1,07		1 +5	55 2	Ϋ́Ϊ		.62	1,09	.95 ?
		17 111			1 32	1,12	*	1 1 +7	4010	VI		,50		-
		х́г_			1 42	1,28		10+0	4010	VIII		•91	1 063	.790 R*
•		XII			1,49	1,35	1	IR+3	50 6	VI			87	-73 7
CD+2	4010	IV			· 98	.84		IR+4	50 5	VI.		68	77	63 R
		vī		.97	1.09	.95	I	K +4	34 0	VI VTI		1.35	1,52	1,38
		VII		•••	1 14	1.00	!			VIII			1.65	1.61 2
		VIII			1,21	1.07	!			IX			1 69	1.55 ?
CE+3	65 1	XII XII		1.07	1 174	1,31				X.,			1 73	1,59 7
4644	0.5 1	VIII		1.0.	1.28	1,14	R	1 443	4010	X1.1 VT		1.14	1 201	1,60 7
		XII			1,43	1,29	ł	6440	4644	vî1		****	1 24	1.10
CE+4	5P 6	VI		.9	94	.80	2			VIII			1,32	1,18 R
CF+3	6D 1	VI			1.09	.97	R			ÎX			1,34	1,20 ?
CL-1	3P 6	VI.		1,81		• - •	<u> </u>	1		XII			1.42	1,28
CL+5	35 2	III			.26	.12	1	LI+1	15 2	ĪV		_	73	59
CL+/	2P 6	IV UT		. 27	, 34	,20		l		VI		•68 •F	.88	.74
CM+3	5F 7	vî		•	1,12	.98	R	LU+3	4F14	VI VTIT		.85	.988	.848 R
CM+4	5F 6	VIII			1,09	.95	Ř	MG+2	2P 6	Ĭv			63	.9/ R .49
C0+2	30 7	γI	LS	72	.79	.65	R			VI_		.66	860	720 R*
C0+3	30.6	νī	15	./.	•0/5 665	6735	R*	MNL 2	30 p	VIII			1,03	.89
	90 0	V #	HS	.63	.75	-61	R	MN+C	30 5	A1	LS HS	- 80	,81 960	.67 E
CR+2	30 4	VI	LŜ	••	87	73	E			VIII	n.a	.00	1_07	.820 K∓ .93 R
Co.3	XD X		HS	41	.96	.82	R	MN+3	3D 4	Ŷ.			72	.58
CR+4	3D 2	TV TV		.00	,755 58	,615 44	R*			VI	LS		.72	.58 R
V 114 ·		ŶĬ			69	.55	R	MNAL	30 3	¥ T	115	(1007	.79	65 R
CR+5	3D 1	İV			490	,350	R	MN+6	3D 1	ΪV		(.41	.540 K
	•• •	VIII			.71	,57	1	MN+7	3P 6	Īv			40	26
CR+D	34 6	1V VT		52	•44	.30		-	~ n •	VI		.46	-	-
CS+1	5P 6	vi		1.67	1.84	1.70		MO+3	40 j 40 j	VI VT		70	.81	.67
-		IX			1,92	1,78	J	MO+5	40 1	Ϋ́Ϊ			:77	.63 E
		x			1,95	1,81		M0+6	4P 6	I۷			56	.42 *

Table	1(a)	(cont.)
Table	1(4)	(00000)

ION EC CN S	SP A-IR	CR	'IR'	ION EC	CN SF	P A-IR	CR	'IR'
M0+6 4P 6 V		.64	.50	5 +6 2P (6 VI	.30		
VI	.62	74	60 *	SB+3 55	2 IVPY		.91	.77
VII		, 85	.71		N ₁	4 761	• 94	•80
N 45 25 2 VI N 45 15 2 TII	(.10)	. 02	12	58+5 4D1		.62	.75	.61
vi vi	.13		•••	SC+3 3P	6 VI	.81	870	,730 R*
NA+1 2P 6 IV		1,13	.99 ?		VIII	4 09P	1_01	.87 R
V	07	1,14	1,00 2	SE-2 4P	6 VI 2 VT	1.90		
VI	4.21	1.27	1.13 7	SE+6 3D1			.43	.29
VIII		1,30	1,16 ?		Ŷİ	.42		
IX		1,46	1,32 ?	5I+4 2P	6 IV	. 4.2	,40 540	•20 ¥ 400 ₽#
NB+2 40 3 VI		.85 84	.71 F	SM+3 4F	5 VI	1.00	1,104	964 R
NB+4 40 1 VI	(.74)	83	69 R		VIII	_ •	1,23	1,09 R
NB+5 4P 6 IV		46	.32 ?	SN+2 55	2 VI	•93	1 84	1 22 B
	.69	•78 80	.64	5N14 4D1		.71	.830	.690 R*
ND+3 4F 3 VI	1.04	1,135	.995 R	SR+2 4P	6 VI	1.12	1 30	1,16
VIII		1,26	1.12 R		VII		1 35	1,21
	60	1,23	1,09 7		VIII		1 46	1,25
NI+2 30 8 VI	LS .07	.70	.56 R		ŶIJ		1 58	1,44
	ĤŜ	74	,60 E	TA+3 5D	2 VI		.81	.67
NP+2 5F 5 VI		1.24	1.10	TA+4 50	1 VI 6 VT	- 68	.80	.65 R
NP+3 5F 4 VI	1,10	1,18	1,04 K		VIII	,	83	.69
VIII	• • •	1,12	.98 R	TB+3 4F	8 VI	.93	1,063	.923 R
NP+7 6P 6 VI	.71		4 75	•0.0 0E	VIII	21	1,18	1.04 R
0 -2 2P 6 II		1.22	1,35		VIII	*0*	1.02	.88
1 11		1 24	1,38	TC+4 4D	зVĪ		78	64 R
VI	1,40	1,26	1.40	TC+7 4P	6 VI	.56		
	.69	1,28	1,42 .630 B			6,24	.66	.52
P +3 35 2 VI	(.44)				IV	_ • •	•	-
P +5 2P 6 IV		.31	.17 *		VI	(.70)		
	.35			TE+0 401	6 VI	1.02	1,14	1.00
PA+5 5P 2 VI	.98				Î VÎL		1 20	1.06
IIIÝ		1,15	1.01		IX		1,23	1,09
PA+5 6P 6 VI	.89	1 10	05	TI+2 50	2 11	(.76)	A1	.67 B
PB-2 65 2 IVPY		1.09	.95	TI+4 3P	6 V	(,,,,,	67	53
VI	1,20	1 32	1,18		VI	.68	,745	.605 R*
VIII		1,45	1.29	TL+1 65	2 VI	1.47	1 74	1,50 K
		1.53	1.39		XII		1 90	1.76 R
XII		1,63	1,49	TL+3 501	LO VI	•95	1,020	.880 R
P8+4 5010 VI	.84	,915	.775 R	TH. 3 (15)		£7	1 000	1.00 K
PD-1 40 9 II		.73	.59 K	1 1M+5 4P	VIII	•07	1,13	.99 R
PD+2 4D 8 IVSQ		78	.64	U +3 5F	3 VĪ		1,20	1,06 R
<u>tv</u>	(.80)	1,00	.86	U +4 5F	2 VI	,97	1.12	. 98
PD+3 40 7 VI PD+4 40 6 VT	.65	.76	62 R		VIII		1,14	1.00 R*
PM+3 4F 4 VI	1,06	1 119	.979 R		IX		1,19	1.05
PO+4 65 2 VIII	· •	1,24	1.10 R	U +5 5F	1 VI		1 10	.70
P0+6 5010 VI	+67 1-06	1,153	1.013 R	U +6 6P	6 II		1.59	45
		1,28	1,14 R	1	IV	~~	.62	• <u>48</u>
PR+4 4F 1 VI	,92	.92	.78 ?		VI	.80	1 0.2	./5 AA
VIII	/ .AĤ1	1,12	. 77	V +2 30	3 VI	(.88)	93	.79
PT+4 50 6 VI	651	,77	.63 R	V ∔3 30	2 VI	(.74)	780	640 R
PU+3 5F 5 VI	1.08	1,14	1.00 R	V +4 30	1 VI	(.63)	.73 40F	.57 K
PU+4 SF 4 VI	,93	.94	.80 7	V +5 3P	0 1V		60	46 *
RA+2 6P 6 VI	1.43	~	1 20	1	VI.	, 59	.68	.54
RB+1 4P 6 VI	(1.47)	1.63	1.49	₩ +4 50	2 VI	,70	.790	.650 R
VII.		1.74	1,60	₩ +6 5P	6 IV VT	.62	.72	.58
XII XII		1,87	1,73	Y +3 4P	6 VI	,92	1,032	.892 R*
RE+4 5D 3 VI	(,72)	.77	.63 R	· ·	VIII		1,155	1.015 R*
RE+5 50 2 VI		,66	.52 7	YD. 3 //E	1X 1X 1X	. 86	.44 .99A	.858 R
RE+6 50 1 VI		, 54	.5∠ .40	¹⁸⁺³ 4 ^r	VIII		1,12	98 R
VI VI	.56	71	57	ZN+2 3D	10 ĪV		.74	.60 *
RH+3 40 6 VI	.68	.805	.665 R	1	V VT	. 74	82	400 .745 R≭
RH+4 40 5 VI		.755	.68	ZR14 4P	6 VI	.79	86	72 R
RU+4 4D 4 VI	.67	760	620 R		VII		.92	.78
5 -2 3P 6 VI	1.84P			1	VIII		.98	.84
5 +4 35 2 VI	(.37)	94	12 +					
1 2 40 24 0 14		, 20		1				

EC – electron configuration, CN – coordination number, SP – electronic spin, A–IR – Ahrens ionic radii (CN = VI) (Ahrens, 1952), CR – crystal radii based on $r(^{vI}F^{-})=1\cdot19$ Å, 'IR' – effective ionic radii based on $r(^{vI}O^{2-})=1\cdot40$ Å.

Table 1(b). References for Table $1(a)^*$

AC+3 VT 8 +3 VI BA+2 VI WANIN,V, AND S.RAMASESHAN, Z.KRIST, 114,200(1960) RA (CL 04)2.3 H2 0 MIGHELLAID, ET AL., ACTA CRYST. 20:A19(1966) RA B2 04 SCHWERINGH.G. ET AL., Z.ANDAG ALLGE.CHEW. 305;241(1960) RA 2N 02 WYCKOFF.R.W.G., CRYSTAL STRUCTURES 1,(1963) BA 0 ZACHARIASEN.W.H., Z.KRIST. 70:139(1980) 0A TI 513 09 AG+1 II PREMITT,C.T. AND R.D.SHANNON, TO 9F PUBLISHED. AG FF 02 WYCKOFF,R.W.G., CRYSTAL STRUCTURES 1.(1963) AG2 0 AG+1 IVS0 BYSTROW,A, AND L.EYEPS, ACTA CHEM,SCAND, 4:613(1950) A62 PG 02 DONOHUFJ, AND L.HELMHOLZ, J.AM,CHEM,SOC, 66:295(194) K A6 C 03 SCATTUPIN,V, AND P.L.BELLON, J. FLFCTROCHEM.SOC, 108,819(1961) A6 D AG+1 V JOST.K.H., ACTA CRYST. 14.770(1961) (AG P 03)X AG+1 VI RODE/H., Z-ANOPG-ALLGEM.CHFM, 267.62(1951) AG 59 F6 KIERKEGAROJF- AND 5.HOLMEN, ARKIV KFMI 23.213(1965) AG MO F 06 WYCKOFF.GM.S. CHY5TAL STRUCTURES 1.(1963) AG F ZACHARIASEM.W.H.Z-ZARIST, 82.HOLMENS 4.69 5 OL AG+1 VII KIERKEGAARD,P. AND S. HOLMEN, ARKTV KEMI 23,713(1965) AG NO P 06 46+1 VIII 541 ¥111 KROBH-MOEJJ., ACTA CRYST.18,77(1965) AG2 B8 013 WELL5×A.F., Z.KRIST. 95,74(1936) AG CO (N H3)2 (N 02)4 4042 V1 AG+3 IV50 SCATTURIN:V. AND P.L.BELLON, J.ELFCTROCHFM.SOC. 108,019(1961) AG+1 AG+3 O2 L-3 IV E-3 IV BROWN-R.E. AND S.W.BAILEY, ACTA (RYST, 17.1301(1964) K AL <13 08 BURNAM-C.W., Z.KRIST, 118.127(1963) AL2 51 05 EULER-F, AND JA, BRUCE, ACTA (RYST, 19.071(1065) M3 AL5 012 MACHATSCHKIFF, Z.KRIST, 04.222(1936) AL A5 0A MACHATSCHKIFF, Z.KRIST, 192,101(1966) AL P 0A SCHWARZEHBACH-0, Z.KRIST, 132,101(1966) AL P 0A SCHWARZEHBACH-0, Z.KRIST, 132,101(1966) AL P 0A STITH-J,V. AND S.W.BAILEY, ACTA (RYST, 17.682(1965) AL-0 VILLTAMS.P.P. AND H.D.HEGAV, ACTA (RYST, 17.682(1965) AL-0 VILLTAMS.P.P. AND H.D.HEGAV, ACTA (RYST, 17.682(1965) ALFTES LIEBAUF, ACTA (RYST, 19.396(1965) LT AL 02 AL+3 V BERTAUT.F. AND J.MARESCHAL: COMPT.REND, 257:667(1963) Y AL 03 BURNHAM.C.W. AND M.J.BUERGER, Z.KRIST. 115:269(1961) AL2 St 05 BURNHAM.C.W. AND M.J.BUENGER, Z.KRIST. 115.269(1961) AL2 ST 05 AL-3 VI BURNHAM.C.W. Z.KRIST. 116.337(1963) AL2 SI 05 ENDERING.T. ET AL., ACTA CRYCT. 221,38(1966) NAL (5 04)2,12 H2 0 ENDERING.T. ET AL., ACTA CRYCT. 221,38(1966) C5 AL (5 04)2,12 H2 0 FALERTIL.E.F. ET ARR.CTA CRYCT. 213,38(1966) AC5 AL (5 04)2,12 H2 0 FALERTIL.E.F. ET ARR.CTA CRYCT. 213,38(1966) ARC AL (5 04)2,12 H2 0 JACK K.H. AND U.S.UTHAN AN HINFGALOTIST 50:2031(1966) MC3 AL2 ST3 017 JACK K.H. AND V.GUTHANN, ACTA CRYST. 4:286(1951) AL F3 MAREZIO.M. AND U.S.PHREIXA J., (FCH.) MAS AL F6 NARAY-52AB0.SI.V. AND K.SSVARI, Z.KRIST. 99,27(1058) M13 AL F6 NEWNHAM.R.E. AND Y.M. DEMAAN. XRIST. 13:7.1255(1962) AL AND 04 PERSEN.B.F. ACTA CHEM, SCAND. 16:421(1962) AL AND 04 PRANDLYM.Z.KRIST. 13:361(1966) AL 2 (3 (51 00)3 SADANAGAIR. ET AL., ACTA CRYST. 15:65(1962) AL& SI 08 BE+2 VI AK+3 VI SHANNON,R.D. AND C.T.PREWITT, TO BE PUBLISHED. B3 VS V (LA F3) 4H+4 VI AM+4 VIII WYCKOFF,R.W.G., CRYSTAL STRUCTURES 1,(1963) AM 02 ACAS VT Ass3 VI Ass3 VI Ass5 IV PINETY.J.J., AM, WINFRALOGIST VARIIGASJ, PR FEP (AS ON)9 (n H)2 FINNETY.J.J., ACTA CRYST, 94711956) (LI AS O3)X UDSTK.H. FT AL, ACTA CRYST, 9140611966) AS2 05,5/3 H2 0 LIEBAUF.ACTA CRYST, 9140611966) AS2 05,5/3 H2 0 LIEBAUF.ACTA CRYST, 9140611966) AS2 05,5/3 H2 0 LIEBAUF.ACTA CRYST, 9140611966) AS2 05,5/3 H2 0 LIEBAUF.ACTA CRYST, 9140611956) AL AS O3 UCKASEVICZ.W., 8ULLACAD.POC.M.SCI.SER.5C.CHIM, 11.50(1961) MG2 AS2 07 MACHATSCHKIJF.I, Z.KRIST. 94-222(1936) AL AS O3 MCONEYR.C.L., ACTA CRYST, 1361194H J AS O3 MCONEYR.C.L., ACTA CRYST, 31(1650) FE3 AS O4 MCONEYR.AD G.SLANGER.Z.KRIST, 124,91(167) ZV2 CU AS2 0A PULETHAK.AND C.CALVO. CAN.J.CHEM, 66437(1963) CU3 AS2 0A SCHULZE.S.E.R.,Z.PHYSTK.CHEW.248125(1941) BA TO4 SHANNON,R.D. AND C.T.PREWITT. TO BE PUBLISHEN. R3 V5 V (ZIECON) BI+5 VI AS+5 VI Ibers, J.A., Acta Cryst. 9,967(1956) K AS ⁶6 Jost, K.H. Et Al., Acta Cryst, 21,800(1966) Asz 05.5/3 H2 0 88-1 VT 88+7 VI AT+7 VI A11+1 VT C +9 IV AU+3 IVSO EINSTEIN+F-W.B. ET AL++ J.CHEM.SOC. \$1967,478 AU F3 C +4 VI AU+3 VI 0 -3 III BLOCK-S. AND A.PERLOFF, ACTA CPYST. 19.297(1965) R4 R4 07 BLOCK-S. AND J.KROGH-MOE. ACTA CPYST. 20,132(1966) CD R4 07 KROGH-MOE-J., ACTA CPYST. 13.689(1967) CS2 86 010 KROGH-MOE-J, ACTA CPYST. 13.70(1965) L2 84 07 KROGH-MOE-J, ACTA CPYST. 18,77(1965) A62 8M 01% KROGH-MOE-J, ACTA CPYST. 18,77(1965) A62 8M 01% KROGH-MOE-J, ACTA CPYST. 16,70(1963) K 85 00 WAREZIO.W. ET AL., ACTA CRYST. 16,79(1963) K2 R4 07.4 M2 0 ZACHARIASEN.W.H. ET AL., ACTA CRYST. 16,1144(1963) ME2 R 03 (0 H) ZACHARIASEN-W.H., ET AL., ALIA CRISI, 10,11-11-11, 207(1365), R. B. O. B +3 IV ELOCK-S. AND A.PERLDFF, ACTA CRYST, 19,207(1365), R. B. O. CLARK-J.R., AMD - MIMERALOSIST 00,150,11061, SS R M (0 M)2,3 H2 O MINARAW, AND J.KCHOH-MOZ, ACTA CRYST, 20,11361, CD M 07 KROGH-MOE-J., ACTA CHEM, SCAND, 16,2055(1964) SR RM 07 KROGH-MOE-J., ACTA CRYST, 13,100(1362) L12 B& 07 KROGH-MOE-J., ACTA CRYST, 13,100(1362) L12 B& 07 KROGH-MOE-J., ACTA CRYST, 13,100(1362) K2 RM 013 KROGH-MOE-J., ACTA CRYST, 13,108(1365) K2 RM 013 KROGH-MOE-J., ACTA CRYST, 13,108(1365) K2 RM 013 KROGH-MOE-J., ACTA CRYST, 13,108(1365) K2 RM 014 KROGH-MOE-J., ACTA CRYST, 13,108(1365) K2 RM 014 KROGH-MOE-J., ACTA CRYST, 10,978(1963) K2 RM 074 H2 O MAREZIO,M. KTAL, ACTA CRYST, 10,978(1963) K2 RM 074 H2 O MAREZIO,M. AND J.P.REWEIXA, J.C.HE-MYTS, 4453361(1963) K2 RM 074 PELOKA, AND S.D.KOK, ACTA CRYST, 10,978(1963) K2 RM 074 H2 O MAREZIO,M. AND J.P.NENETAN, J.P.MYS, J. MASSIGI 1966] LT R 02 PELOKIA, J.P. AND M.MAREZIO J.P.MYS, SCHWAGOLIDS 26,2031(1963) LT R 02 ZACHARIASEN-W.H., ACTA CRYST, 16,385(1963) H B 02 ZACHARIASEN-W.H., ACTA CRYST, 16,385(1963) H B 02

BA+2 VII HILMER¥., ACTA CRYST. 15,1101(1962) BA GE 03 Schnerkoff.g., 2,andrg.allgew.chem. 314,124(1962) BA CD 02 WYCKOFF.R.W.G., CRYSTAL STPUCTURES 3,[1965] BA3 NI 04 BA-2 VIII BLANDJ.A., ACTA (RYST, 14:875(1961) RA2 TI 04 HILMERW, ACTA (RYST, 15:101(1962) BA GE 03 LANDERJ.J., ACTA (RYST, 4:108(1951) BA NI 02 MANDHARM, AND J.RAMASESHAN, 2:KRIST, 119:357(1964) BA (0 H)2.6 H2 0 NARDELLINL AND G.FAVA, ACTA (RYST, 15:477(1962) BA 52 03.H2 0 RADORL, G.S. ET AL., 2:KRIST, 110:23(1958) CU BA (0 H)6.4 H2 0 WYCKOFF, R.W.G., CRYSTAL STRUCTURES 1,(1963) BA F2 BA-2 IX BLOCK-S, AND A.PERLOFF, ACTA CRYST. 19.297(1965) BA B& 07 COLBY:H,Y. AND L.J.BLACOSTE, Z.KRIST 90.1(1935) BA C 03 F05SiO, AND 0.7JDMSLAND, ACTA CHEM.SCAND, 10.284(1956) BA 5 (52 03)2.2 H2 0 F0SSIO, AND 0.7JDMSLAND, ACTA CHEM.SCAND, 11.252(1958) BA TF (52 03)2.2 H2 0 HIGHELL,A.D. ET AL., ACTA CRYST, 20.819(1966) RA H2 DB BA+2 X BLOCK.S. AND A.PERLOFF, ACTA CRYST, 19:297(1965) BA B4 07 BURLEY.G., J.RES.NATL.BUR.STD. 60:23(1954) BA H P 09 NEWHAWAR.E. AND H.D.MESAW, ACTA CRYST, 13.503(1965) BA AL2 512 08 NEWHAWAR.E. AND H.D.MESAW, ACTA CRYST, 13.503(1965) BA AL2 512 08 TEMPLITON.D.H. AND C.H.DAUBEN, U.CHEM.PHYS. 32:1515(1966) RA TI4 09 ZACHARIASEN.W.H., ACTA CRYST, 1:263(1946) RA3 (P 04)2 DA-2 XII EVANS, H.T., JR., ACTA CRYST. 4,377(1951) BA TI 03 HOARD-JL. AND W. VINCENT, J.AM.CHEM.SOC. 52,3126(1940) BA SI F6,BA 6E F6 SAHL,K., BEITA, MINEMAL-PETROG. 9,111(1963) BA S 04 SCHNERINGH.G., Z. ANORG.ALLECH.CHEM. 353:151(967) BA2 M F6 SCHNERINGH.G., ET AL., PHYS.REV. 105:056(1957) BA TI 03 ZACHARIASEN.W.H., ACTA CRYST. 1,263(1948) BA3 (P 04)2 BE+2 III HARRIS,L.A. AND H.L.YAKEL, ACTA CRYST. 22,354(1967) Y2 BE 04
 BE+2 1Y

 BEEYERS,C.A. AND H.LIPSON, Z.KRIST. 82,297(1932)
 BE S ∩ 4,4 H2 O

 BURNSJ,J.H. AND E.K.GORDON, ACTA CRYST. 20135(1966)
 L12 BE F4

 FARRELL,E.F. ET AL., AM. MINERALOBIST 48,8094(1963)
 RE AL2 D4

 GOLOVASTIKOVINI.I., SUITE PHYS.-CRYST. 6373(1962)
 NA RE D O4

 MARRISILA. AND H.L.YAKEL. ACTA CRYST.20,295(1966)
 CA12 BE17 O29

 MROSE.M.E. AND J.A.BUPLENAN.ZKRIST. 117,16(1962)
 AL RE ST 04 O H

 SARTHO.K. ET AL., J.LECTROCHM.SCG.111,78(1964)
 BE 7 ZR F8

 SHITHO.K. ET AL., J.ELECTROCHM.SCG.111,78(1964)
 BE 03 (0 H)
 B1+3 V Arrahams,s.C. et al., J.CHEN.Phys, 47:4034(1967) B112 GE 020 Aurivillius.acta CHEN.5Cand, 18:2375(1964) 812 02 5 04,H2 0 BI+3 VI AURIVILLIUS:B., ACTA CHEM.SCAND, 18:2375(1964) BI2 02 5 04,H2 0 AURIVILLIUS:B. AND 1.JONSSON, ARKIV KEMI 19:271(1962) BI 0 H CR 04 AURIVILLIUS:B. ET AL., ACTA CHEM.SCAND, 18:1555(1966) BI2 6E 05 SEGAL-0.J. ET AL., Z.KRIST, 12:373(1966) BI4 SI3 012 ZEMANN.J., HEIDELBERGER BEITR.MINERAL.PETROG.MITT. 5:139(1956) BI2 M0 06 BI+3 VIII ANRIVILLUS-8+, ARKIV KENT 3,153(1951) BI 58 04 MOOMEY-R.C.L., ACTA CRYST. 1,163(1948) BI AS 04 MOOMEY-SLATERR.C.L., Z.KRIST. 117.371(1962) BI P 04 GURASHI,M.M. AND M-H.BARNES. ANM.HTEALOGIST 38+889(1953) BI V 04 SHANNON,R.D. AND C.T.PREWIT, TO BE PUBLISHED. R3 V5 V (ZIPCONI TOMASMPCJISKIIT. ET AL., SOVIET PHYS-CRYST, 4,713(1965) RI FE 03 BK+3 YI PETERSON,J.R. AND B.B.CUNNINGHAM, INORG.NUCL.CHEM.LETTER5 3,327(1967) 8K2 03 BK+4 VIII PETERSON,J.R. AND B.B.CUNNINGHAM, INORG.NUCL.CHEM.LETTER5 3,327(1967) BK 02 C +4 III CHESSIN,H.,W.C.HAHILTON, AND B.POST, ACTA CRYST, 18,689(1965) CA C 03 C 44 VI CA+2 VI CA+ Construction of the second sec

CA-2 VIII BERTAUT.E.F. AND P.BLUM. ACTA CRYST. 9.121(1956) CA TIZ QA BUBBANK.D., ACTA CRYST. 18.86(1955) CA W QA CLARK.J.R. AND C.L.CHRIST. 2.KRIST. 112.213(1964) CA B3 03.(D H)5.2 H2 O CRUICKSHANK.D.W.J. ACTA CRYST. 17.685(1964) C2 SI QA DECKRENG.F. AND J.S.KASPER. ACTA CRYST. 10.332(1957) CA FE2 QA

Table 1(b) (cont.)

HARRISIL.A. AND H.L.YANEL, ACTA CRYST, 20,295(1966) CA12 BF17 029 HILLIPJ.W. ET AL, ACTA CRYST, 9,981(1955) CA FEP 0a HULLIPJ.W. ET ALL, ACTA CRYST, 559(1963) CA FEP 0a MAREZION.W. ET ALL, ACTA CRYST, 16,590(1963) CA F2 0a MULLER-BUSCHBAUMH.H. H.G.SCHWERING, 7,4A086ALLEFM.CHEM, 336,295(1965) CA SC? 04 PAULOV.P.V. AND N.V.BULOV. SOVIET PHYS.-CHYST, 4,300(1964) CA RE P 0a F PAULOV.P.V. AND N.V.BULOV. SOVIET PHYS.-CHYST, 4,300(1964) CA ME P 0a F PEACOPID.R. AND M.J.BUERGEN, Z.KRIST, 117,331(1962) CA MN J.LEBLOV. SOVIET PHYS.-CHYST, 4,300(1964) MULLER-MUSCHBAUMY, 21,992(1966) CA P 207 WILHELMIN, AND O.JONSSON, ACTA CHFM.SCAND. 19,177(1965) CA 5 0 H (CR 04)3 WYCKOFF.R.M.G. CRYSTAL STRUCTURES 1,1(043) ZALKINJA. AND D.M.TEMPLETON, J.CHEM.PHYS, 40,501(1964) CA W 04 CA+2 IX JOHANSSON:G., ACTA CRYST, 12,522(1959) CA R2 S12 OA POSNERA.S. ET AL., ACTA CRYST, 11,300(1958) CA10 (P 04)6 F2 RODI.F. AND D.BABEL, Z.ANOPG.ALLGEW, CHEW, 336,17(1965) CA IR 03 CA+2 X MAREZIO, M. ET AL., TO BE PUBLISHED CA R2 04 CA+2 XII MAREZIO,M. ET AL., TO BE PUBLISHED CA 92 04 CO+2 IV KOKKORG5,P.A. AND P.J.RENTZEPERIS, Z.KRIST. 119,734(1963) CD 5 04 SCHNERING+N.G.- Z.ANORG.ALLGEM.CHEM. 314+144(1967) 94 CO 02 CO+2 V CALVO.C. AND J.S.STEVENS; CAN.J.CHEM, 46;903(1968) & CD 7N2 (P 08)2 & CALVO.C. AND J.S.STEVENS; CAN.J.CHEM, 46;903(1968) & CD2 7N (P 08)2 & C0+2 VI AJPP,K.L. AND C.CALVO; CAN.J.CHEM, %5,2297(1967) CD2 V2 07 DEWOLFP,P.M., ACTA CRYST, 21,432(1966) CD (0 H)2 LIPSONNH, PROCARD,SOC.L(DNDON)SEPA, 156,467(1936) CD 5 0%,A H2 0 POSNJAK, CAND J.F.W.BARTH, Z,KHIST, M6,271(1930) CD TI 03 WTCKOFF,R.M.G. CAYSTAL STRUCTURES J.(1963) CD 0 WTCKOFF,R.M.G. CAYSTAL STRUCTURES J.(1963) K CD F3 CD+2 VII CALVOIC. AND J.S.STEVENS, CAN.J.CHEM, 46/903(196A) CD 7N2 (P 04)2 CALVOIC. AND J.S.STEVENS, CAN.J.CHEM, 46/903(196A) CD2 7N (P 04)2 CD+2 VIII BRISSE.F., PH.D.THESIS.DALHOUSIE UNIV..HALIFAX.N.S.(1967) CD2 742 07 DRISSE.F., PH.D.THESIS.DALHOUSIE UNIV..HALIFAX.N.S.(1967) CD2 RF2 07 MATKOVIC.B. ET AL., INDRE.CHEM. 4.152(1964) CD (N 03)2.4 H20 WTCKOFF.R.W.G., CRYSTAL STRUCTURES 1.(1963) CD F? CD+2 XII POSNJAK,E, AND R.F.W.BARTH, Z.KRIST. 88,265(1934) CD TT 03 CE+3 VI SLEIGHT,A,W., TO BE PUBLISHED (RE)2 53 TEMPLETONID.H. AND C.H. DAUBEN, J.AMER.CHEM.SOC. 76,5237(1954) CE+3 VIII Mooney.r.C.L., Acta Cryst, 3,337(1950) CE P 04 Shannon.r.D. And C.T.Prewitt, to be published. R3 V5 V (Zircon) CE+3 XII ZALKIN.A. ET AL., J.CHEM.PHYS, 39.2881(1963) CF2 MG3 (N 03)12.24 H2 0 CE+4 VI WTCKOFF,R.W.G., CRYSTAL STRUCTURES 1,11963) BA CE 03 WTCKOFF,R.W.G., CRYSTAL STRUCTURES 1,11963) SR CE 03 CE+4 VIII WYCKOFF,R.W.G., CRYSTAL STRUCTURES 1,(1963) CE 02 CF-3 VI GREEN-JL, AND B.B.CUNNINGHAM, INORG, NUCL, CHEM, LETTERS 3, 343(1967) CF2 03 PETERSON-J.R. AND B.R. CUNNINGHAM, INORG, NUCL, CHEM, LETTERS 3, 327(1967) CF2 03 CI -1 VI CL+5 111 ZACHARIASEN.W.H., Z.KRIST. 71.517(1929) K CL 03 CL+7 IV GOTTFRIED.C. AND C.SCHUSTERIUS, Z.KRIST. 84:65(1932) K CL 04 ZACHARIASEN.W.H., Z.KRIST. 73:141(1930) NA CL 04 CL+7 VI CM+3 VI PETERSON, J.R. AND 8-8, CUNNINGHAM, INORG, NUCL. CHEM. LETTERS 3, 327(1967) EM2 03 CM+4 VIII OFTERSON.J.R. AND B.B.CUNNINGHAM, INDRG.NUCL.CHEM.LFTTER5 3,327(1967) CM 02 CO+2 VI LS COP2 VI US CO22 VI US BAUR, W.H., ACTA CRYST. 11,488(1958) CO F2 BAUR, W.H., ACTA CRYST. 11,190(1964) NA CO2.3 MO3 012 MONTGOMERY.H. ET AL., ACTA CRYST. 22,775(1967) CO (M H412 (5 04)2.6 H2 0 MONTGOMERY.H. ET AL., ACTA CRYST. 22,775(1967) A2 CO (M H412 (5 04)2.6 H2 0 SCHKERING.H.G., Z,ANDEG,ALLGEW, CHEW, 353,131(967) M2 CO (5 SCHKERING.H.G., Z,ANDEG,ALLGEW, CHEW, 353,131(967) M2 CO (5 SCHKERING.H.G., Z,ANDEG,ALLGEW, CHEW, 353,131(967) M2 CO (5 SCHKERING.H.G., Z,ANDEG,ALLGEW, CHEW, 353,131(967) M2 CO (5 SCHKERING.H.G., Z,ANDEG,ALLGEW, CHEW, 353,131(967) M2 CO (5 SCHKERING.H.G., Z,ANDEG,ALLGEW, CHEW, 353,131(967) M2 CO (5 SCHKERING.H.G., Z,ANDEG,ALLGEW, CHEW, 353,131(967) M2 CO (5 SCHKERING.H.G., Z,ANDEG,ALLGEW, CHEW, 353,131(967) M2 CO (5 SCHKERING.H.G., Z,ANDEG,ALLGEW, CHEW, 353,131(967) M2 CO (5 SCHKERING.H.G., Z,ANDEG,ALLGEW, CHEW, 353,131(967) M2 CO (5 SCHKERING.H.G., Z,ANDEG,ALLGEW, CHEW, 353,131(967) M2 CO (5 SCHKERING.H.G., Z,ANDEG,ALLGEW, CHEW, 353,131(967) M2 CO (5 SCHKERING.H.G., Z,ANDEG,ALLGEW, CHEW, 354,131(967) M2 CO (5 SCHKERING.H.G., Z,ANDEG,ALLGEW, CHEW, 354,131(967) M2 CO (5 SCHKERING.H.G., Z,ANDEG,ALLGEW, CHEW, 354,131(962) CO S 04,6 M2 O SCHKERING, SCHKERING, SCHKERING, 21,1962) CO S 04,6 M2 O SCHKERING, SCHKERING, SCHKERING, 21,1962) CO S 04,6 M2 O SCHKERING, SCHKERING, SCHKERING, 21,1962) CO S 04,6 M2 O SCHKERING, SCHKERING, SCHKERING, 21,1962) CO S 04,6 M2 O SCHKERING, SCHKERING, SCHKERING, 21,1962) CO S 04,6 M2 O SCHKERING, SCHKERING, 21,1962) CO S 04,6 M2 O SCHKERING, SCHKERING, 21,1962) CO S 04,6 M2 O SCHKERING, SCHKERING, 21,1962) CO S 04,6 M2 O SCHKERING, SCHKERING, 21,1962) CO S 04,6 M2 O SCHKERING, SCHKERING, SCHKERING, 21,1962) CO S 04,6 M2 O SCHKERING, SCHKERING, SCHKERING, 21,1962) CO S 04,6 M2 O SCHKERING, SCHKERING, SCHKERING, 21,1962) CO S 04,6 M2 O SCHKERING, SCHKERING, SCHKERING, 21,1962) CO S 04,6 M2 O SCHKERING, SCHKERING, SCHKERING, 21,1962) CO S 04,6 M2 O SCHKERING, SCHKERING, SCHKERING, SCHKERING, SCHKERING, SCHKERING, SCHKERING, SCHKER CO+3 VI LS 8LASSE:6., J.INORG. NUCL. CHEM. 27,748(1965) LI CO 02 SHANNON,R.D., INORG.CHEM. 6.1474(1967) R3 V5 V (PFROVSKITE) CO+3 VI HS BLASSErG. J.INORG. NUCL. CHEM. 27.748(1965) K3 CO F6 HEPWORTH.M.A. ET AL., ACTA CRYST. 10,63(1957) CO F3 CR+2 VI L5 CR+2 VI H5 JACK+K-H. AND R. MAITLAND, PROC.CHEM,SOC.LONDON 232(1957) CP F2 SHANNOM,AD. AND C.T. PREWITT. TO BE PUBLISHED, R3 V5 V (RVIILE) STEINFINK.H. AND J.H.BURNS, ACTA CPYCT. 17.823(1964) CP2 F5 CR33 VI DOUGLASS, R.M., ACTA CRYST, 10,42311957) Η CR 02 KNOXK, ACTA CRYST, 13,507(1960) C^R F3 KNOXK, ACTA CRYST, 13,507(1960) C^R F3 NENNAM, R.C. AND Y.E. DEHAAN, Z.RRIST, 117,235(1962) CR2 03 SHANNON, R.D., INDEG, CHEM. 6,1174(1967) R3 V5 V (PFROVSKITF1) STEINFINK-H, AND J.H., TURNS, ACTA CRYCT, 17,873(1964) CR9 F6 VILHELWIK, ACTA CHEM.SCAND, 12,1055(1938) K CR (CR 041) VILHELWIK, ACTA CHEM.SCAND, 12,1055(1938) K CR (CR 041) VICKOFF, W.G., CRYSTAL STRUCTURES 2,119941) CE CR 03

CR+4 IV WILHELMI,K.A., ARKIV KEMI 26,157(1967) SR2 CR 04 CR+W VI CHAMBELAND.B.L., SOLID STATE COMMUN, 5,66311967) SR CP 03 CLOUDIW.H. ET AL., J.APPL.PHYS.,SUPPL, 33:1103(1962) CP 02 SHANNON,R.D. AND C.T.PREWITT, TO BE PUBLISHED, R3 V5 V (RIITLE) CR+5 IV GREENBLAITIN, ET AL., ACTA CRYST, 23,166(1967) CA2 CR ^4 CL Shannonir.D, and C.T.PREWITT, TO BE PUBLISHED, P3 VS V (710CON) WILHELMIK.A, AND O.JONSSON, ACTA CHEM.SCAND, 19,177(1965) CA5 O H (CR 04)3 CR+5 VIII STRONBERG,R. AND C.BROSSET; ACTA CHEM.SCAND, 14,441(1960) #3 CR 08 SHOWELROWS AND CLOUDSEN AND CLOUDSEN AND CLOUDSEN AND CLOUDSEN AND CLOUDSEN AND CLOUDSEN AND CLOUDSEN AND CLOUDSEN AND CLOUDSEN AND CLOUDSEN AND CLOUDSEN AND CLOUDSEN AND CLOUDSEN AND CLOUDSEN AND CLOUDSEN AND CLOW AND CLOUDSEN AND CLOW AND CLUMDRENGS. AND CLOW AND CLUMDRENGS. AND CLUMDRENGS AND CLUMDRENGS. AND CLUMDRENGS AND CLUMDRENGS. AND CLUMDRENGS AND CLUMDRENGS. AND CLUMDRENGS AND CLUMDRENGS. AND CLUMDRENGS AND CLUMDRENGS. AND CLUMDRENGS AND CLUMDRENGS. 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CU FE 02 WYCKOFF.R.W.G., CRYSTAL STRUCTURES 1,(1963) CU2 0 CU+1 VI CUP: 141 CUP: 2145 BACHMANN-G. AND J.2CMANN, ACTA CRYST, 14-747(1961) PR CU 5 04 (0 H)2 BACNTON-G. ET AL.. ACTA CRYST. 11:169(1958) CUP MGR (C 03)4 (0 H)24 ,A H2 0 BUKNOWSKA-STRZYZEWSKAFM., ACTA CRYST. 10:357(1965) CU (H C 0 0)2.2 H2 0 FINNETJJ. AND T.ARAKI! NATURE 107:70(1951) CU (D 000 0 M FLUGEL-XAMLEDAT., ACTA (TA 1577 10 56) CU SC 03.2 H2 0 ANTOWICA. AND J.2CMANN, ACTA CRYST. 11:66(1968) CU SC 000 0 M GHOSE-S., ACTA (CRYST. 16:1105(1962) CU SC 03.2 H2 0 GHOSE-S., ACTA (CRYST. 16:121(1962) CU I 0 3 0 H GHOSE-S., ACTA (CRYST. 16:121(1962) CU I 0 0 0 H2 GHOSE-S., ACTA (CRYST. 16:121(1962) CU I 0 0 H2 GHOSE-S., ACTA (CRYST. 16:121(1962) CU I 0 0 H2 GHOSE-S., ACTA (CRYST. 16:121(1962) CU I 0 0 H2 GHOSE-S., ACTA (CRYST. 16:121(1962) CU I 0 0 H2 GHOSE-S., ACTA (CRYST. 16:121(1962) CU I 0 0 H2 GHOSE-S., ACTA (CRYST. 16:121(1962) CU I 0 0 H2 GHOSE-S., ACTA (CRYST. 16:121(1962) CU I 0 0 H2 GHOSE-S., ACTA (CRYST. 16:121(1962) CU I 0 0 H2 GHOSE-S., ACTA (CRYST. 16:121(1962) CU I 0 0 H2 GHOSE-S., ACTA (CRYST. 16:121(1962) CU I 0 0 H2 GHOSE-S., ACTA (CRYST. 16:121(1962) CU I 0 0 H2 GHOSE-S., ACTA (CRYST. 16:121(1962) CU I 0 0 H2 GHOSE-S., ACTA (CRYST. 16:121(1962) CU I 0 0 H2 GHOSE-S., ACTA (CRYST. 16:120(1963) CU I 0 0 H2 GHOSE-S., ACTA (CRYST. 16:120(1962) CU I 0 0 H2 GHOSE-S., ACTA (CRYST. 16:120(1962) CU I 0 0 H2 GHOSE-S., ACTA (CRYST. 16:120(1962) CU I 0 0 H2 GHOSE-S. GHOSE-S., ACTA (CRYST. 16:120(1951) CU I 0 0 H2 GHOSE-S. CUL2 X ARRANAMS.S.C. ET AL.: J.CHEM.PHYS. 4A.261911968) CU MO 04 FLUGEL-KAHLERIE.: ACTA CHYST. 16:1109(1965) CU2 O(5 04) HERITSH.H. Z.KRIST. 99:166(1935) CU2 O(4 A 5 04 HERITSH.H. Z.KRIST. 102:111339) CU2 O(4 P 04 POLLSEN.S.J. AND C.CALVO CAN.J.CHEM. 46:917(196A) CU3 A52 08 POILSEN,S.J. AND C.CALVG, CAN.J.CHEM. 46-41/11961) (US A2 VS CU-2 VT BACHMANN,H.G. AND J.ZEMANN, ACTA CPYST. 14-747(1961) (PR CU S 04 (0 H)2 BILLY-C. AND H.H.HARDLER, J.M. CHEJ.SC. 74-1094(1967) (UJ F2 BUKONSKA-STRZYZEVSKAM, ACTA CRYST. 19-357(1965) (U H C 0 0)2,7 H2 0 FINETJ.J. ACTA CATST. 21-437(1966) (UZ (AS 0A) (O H). H2 0 FINETJ.J. ACTA CATST. 21-437(1966) (UZ (AS 0A) (O H). HIGTIT.T. BULCST. 94-461(1535) (UZ (AS 0A) (O H). HIGTIT.H. BULCST. 94-461(1535) (UZ (O H) (AS 04) HIGTIT.H. ADV.ASI. 94-461(1535) (UZ (O H) (AS 04) HIGTIT.H. ADV.ASI. 94-461(1535) (UZ (O H) (AS 04) HIGTIH.H. ADV R.G.CANALD.ACTA CATST. 14-101(1961) (U (A H)? KNOXKK, J.CHEM.PHYS. 30-991(1959) K2 (U F4 NOWACKYW. AND R.G.CHEDEGGER, HELV.CLIN.ACTA 58) TCHE 20, (UZ (O H)3 N 03 RAO,R.V.G.SONDRA. ET AL.-7.X81575110/21/5 08) (U RA? (C 0 O H)6.4 H? 0 RAOGRAW.AATA (CTAT. 14-751164) NU IS 04 RAOGRAW.AATA (CTAT. 14-751164) (U IS 04) (U F3 DY+3 VI TEMPLETON+D.H. AND C.H.DAUBEN, J.AM.CHEM.SOC. 76,5237(1954) DY2 03 DY+3 VIII ARRAHAMSISIC, AND JLLBERNSTEIN, JCHEM,PHYS, 46,3776(1967) DY MN2 05 EULERIF, AND BRUCEJJAI, ACTA CRYST, 19,971(1965) DY3 FF2 FF3 012 SWANNON.R.D. AND C.T.PREWITT, TO BF PURLICHFD, R VS A (PYROCHLORF) ER+3 VI FERITA. BULL-SOC.FRANC.WINERAL.CRIST. 85.267(1962) ER2 03 TEMPLETON.D.H. AND C.H.DAUMEN. J.AM.CHEM.SOC. 76.5237(1954) ER2 03 ER+3 VIII KNOP+0. (*5) VII KNOPJG. ET AL., CAN.J.CHEM. 43,2812(1965) FR2 TIP 07 PATSCHEKE.F. ET AL. CHEM.PHYS.LETTERS 2,497(1968) FR M 04 SHANNON.H.G. AND C.T.PREMITT. TO RF DURLISHED. R VS A (PYROCHLORE) EU+2 VI WYCKOFF,R.W.G., CRYSTAL STRUCTURES 1,(1963) EU 0 EU+2 VIII RAUJR.C., ACTA CRYST. 20,716(1966) EU3 04 WYCKOFF,R.W.G., CRYSTAL STRUCTURES 1,(1963) EU F? EU+3 VI RAU/R.C., ACTA CRYST. 20,716/1966) FU3 04 TEMPLETONIDIH, AND C.H.DAUBEN, J.AM.CHEM.SOC, 76,5237(1954) EU2 03

EU+3 YIII MAREZIO:M, ET AL., J.CHEM.PHYS, 48:1094(1968) EU3 FE2 6A3 012 SHANNON:R.O. AND C.T.PREWITT, TO RE PUBLISHED. R VS & IMYROCHLORE) TEMPLETONID.H. AND A.ZALKIN, ACTA CRYST, 10:762(1965) FU2 (W 04)3

EFFECTIVE IONIC RADII IN OXIDES AND FLUORIDES

Table 1(b) (cont.)

F -1 II HG+1 III GRDENIC.D. J.CHEM.SOC. 1312(1956) H82 (N 03)2.2H2 0 F -1 III H9+2 II Aurivilius,k., Acta Cryst. 9,685(1956) H6 0 Aurivilius,k., Acta Chem.Scand. 18+1385(1968) H6 0 Roth,w.L., Acta Cryst. 9,277(1956) H6 0 F -1 IV F -1 VI F +7 VI H6+2 IV BONEFACIC+A., ACTA CRYST, 14,116(1961) H6 5 04,H2 0 Kokkorgs-P.A. AND P.J.RENTZEPERIS, Z.KRIST, 119-234(1963) H6 5 04 FE+2 IV HS HANISCH+K.NEUES JAHRB.MINERAL.MONATSH. 128+362(1966) H FE2 AL8 SI4 024 HG+2 VI BONEFACIC, A., ACTA CRYST. 14,116(1961) HG S 04.H2 0 FE+2 VI LS FEF2 VI L3 FEA2 VI L3 BAUR:4.H., ACTA CRYST, 11:4886(1958) FE F2 BAUR:4.H., ACTA CRYST, 15:615(1962) FE S ON,4 H2 O BURNHAH,C,Y., CARNEGTE INST,VASH,YEAR BOOK 65:285(1966) FE SI O3 61885:6(9,V; ET AL, M.M.MINERALOGIST, IN PRESS. FF2 SI O4 HAMILTON.H.C,Y., ACTA CRYST, 15:353(1962) FE SI F6.6 H2 O HAMILTON.H.C, ACTA CRYST, 15:353(1962) FE SI F6.6 H2 O HAMILTON.H.C, ACTA CRYST, 15:050(1965) FE F53 RUMANOVA:I.H., HUNZANAENSKAYA, SOVJEMYS.-CRYST, 5:650(1961) CA2 FE (P O8)2.H2 O SHTIH-O.K.E TAL., ACTA CRYST, 18:787(1965) FE SI O4 STRUMZCH.AK ET AL., ACTA CRYST, 19:7205(1965) FE GE IO H16 UCK/DFF,R.K.S., CRYSTAL STRUCTURES 2:(1963) K FE F3 ZEMANNIJ., NUEVES JAHRB. MINERAL,MONATSH. 67(1959) FE GF (O H16 HG+2 VIII Shannon,R.D. And C.T.PREWITT, TO BE PUBLISHED. R VS A (PYROCHLORE) SLEIght,A.W., TO BE PUBLISHED. H32 M2 O7 WYCKOFF,R.W.G., CRYSTAL STRUCTURES 1,(1963) H6 F2 H0+3 VI FERT,A., BULL.SOC.FRANC.MINERAL.CRIST. 85,267(1962) H02 03 TEMPLETON.D.H. AND C.H.DAUBEN, J.AM.CHEM.SOC. 76,5237(1954) H02 03 H0+3 VIII Shannon,R.D. and C.T.PREWITT, TO BE PUBLISHED. R VS A (PYROCHLORE) 1 -1 VT I +5 VI WYCKOFF.R.W.6., CRYSTAL STRUCTURES 2,(1964) CS I 03 . WYCKOFF.R.W.6., CRYSTAL STRUCTURES 2,(1964) RB I 03 FE+3 IV MS BATTIA. AND B-POST, ACTA CRYST, 15,1268(1962) V3 FE5 012 BERTAUT, E,F ET AL., ACTA CRYST, 12,149(1959) C42 FE2 05 BERTAUT, F. ET AL., COMPT.AEND, 257,421(1963) NA FF 02 EULER, F. AND J.A. BRUCE, ACTA CRYST, 19,071(1965) M3 FES 012 GELLERNS. AND M.A. GILLEO. ACTA CRYST, 10,239(1957) V3 FES 012 ROMERSIC. ET AL., ACTA CRYST, 24,766(1967) NA3 FE5 09 WEIDENRGRNER, J.E., ACTA CRYST, 14,1051(1961) G03 FE5 012 1 +7 VI IN-3 VI IN-3 VI BERGERNOFF-6. AND H.KASPER. ACTA CRYST. B24-386(1968) CU2 IN2 05 JOHANSSON-6., ACTA CHEM.SCAND. 15,1437(1961) IN 0 H 5 04 (H2 0)2 MAREZIO.H., ACTA CRYST. 210-723(1966) IN2 03 MOONEY-N.C.L., ACTA CRYST. 91,113(01)661) IN P 04,2 H2 0 MOONEY-SLATER.R.C.L., ACTA CRYST. 14,1140(1961) IN P 04,2 H2 0 REID.A.F. INORG.CHEM.S.6.53(1647) LA UP 044,300(1960) R3 V5 V (MN2 0X) ROMAN.F.N.O. AND C.T.REWITT. VO BE PUBLISHED. IN2 03 FE+3 VI LS BLASSE.g., J.INORG.NUCL.CHEM. 27,748(1965) SHANNON.R.D. AND C.T.PREWITT, TO BE PUBLISHED. R3 VS V (K3 MIII (CN)6) SUBMILITY IN THE STATE S FE+3 VI IN+3 VIII Shannon,R.D. and C.T.PREWITT, TO BE PUBLISHED. R VS & (PYROCHLORE) IR+3 VI Hepworth,M.A. et al., Acta Cryst. 10,63(1957) IR F3 IR+S VI Babel-d. Et al., Z.Anorg.Allgem.CHEM, 347/282(1966) CA2 IR 04 Rodi, F. And D.Babel, Z.Anorg.Allgem.CHEM. 336,17(1965) CA IR 03 Snannon.R.D. And C.T.PREWITT, TO BE PUBLISHED. R VS A (PYROCHLORE) K +1 VI +1 VI BODEH, AND E-VOSS, ZIANORG.ALLGEN.CHEN, 264,144(1951) K SB F6 MOPPER, AND M.SABROMSKY, ZIANORG.ALLGEN.CHEN, 339:144(1965) K SC 02 STANLET:G. ACTA (CRYST, 9:07(1956) K S2 06 WYCKOFF.R.W.G., CRYSTAL STRUCTURES 1,(1963) K S 06 ZACHARIASEWI.H.H., ACTA (CRYST, 7:782(1954) K S U 02 F5 ZACHARIASEWI.H.H., ACTA (CRYST, 7:782(1954) K S U 07 ZIEGLER.0:C. ZIARIST, 94:491(1356) K N 02 FR+1 VI 6A+3 IV ARRANAMS,S.C. ET AL., J.CHEM,PHYS. 42,3957(1965) 6A FE 03 EULER+5, AND J.A.DRUCE, ACTA CRYST, 19,971(1965) M3 0A5 012 GELLER+5, J.CHEM,PHYS, 33,676(1960) 6A2 03 MAREZIO,M., ACTA CRYST, 10,461(1965) L1 6A 02 MOOMET,R.C.L.+ ACTA CRYST, 9,728(1956) GA P 04 K +1 VII K +1 VII EAMES; C.O. AND H.H.ONDIK, ACTA CRYST, 15,1280(1962) LI K2 P3 09,42 0 EUMARDS; C.A., Z.KRIST, 80,154(1931) K N 03 LINGWISS, A.T. 200 H.H.OTISELL, ACTA CRYST, 15,10,406(1957) K2 52 05 LINGWISS, ATH ND H.H.OTISELL, ACTA CRYST, 10,406(1957) K2 52 05 ZACHARTARENNH, J. J.CHEM, HYTS, 5191(1957) K B 02 ZACHARTARENNH, J. J.CHEM, HYTS, 5191(1957) K B 02 ZHUKOVA, L.A. AND Z.G.PIMSKER, SOVIET PHYS,-CRYST, 9,31(1964) K2 CR2 07 GA+3 V SHANNON,R.D. AND C.T.PREWITT, J.INORG.NUCL.CHEM. IN PRESS. IN GA 03 GA+3 VI 443 VI BREWERFF.M. ET AL,, J.INORG.NUCL.CHEM. 9,55(1959) GA F3 EULERIF. AND J.A.BRUCE. ACTA CRYST. 19,971(1965) M3 GA5 012 GELLERIS. J.CHEM.PHYS.336/65(1960) GA2 03 MAREZIGIM. AND J.F.REMEIKA. J.PHYS.CHEM.SOLID5 266,1277(1965) GA MORESTLATERRIC.L. ACTA CRYST. 20,326(1965) GA P04,214 20 MORESTH.S. AND A.ROSENZKEIGN. ACTA CRYST. 15,678(1965) GA NN 04 WICKOFF.M.G.F. (ATSTAL STMUCTURES 2,11064) CE GA 03 ZHAKOVALLA. AND Z.G.PINKER, SOVIET PHYS.-ERYST. 9.31(1960) K2 CR2 07 K 1.VIII. NDERESSNIS. AND A.D.WADSLEY, ACTA CHEM.SCAND. 15.46311961) K2 CR2 07 MELTARE. AND M.C.BARZNIGER, J.M.CHEW.SCAND. 15.46311961) K2 TI2 05 MELTARE. AND M.C.BARZNIGER, J.M.CHEW.SCA. 79.31611957) K 5 03 (N H 0 H) CHRISTIC.L. ET AL., ACTA CRYST. 7.401(1954) K V 03.H2 0 FORRESTER.J.D. ET AL., ACTA CRYST. 16.58(1965) K H2 F3 LINGOVIS.C. AND H.M.ONDIK. ACTA CRYST. 16.58(1965) K H2 F3 LINGOVIS.C. ACTA CRYST. 13.443(1965) K RE 04 MOROWIJ.C. ACTA CRYST. 13.443(1966) K RE 04 SIEGEL.S., ACTA CRYST. 13.443(1966) K RE 04 SIEGEL.S., ACTA CRYST. 13.443(1966) K RE 04 SIEGEL.S., ACTA CRYST. 13.443(1966) K RE 04 SIEGEL.S., ACTA CRYST. 9.443(1966) K RE 04 SIEGEL.S., ACTA CRYST. 7.433(1966) K RE 05 ZACHARIASEN.W.H., ACTA CRYST. 7.783(1954) K3 U 02 F5 ZACHARIASEN.W.H., ACTA CRYST. 7.783(1954) K3 U 02 F5 ZACHARIASEN.W.H., ACTA CRYST. 7.782(1954) K3 U 07 ZACHARIASEN.W.H., AND H.A.PLETTINGER, ACTA CRYST. 16.376(1963) K B5 08.4 H2 0 GA2 03 BD+3 VI TEMPLETON+D.H. AND C.H.DAUBEN+ J.AM.CHEN.SOC, 76+5237(1954) GD2 03 6D+3 VIII BRISSEFF., PN+D.THESIS,DALHOUSIE UNIV.,HALIFAX;N.5.(1967) 6D2 TI2 07 EULER:F, AND J.A.GRUCE, ACTA CRYST, 19:071(1965) 6D3 AL2 AL3 012 SNANNON:R.D. AND C.T.PREWITT, TO BE PUBLISHED. R V5 A (PYROCHLORE) WEIDENBORRENJLE. ACTA CRYST. 19:105(1061) 8D3 FF5 012 6F42 V1 ZALMARIASCHIPHA AND MAARLIIINDEN AND SALANDEN AND SALANDARIASCHIPHA AND MARIALIIINJASCHIPHA 1 1X RAAJINDAKSHANIC, Z.KRTST, 111,35(1958) K CL OS BALZJD. AND K.PLIETH. Z.ELKTROCHEM, 59,553(1955) K 2 MT F4 BODE.H. AND G.TEUFER. ACTA CRYST, 9,229(1956) K 2 MF F6 BODE.H. AND G.TEUFER. ACTA CRYST, 9,229(1956) K 2 MF F7 DONDMUE.J. AND CL.M.KER. ACTA CRYST, 20,220(1966) K 2 MF F7 DONDMUE.J. AND CL.M.KER. ACTA CRYST, 20,220(1966) K 2 MF F7 KNOXK., J.CHEM.PHTS, 30,991(1959) K 2 CU F6 KNOXK., J.CHEM.PHTS, 30,991(1959) K 2 CU F6 KNOKK., J.CHEM.PHTS, 18,1086(1965) K 2 MO G4 SCHMEIDER.W., ACTA CRYST, 18,1086(1967) K 2 MO G4 SCHMEIDER.W., ACTA CRYST, 18,1086(1967) K 2 MO G4 SCHMEIDER.W., ACTA CRYST, 18,734(1967) K 2 MO G4 ZACHARIASEN.W., ACTA CRYST, 18,239(1958) K 2 CO G4 ZACHARIASEN.W., ACTA CRYST, 18,239(1958) K 2 CO G4 ZACHARIASEN.W., AND G.E.ZIEGLER.Z.KRIST, 50,164(1931) K2 CR G4 GE+4 IV ABANAMS/S.C. ET AL., J.CHEM.DHYS. 47:4034(1967) B112 BF. 020 DURIF-A., ACTA CRYST. 9:533(1956) GE U 04 GIVETTI,Y., BULL-SOC.CHIM.BELGES 63:200(1954) TILMER*W. ACTA CRYST. 15,101(1962) B4 GE 03 HILMER*W. ACTA CRYST. 15,101(1962) A4 GE 03 HILMER*W. AND GLUNGREW. ACTA CRYST. 25:57(1967) NA2 GE 03.6 M2 0 LAMESNAF.K. ET AL., ACTA CHEM.SCAND. 21.1281(1967) NA8 GH4 GE10 030 (0 H)4 ROBBINS-C. ET AL., ACTA CHEM.SCAND. 21.1281(1967) NA8 GE4 09 SMITH-6.5. AND P.B.ISAACS, ACTA CRYST. 17,842(1964) GE 02 CE44 VI BAUR.W.H., ACTA CRYST. 9.515(1956) GE 02 HOARD-JL. AND W.B.VINCENT.J.AM.CKEM.SDC. 61,2849(1939) K2 GE F6 INORI.W. AND G.LUNDOREN. ACTA CHEM.SCG.MD. 17.617(1963) NAN 026 020 ROBEINS.C. ET AL. J.MES.WATL.BURSID. 703.385(1966) URB GE 03 STEUNZ.H. AND M.GIGLIO. ACTA CRYST. 14.205(1961) FE GE (0 H)6 ZEMANN.J., NEUES JAHRB.WINERAL.MONATSH. 67.(1959) FE GE (0. H)6 K +1 X BYSTROWAA. ET AL., ARKIV KEMI #,175(1952) K SB F4 CAMMILLOFF. ET AL., ACK CHIST 21:200(1966) K NA3 FE2 T12 (S14 012)2 EXEMPLA. C. Z.KRISTROM CHIST 21:200(1965) K S2 03 PALENIK.J., INORGACHEME, 6:507(1967) K 2M 04 ZACHARIASEN.W.H. AND G.E.ZIEGLER, Z.KRIST, 80,164(1931) K2 CR 04 H +1 I +1 I Abrahamsis.C., J.CHEM.Phys. 36/56(1962) CU F2.2 H2 O Abrahamsis.C., J.CHEM.Phys. 44/2230(1964) F53 P2 O64 H2 O Cromerol.T. ET AL., ACTA CRYST.21331(366) ALUM Kuipers, ET AL., J.CHEM.Phys. 25/275(1956) HF Pimentel.Ge. And Al..ME Clellan. The Mysober Bond, Reinhold (New York) 1960 LACHARALSENTUIN, AND OILLIELELER, INAILIST OUTGOTISTIT AL CALO & 1 XII BODE, A. AND H.V.DOHREN, ACTA CRYST, 11.80(1958) K TA F6 BODE, A. AND H.V.DOHREN, ACTA CRYST, 11.80(1958) K NB F6 ELLINGER, F.H. AND ZACHARIASENNU, J. JPHYS, CHEM, 50:051(955) K PU 02 C 03 EVANSH, T., VR.F. AND SACHARIASENNU, J. JPHYS, CHEM, 50:051(955) K V3 08 gOTTFRIED.C. AND C. SCHUSTERUS, ZIKTST, 84.65(1952) K V3 08 GOTTFRIED.C. AND C. SCHUSTERUS, ZIKTST, 84.65(1952) K V3 08 HEPMORTH.M.A. ET AL., J, HNORG, HUCL, SCHU, SCHUB, SCHOOL, SCH 65 F6 HORROJ, A. ATTA CRYST, 9.66(1)55) K AS F6 HORROJ, A. ATTA CRYST, 9.66(1)55) K AS F6 ITTT, ET AL., ACTA CRYST, 9.66(1)55) K AS F6 SIEGEL, SA, ACTA CRYST, 5.683(1952) K AC SIEGEL, SA, ACTA CRYST, SCHUBSINGEN K CA SUBJOEN, P., ACTA CRYST, SCHUBSINGEN K CA SUBJOEN, ACTA CRYST, SCHUBSINGEN K CA WILHELMIK, A. ACTA CHYST, SCHUBSINGEN K CA WILHELMIK, A. ACTA CRYST, STAJSISSI K TA 03 WILHELMIK, A. GA, CRYSTAL STRUCTURES 3,(1965) K AL3 (0 H)6 (S 04)2 H +1 II ELLISON,R.D. AND H.A.LEVY, ACTA CRYST, 19;260(1965) K H (C2 H 04 CL) MAMILTON.W.C. AND J.A.IBERS, ACTA CRYST, 16:1209(1963) H CR 02 HC DONALD.T.R.R., ACTA CRYST, 13:13(1960) (W H3) H F2 HC GAVNBLL, AND J.A.IBERS, J.C.HEM,PHYS, 39:2677(1963) NA H F2 PETERSON.S.W. AND H.A.LEVY, J.CHEM,PHYS, 20:704(1952) K H F2 PETERSON.S.W. AND H.A.LEVY, J.CHEM,PHYS, 29:704(1952) K H (C2 H2 04) HF+4 VI BRISSE,F., PM.D.THESIS,DALHOUSIE UNIV.,HALIFAX,N.S.(1967) M2 HF2 07 SHANNON.R.D. AND C.T.PREWITT, TO BE PUBLISHED. R VS A (PYROCHLORE) HF+4 VIII WYCKOFF.R.W.G., CRYSTAL STRUCTURES 1,(1963) HF 02 LAHS VI

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Table 1(b) (cont.)

SILLEN, L.G. AND K, LUNDBORG, Z.ANORG, ALLGEM. CHEM, 252,2(1943) LA2 MO 06 TEMPLETON, D.H. AND C.H.DAUBEN, J.AM.CHEM.SOC. 76,5237(1954) LA2 03 LA+3 VII GUILLEN.M. AND E.F.GERTAUT, COMPT.REND. SER.A.B 2628,962(1966) LA2 TI 05 LA+3 VIII BRISSE+F., PH-D.THESIS.DALHOUSIE UNIV.,HALIFAX,N.S.(1967) LA2 M2 D7 SHANNON,R.D. AND C.T.PREWITT, TO BE PUBLISHED. R VS A (PTRACHLORE) LA+3 IX Hunt-6.8.,JR., ET AL., ACTA CRYST. 7,106(1954) LA2 (S 04)3.9 H2 D ZALKIN'AA, ET AL., 1MORG.CHEM, 5.1466(1966) LA F3 LA+3 X LONGO+J.M. AND A.W.SLEIGHT, INDRG.CHEM. 7+108(1968) LA4 REA 019 LA+3 XII HUNT+E-B.+JR.+ ET AL.+ ACTA CRYST. 7,106(1954) LA2 (S 94)3.9 H2 0 I*1 IV BURNS-J.H. AND W.R.BUSING, INDRG.CHEW, A.1510(1965) RB LI F? BURNS-J.H. AND W.R.BUSING, INDRG.CHEW, A.1510(1965) C5 LI F? BURNS-J.H. AND E.K. GORDON, NOTG.CHEW, A.1510(1965) LI 7 RF FA EANESFED, AND H.M.ONDIK. ACTA (CRYST, 15,1280(1962) LI X P3 09,H2 O GALYJ, AND A.HAROYA ACTA (CRYST, 19,432(1965) LI X 05 XEFFER:C. ET AL, INDRG.CHEW, G.119(19657) LI S 04.H2 O LIEBAUFF. ACTA (CRYST, 10+32(1965) LI Z 504.H2 O LIEBAUFF. ACTA (CRYST, 10+392(1965) LI Z 50 LIEBAUFF. ACTA (CRYST, 10+392(1965) LI Z 50 HAREZIO, A. ACTA (CRYST, 10+392(1965) LI Z 50 HAREZIO, A. ACTA (CRYST, 10+392(1965) LI Z 50 HAREZIO, A. ACTA (CRYST, 10+396(1965) LI GA 02 HAREZIO, A. ACTA (CRYST, 10+396(1965) LI GA 02 HAREZIO, A. ADI (A.HURKS) J.CHEN, JUNYS, LIXJAT(1964) LI R 02 HEREKA, J.P. AND W.MAREZIO, J.CHENS, J.CHEN, JONS, ACTA/CRYST, 10+28(1961) LI Z 904 ZACHARIASENNWH, AND H.A.PLETTINGEF. ACTA (CRYST, 10+28(1961) LI Z 904 ZIMANNJA, ACTA (CRYST, 13+86(1960) LI J D 04 ZIMANL, ACTA (CRYST, 13+86(1960) LI J D 04 ZIMANL, ACTA (CRYST, 13+86) LI B 05 ZACHARIASENNWH, AND H.A.PLETTINGEF. ACTA (CRYST, 10+28(1961) LI 2 904 ZIMANL, ACTA (CRYST, 13+86(1960) LI J D 04 XIML, ET AL., Z, ELEKTROCHEM, 405,586(1934) LI 2 0 LI+1 IV

ZIMILFL ET ALLY ZIELENTUGERT, SUPPORTES. L. C. C. LITI VI ABRAMME,S.C. AND J.L.BERNSTEIN, J.PHYS,CHEM, SOLIDS 20,1445(1967) LI TA D3 ABRAMME,S.C. AND M.JUKILIAWE, J.CHEM, MYNS, 35,2723(1967) LI CU CL3,2 H2 O ABRAMMS,S.C. ET AL., J.PHYS,CHEM,SOLIDS 27,097(1966) LI NA D3 BRUNTON.G., ACTA CRYST. 21,819(1966) LI U F5 BRUNTON.G., J.INGO, VUCL.CHEM, 20,105 20,1073(1967) LI V D5 BRUNTON.G., J.INGO, NUCL.CHEM, 20,105(1967) LI V D78 BURNS.J.H., ACTA CRYST. 21,80104601 K NAS FE2 TI2 (514 D12)2 GALY, J. AND A.HARDY, ACTA CRYST, 13,9432(1965) LI V2 D5 GELLEF.S. AND J.L.DURAND, ACTA CRYST, 13,9432(1965) LI V2 D5 GELLEF.S. AND J.P.REMIKA, J.PHYS,CHEM,SOLIDS 26,1277(1965) LI GA 02 MAREZIOLM, AND J.P.REMIKA, J.PHYS,CHEM,SOLIDS 26,1277(1965) LI GA 02 WYCKOFF.R.W.G., CRYSTAL STRUCTURES 1,119631 LI F

LU+3 VI BAATRAM-S.F., INORG.CHEM. 5,749(1966) LU6 U 012 TEMPLETONND.H. AND C.H.DAUBEN. J.AM.CHEM.SOC. 76-5237(195%) LU2 03

LU+3 VIII BRISSE/F., PH.O.THESIS/DALHOUSIE UNIV.,HALIFAX;N.S.(1967) LU2 TI2 O7 Euler,F. And J.A.BRUCE, ACTA CRYST, 19,971(1965) LU3 MF5 O12 SHANNON,R.O. AND C.T.PREWITT, TO BE PUBLISHED. R VS A (PYAOCHLORE)

MG+2 IV SmITH,J,V,, AM,MINERALOGIST 38,643(1953) CA2 MG SI2 07

SHITH.J.V., AM, HINERALOGIST 38,643(1953) CA2 MG 512 07 MG-2 VI BAUR.W.H., ACTA CRYST, 9,515(1954) MG F2 BAUR.W.H., ACTA CRYST, 15,815(1962) F2 S 00.4 H2 O BAUR.W.H., ACTA CRYST, 15,815(1962) MG F2 O7 CORBRIDGE.D.E.C., ACTA CRYST, 11,160(1958) MG H P 03.6 H2 O CORBRIDGE.D.E.C., ACTA CRYST, 9,991(1956) MG F2 O7 CORBRIDGE.D.E.C., ACTA CRYST, 9,991(1956) MG H P 03.6 H2 O GIBBS:06, V ET AL. AM.HINERALOGIST, IN PESS. MG F3 Co KASPER.J.S. AND J.S.PRENER, ACTA CRYST, 7,246(1954) MG 6 MV 08 LUKASZEWICZK.K. BULL.ACAD.POLONISCI.SFR.SCI.CHIM. 11,365(1954) MG 2 A52 07 RENTZEPERIS.F.J., AND C.T.SOLDATOSI, ACTA CRYST, 11,4585(1954) MG 5 04 SITEINFING., ACTA CHYST, 51574(1952) MG 2 M2 O TABEUCHIJY., ACTA CRYST, 51574(1952) MG 2 M2 O WYCKOFF.R.M.G., CRYSTAL STRUCTURES 1,1(1963) MG 0 WYCKOFF.R.M.G., CRYSTAL STRUCTURES 1,1(1963) MG 0 WYCKOFF.R.M.G., CRYSTAL STRUCTURES 1,1(1964) MG 0 ZEMANNAA, AND J.ZEMANN, ACTA CRYST, 10,409(1957) K2 MG 7 504)3

MG+2 VIII GIBB5-6.V, AND J.V.5MITH: AM.MINERALOGIST 50,2023(1965) MG3 ALZ 513 012 Zemanna, And J.Zemann. Acta Cryst, 19-835(196)) MG3 ALZ 513 012

MN+2 VI LS

WHE'Y L HS ARRAHAMS,S.C. AND J.W.REDDY, J.CHEM.PHYS. 43,2535(1965) MN MO 04 BAUR.M.H. ACTA CRYST. 11:486(1958) MN F2 GELLER.S. AND JL.DURAND, ACTA CRYST. 13:325(1960) LI WN P 04 MOSE-M.E. AND 04.J.BUERGER 2.KRIST, 11:15(161962) (MN,F7) BE (P 04) (O H) PEACOR.D.R. AND M.J.BUERGER 2.KRIST, 11:15(161962) (MN,F7) BE (P 04) (O H) PEACOR.D.R. AND M.J.BUERGER 2.KRIST, 11:15(161962) (MN,F7) BE (P 04) (O H) PEACOR.D.R. AND M.J.BUERGER 2.KRIST, 11:15(1916) RENTZEPERIS.G., N. BUELS JAHROBHTALGOONTA 2010 (MN,F7) BE (P 04) (O H) PEACOR.D.R. AND M.J.BUERGER 2.KRIST, 11:15(1916) SHITANES, G.H. ALT, D.HYS,SOCJAPAN 14,1155(1950) MN TI 03 WTCKOFF.R.W.G.R. CRYSTAL STRUCTURES 1.(1963) KN 0 WTCKOFF.R.W.G., CRYSTAL STRUCTURES 2.(1964) K MN F3 ZALKIN.A. ET AL., J.CHEM.PHYS, 376697(1962) CS MN F3

MN+2 VIII SHANNON,R.D. AND C.T.PREWITT, UNPUBLISHED DATA. R VS A (GARNET)

MN-3 V Abrahuksis.c. and J.L.Bernstein, J.Chem.Phys. 46:3776(1967) dv MN2 05 Yakel.H.L. et al., acta cryst, 16:457(1963) v MN 03

MN+3 VI LS BLASSE.G., J.INORG.NUCL.CHEM, 27,748(1965) SHANNON.R.O. AND C.T.PREMITT, TO BE PUBLISHED. R3 VS V (K3 MII] (C N)6)

MH-3 VT HS COLLINER-L. AND W.N.LIPSCOMB, ACTA CRYST. 2/104(1949) H MN 02 DACHS.H., Z.KRIST. 118.303(1063) MN 0 0 H FERT.A., BULL.SOC.FRANC.HIMERAL.CRIST. 85.257(1962) MN2 03 HASE.W.S.T AL., Z.KRIST. 124.428(1967) MN2 03 HEPWORTH.M.A. AND K.H.JACK: ACTA CRYST. 10.345(1957) MN F5 NORRESTAM.R., ACTA LCHM.SCAND, 21.2871(1967) MN2 03 ROTH.R.S., S.J.SCHNEIDER. J.RES.NATL.RUA.STO. 64A.306(1960) R3 V5 V (MN2 03) SHANGON, R.Q., INDG.CHEM. 61/147(1967) R3 V5 V (FEROVSKITE) SHANGON,R.Q., AND C.T.PREWITT, TO BE PUBLISHED. R VS A (MN2 03)

MN+4 VI ABBAHANS,S.C. AND J.L.BERNSTEIN, J.C.KEW,PHYS, 46,3776(1967) DY MN2 05 Abbahans, and W.Lendt. Z.Andrég.Allcem.(Kem. 269(165(1952) RB2 MN F6 Kasper,J.S. And J.S.PREMER, ACTA (ATST. 7.246(1958) MG6 MN 08 Shannon, R.D. And C.T.PREWITT, TO BE PUBLISHED. R3 VS V (MUTILE) WADSLEY.ALD., ACTA (CRYST. 8.165(1955) ZY MN3 07,3 H2 0

MN+6 IV PALENIK,G.J., INORG.CHEM. 6+507(1967) 82 MN 04 MN+7 IV FERRARIA, ET AL., ACTA CRYST, 21,681(1966) SR (MN 04)2,3 H2 0 JAVANIA, AND A. ENGELBRECHT, PHYS.REV, 96,649(1956) MN 03 F MOOMEYIR.C.L., PHYS.REV, 37,1306(1931) K MN 04 PALENIKY,J.J. RINGR.CHEM, 6,533(1967) K MN 04 SASVARIK,, 2.KRIST, 99,9(1938) AG MN 04 -----M0+3 VI HEPWORTH+M-A, ET AL., ACTA CRYST, 10,63(1957) Mo F3 NOA VI Ansellg.B. And L.Katz, acta Cryst. 21.482(1966) ZN2 MOS OR Brandteb.g. and A.C.Skapski. Acta Chem.Scand. 21.661(1967) No O2 Shannon.ad. And C.T.Premitt. To be published. Rs VS (Ruyile) VyCKOFF.R.V.G. (Crystal Structures 2.(1964) SR MO O3 VyCKOFF.N.G.G. Crystal Structures 2.(1964) SR MO O3 M0+5 VI Edwards,A.J. And R.D.PEACOCK, J.CHEM.Soc. 1961,4253 NA MO F6 Kierkegaard.d. And M.Westerlind. Acta Chem.Scand. 18,2217(1964) NO P 05 Mode IV ARRAHAMS.S.C., J.CHEM.PHYS. 46,2052(1967) ZN MO O4 ARRAHAMS.S.C., AND M.REDOY, J.CHEM.PHYS. 43,2533(1665) MN MO O4 ARRAHAMS.S.C. AND M.REDOY, J.CHEM.PHYS. 48,2619(1968) CH MO O4 ARRAHAMS.S.C. ET AL., J.C.EM.PHYS. 48,2619(1968) CH MO O3 IBERS.J.A.MO G.W.SMITH, ACTA CRYST, 17,100(1964) NA CO2,3 MO3 012 KIHLBORG.L., ARKIV KEMI 21,357(1965) MO O3 IHLBORG.L., ARKIV KEMI 21,357(1965) MO O3 IHLBORG.L., ARKIV KEMI 21,357(1965) SELEBORG.H., ACTA CHEM.SCAND, 20,2195(1966) K2 MO3 010 SELEBORG.H., ACTA CHEM.SCAND, 21,2199(1967) NA2 MO3 07 SILLEN.L.G. AND K.LUNDBORG, 2,ANORG.ALLGFM.CHEM. 252,211943) L.2 MO O6 MO+6 V EICK.M.A. AND L.KIMLBORG, ACTA CHEM.SCAND. 20,1658(1966) V2 MO 08 GATEHOUSE.B.M. AND P.LEVERETT, CHEM.COMM. 1967,374 K2 MO3 010 SELEBORG,M., ACTA CHEM.SCAND. 20,2195(1966) K2 MO3 010 SELEDORGATI ACTA CHEM.SCAND. 20121314007 AC HOS OTO GATENOUSE:B.H. AND P.LEVERETT, CHEM.COMM. 1967.374 K2 M03 010 KIERKEGARD.P., ARKIV KEMI 18:521(1962) NA M0 02 P 04 KIERKEGARD.P., ARKIV KEMI 18:521(1962) NA M0 02 P 04 KIERKEGARD.P., ARKIV KEMI 18:13635 (NO 02)2 P 20 A7 KIERKEGARD.P., ARKIV KEMI 21:35(1965) NG 03 (NO 02)2 P 04 KIHLORG.L., ARKIV KEMI 21:35(1965) M0 03 KIHLORG.L., ARKIV KEMI 21:35(1965) M0 03 KIHLORG.L., ARKIV KEMI 21:35(1966) KIHLORG.L., ARKIV KEMI 21:35(1966) M0 03 KIHLORG.L., ARKIV KEMI 21:45(1966) M0 03 KIHLORG.L., ARKIV KEMI 21:45(1966) M0 03 SELEDORG.H., ACTA CHEM.SCAND. 20:2195(1966) K2 M03 010 SMITH.G.W. AND J.A.IBERS; ACTA CRYST, 19:269(1965) C0 M0 04 MO+6 VII KIHLBORG,L., ARKIV KEMI 21.427(1964) MO5 014 N +3 VI N +5 III Cherin,p. Et Al., Acta Cryst, 23,455(1967) NA N 03 N 45 VT NA+1 IV Corbridge.d.e.c., Acta Cryst, 13,263(1968) NA5 P3 010 Wunderlich.j.a., Acta Cryst, 10,462(1957) NA 0 H.M2 0 Wyckoff.r.w.g., Crystal Structures 1,(1963) NA2 0 NA-1 V HENILYD,W., ACTA CRYST, 10:37(1938) NA O N.* M2 D LARSENIF,K. ET AL., ACTA CHEM,SCAND, 21,1281(1967) NA3 5N% GE10 030 10 H14 MCDONALDWYS, AND D.W.JCRUICKSMANN, ACTA CRYST, 22-37(1967) NA2 5T 03 ONDIKH,M., ACTA CRYST, 18,226(1965) NA3 P3 09 ONDIKH,M., ACTA CRYST, 18,226(1965) NA3 P3 09-00 PANT,A.K. AND O.W.JCRUICKSMANK, ACTA CRYST, B24-13(1968) NA2 512 05 ROMERS/C. ET AL., ACTA CRYST, 22-786(1967) NA3 P5 09 A+1 VI DUBBS J.H.. INDRO,CHEW, 0,081(1965) NA NO FA DUBBS DDE.D.E.C., ACTA CRYST, 13-265(1960) NA5 P3 D10 CROMER,D.T. ET AL, ACTA CRYST, 27:182(1975) NA AL (5 Obj2,12 H2 O CRUICKSHANK,D,W,J,, ACTA CRYST, 27:1957(1965) NA NA P2 07:10 H2 O DYERAL,D. ET AL, J.AM,CHEW,SGC, 76:1995(1959) NA NI 0? 0:ESE;R,F,J,R.: SCIENCE 150;1435(1966) NA2 P8 O6 (0 H)2,3 H2 O WECOMALD.R.R. ET AL., ACTA CRYST, 17:1104(1964) NA S02 P5 MCDOMALD.R.R. ET AL., ACTA CRYST, 17:104(1964) NA S02 P5 MCDAMALD.R.R. ET AL., ACTA CRYST, 17:104(1964) NA S02 P5 SHARMAS,D.T. ACTA CRYST, 16:35(1961) NA NO 02 P0 SHARMAS,D.T. ACTA CRYST, 17:5104(1964) NA S02 P5 SHARMAS,D.T. ACTA CRYST, 17:5104(1964) NA S17 F8 WYCKOFF,R.M.A., CYST, 15:390(1999) NA TH2 F9 ZACHARIASCH.W.H., ACTA CRYST, 27:390(1999) NA TH2 F9 ZACHARIASCH.W.H., ACTA CRYST, 17:1908(1964) NA2 SI F6 N841 VT NA-1 YII ANDERSSON;S, AND A.O.WADSLEY, ACTA CRYST. 14,1245(1961) NA2 TI3 07 CANNILLOFF. ET AL., ACTA CRYST. 21,220(1966) K NA3 FE2 TI2 (514 012)2 FANGIS.N., Z.KRIST. 99A.1(1938) NA B 02 KIERKEGARD.P., ARKIV KEMI LA:535(1962) NA V 02 P 04 KOKKOROS.P., Z.KRIST. 99A.38(1938) NA AL F AS 04 MAREZION, ET AL., ACTA CRYST. 16,354(1963) NA B 02 RAO:B.RAMA, ACTA CRYST. 14,738(1961) NA2 CU (5 04)2.2 H2 NA+1 VIII ** I VII MCDONALDER.R. ET AL., ACTA CRYST. 17,1104(1964) ΝΑ SN2 F5 MCNZER.G., Z.KRIST. 75.255(1950) ΝΑΣ [13 ALZ F12 Rumanoya: Int., G.F.VOLDTNA; DOLLARADINAUX SSSR 123,78(1956) ΝΑ CU2 Ο Η (5 04)2.Η2 WEISSFR, ET AL., ACTA CRYST. 20.534(1965) ΝΑΣ Η ΤΙ F8 Zacharlashwih, Z.KRIST. 73A191(1930) ΝΑ CL 04 NA+1 IX ANDERSSON,S. AND A.D.WADSLEY, ACTA CRYST, 14+1245(1961) NA2 TI3 07 NB+2 VI ANDERSSON, G. AND A.MAGNELI+ ACTA CHEM.SCAND. 11,1065(1957) NB 0 NB+3 VI HEPWORTH.M.A. ET AL., ACTA CRYST. 10,63(1957) NB F3

NB+4 VI Marinder,8., Arkiv Kemi 19,435(1962) NB 02 Skannon,r.D. And C.T.Prewitt, to be published. R3 V5 V (Rutile)

N8+5 IV BATEHOUSE, B.M. AND A.D. WADSLEY, ACTA CRYST. 17,1545(1964) NB2 05

Table 1(b) (cont.)

ANDERSSON,S., ACTA CHEN,SCAND, 18,2339(1964) NB3 07 F BODE.H., AND H. DOHMEN, ACTA CRYST, 11,80(1958) K NB F6 SATEHOUSENEN, AND ALO,MADSLEY, ACTA CRYST, 17,1545(1964) NB2 (KATZ.L. AND H.D. MEGAN, ACTA CRYST, 22,639(1967) K NB 03 MOROSIN.8, AND A.ROSENZUEIG, ACTA CRYST, 18,437(1965) G NG 04 PEDERSEN.8.F., ACTA CHEM.SCAND, 16,421(1962) AL NP 04 SLEPSHIPALC, AND P.ROSENS, CHEM.COWM, (1965)611 SB NB 04 SLEIGHT,A.V., TO BE PUBLISHED, HG2 NB2 07 N82 05 NB+5 VII BROWN-6.M. AND L.A.WALKER, ACTA CRYST. 20,220(1966) K2 NR F7 HOARD,J.L., J.AM.CHEM.SOC. 61,1252(1939) K2 NB F7 ND+3 VI TEMPLETON;D.H. AND C.H.DAUBEN; J.AM.CHEM.SOC. 76;5237(1954) ND2 03 ND+3 V111 HILLIGAN,W.O. AND L.W.VERNDN, J.PHYS.CHEM. 56,145(1952) ND V 04 Shannon,R.D. And C.T.PREWITT, TO BE PUBLISHED. R VS A (PYROCHLORE) ND+3 IX BURNS+J.H., INORG.CHEM. 4+881(1965) NA ND F4 NI-2 VI BALZ-D. AND K.PLIETH, Z.ELEKTROCHEW. 59,545(1955) K2 NT F4 BAURAWH. ACTA CRYST 114486(1958) NI F2 BJLLY.C. AND M.H.MAENDLER. J.AM.CHEM.SOC. 79,1049(1957) NI F2 DYER.L.D. ET AL., J.AM.CHEM.SOC. 76,1496(1951) NA NI 02 KEELING.R.O.JM. ACTA CRYST. 10,209(1957) NV 04 LANDERJ.J.J. ACTA CRYST. 10,209(1957) COMMERING.H.G. Z.ANDG.ALLGEM.CHEM. 353,13(1967) A2 NT F6 SHIRAKE.G. ET AL., J.ANDG.ALLGEM.CHEM. 353,13(1967) A2 NT F6 SHIRAKE.G. ET AL., J.ANDG.ALLGEM.CHEM. 353,13(1967) A2 NT F6 VAN NIEKERK.J.N., F.R.SCHDENING. ACTA CRYST. 6609(1953) NIIC H3 C 0 0)2.4 H2 O WYCKOFF.R.W.G. CRYSTAL STRUCTURES 1,(1963) NI F3 NI+3 VI LS BLASSE,g., J.INORG.NUCL.CHEM, 27,748(1965) K3 NI F6 Shannon.r.D., INORG.CHEM. 6.1474(1967) R3 V5 V (PEROVSKITE) NI+3 VI HS BLASSE.G., J.INORG.NUCL.CHEM, 27,748(1965) NP+2 VI WYCKOFF,R.W.G., CRYSTAL STRUCTURES 1,(1963) NP 0 NP+3 VI SHANNON,R.D. AND C.T.PREWITT, TO BE PUBLISHED. R3 V5 V (LA F3) NP+4 VI NP+4 VIII Shannon,r.D. and C.T.PREWITT, TO BE PUBLISHED. R VS A (FLUGRITE) WYCKOFF.R.W.G., CRYSTAL STRUCTURES 1.(1963) NP 02 NP+7 VT 0 -2 11 0 -2 111 0 -2 IV 0 -2 VT 0 -2 VIII 05+4 VI SHANNON,R.D. AND C.T.PREWITT, TO BE PUBLISHED. R3 VS V (RUTILE) P +3 VI P •3 VI P •5 IV CRUICKSHANK:D.W.J., ACTA CRYST. 17.671[1964] NA P 03 N H3 CRUICKSHANK:D.W.J., ACTA CRYST. 17.671[1964] NA P 03 N H3 CRUICKSHANK:D.W.J., ACTA CRYST. 17.672[1964] NA P 03 D11 CRUICKSHANK:D.W.J., ACTA CRYST. 17.675[1964] NA P 03 D12 CRUICKSHANK:D.W.J., ACTA CRYST. 17.675[1964] P 05 CRUICKSHANK:D.W.J., ACTA CRYST. 17.675[1964] P 05 CRUICKSHANK:D.W.J., ACTA CRYST. 17.675[1964] P 05 CRUICKSHANK:D.W.J., ACTA CRYST. 17.675[1964] P 05 CRUICKSHANK:D.W.J., ACTA CRYST. 17.675[1964] P 05 CRUICKSHANK:D.W.J., ACTA CRYST. 17.621[1964] NA P 05 KEFNER,C. ET AL., INDREG.CHEW., 6.119(1967) LI3 P 04 KIERKEGARD:P., ARKIY KCHT, 18.521[1961] NA M0 02 P 04 KIERKEGARD:P., AND S.HOLWEN, ARKIY KCHT 23.213[1965] AG NO P 06 KIERKEGARD:P. AND S.HOLWEN, ARKIY KCHT 23.213[1965] AG NO P 06 KIERKEGARD:P. AND S.HOLWEN, ARKIY KCHT 23.213[1965] AG NO P 06 KIERKEGARD:P. AND S.HOLWEN, ARKIY KCHT 23.213[1965] AG NO P 06 KIERKEGARD:P. AND S.HOLWEN, ARKIY KCHT 23.213[1965] AG NO P 06 KIERKEGARD:P. AND S.HOLWEN, ARKIY KCHT 23.213[1965] AG NO P 06 KIERKEGARD:P. AND S.HOLWEN, ARKIY KCHT 23.213[1965] AG NO P 06 KIERKEGARD:P. AND S.HOLWEN, ARKIY KCHT 23.213[1965] AG NO P 06 KIERKEGARD:P. AND S.HOLWEN, ARKIY KCHT 23.213[1965] AG NO P 06 KIERKEGARD:P. AND S.HOLWEN, ARKIY KCHT 23.213[1965] AG NO P 06 KIERKEGARD:P. AND S.HOLWEN, ARKIY KCHT 23.213[1965] AG NO P 06 KIERKEGARD:P. AND S.HOLWEN, ARKIY KCHT 23.213[1965] AG NO P 06 KIERKEGARD:P. AND S.HOLWEN, ARKIY KCHT 23.213[1965] AG NO P 06 KIERKEGARD:P. AND S.HOLWEN, AFT 06 AFT 07 MULING, ACTA CRYST. 13.463(1966) AL P 04 MEBB.N.C., ACTA CRYST. 13.463(1966) LI3 P 04 P +5 VI PA+T VT PA+9 VT PA+8 VIII WYCKOFF,R.W.G., CRYSTAL STRUCTURES 1,(1963) PA 02 Pa+5 VI PA+5 IX BROWN+D. ET AL., J.CHEM.SOC. 1967A+1429 K2 PA F7 PB+2 IVPY BUCHER.H.L. AND D.R.PEACOR, Z.KRIST, 126+98(1968) P8 51 03 BYSTROM.A. AND L.EVERS, ACTA CHEM.SCAND, 4+613(1950) A42 P8 02 KAY.H.I., ACTA CRYST, 14+80(1961) P8 0 PB+2 VI MAGNELIA, ARKIV KEMI, MINERAL.GEOL. 159, 3(1942) PB 5B? 06 HOLLER.C.K., ACTA CHEM.SCAND. 6.81(1954) PB (N H4)2 (5 04)2 WYCKOFF,R.W.G., CRYSTAL STRUCTURES 2,(1964) C5 PB F3 P9:2 VIII FINNEYIJJ., AM.MINERALDGIST 46.1(1963) P8 F22 (AS 04)2 (O H)2 LECTEUEWICZ.J., Z.KRIST. 121.158(1965) P8 0 KAYIMII, ACTA CRYST. 14.80(1961) P8 0 QUARENIS, AND R.DEPIERI, ACTA CRYST, 19.287(1965) P8 CR 04 WYCKOFF,R.W.G., CRYSTAL STRUCTURES 1,(1963) P8 F2 PB+2 IX COLBY:M.Y. AND L.J.B.LACOSTE, Z.KRIST, 84,299(1932) PB C 03 PB+2 XI NARAY-SZABO+I. AND 6.ARGAY+ ACTA CHIM.ACAD.SCI.HUNG. 40,283(1964) PB CR 04

PB+2 XII HAMILTON:W.C., ACTA CRYST, 10,103(1957) PB (N 03)2 SAHLIK., BEITR.MINERAL,PETROG, 9,111(1963) PB 5 04 PB+4 VI BYSTROH,A., ARKIV KEMI,MINERAL,GEOL, 25A,13(1947) PG 02 SHANNOH,R.O. AND C.T.PREWITT, TO BE PUBLISHED. R3 VS V (RUTILE) SHANNOH,R.O. AND C.T.PREWITT, TO BE PUBLISHED. R VS A (PYRCHLORE) WYCKOFF.M.W.G., CRTSTAL STRUCTURES 2.(1964) BA PB 03 B0+4 VIII Shannon,R.D. and C.Y.PREWITT, TO BE PUBLISHED. R VS & (FLUOPITE) PD+1 II ROGER5+0.B. AND R.D.SHANNON, FALL A.C.S. MTG.+ATLANTIC CITY(1968) PD CO 02 PD+2 IV50 WASER;J. ET AL.+ ACTA CRYST. 6,661(1953) PD 0 PD+2 YI RARTLETT,N. AND R.MAITLAND, ACTA CRYST, 11,747(1958) PD F2 PD+3 VI HEPWORTH+M+A, ET AL., ACTA CRYST, 10,63(1957) PD F3 PD+4 VI SLEIGHT, A. W., TO BE PUBLISHED. R VS A (PYROCHLORE) PH+3 VI TEMPLETON.D.H. AND C.H.DAUBEN, J.AM.CHEM.SOC. 76:5237(1954) PM2 03 P0+4 VIII SHANNGN.R.D. AND C.T.PREWITT, TO BE PUBLISHED. R VS A (FLUORITE) PO+6 VI PR+3 VI FERT.a., BULL.SOC.FRANC.MINERAL CRIST. 85:267(1962) PR2 03 TEMPLETONID.H. AND C.H.DAUBENI J.AM.CHEM.SOC. 76:5237(1954) PR2 03 PR+3 VIII SMANNON,R.D. AND C.T.PREWITT, TO BE PUBLISHED. R VS & (PYROCHLORE) PR+4 VI MYCKOFF,R.W.G.+ CRYSTAL STRUCTURES 2,(1964) BA PR 03 PR+4 VIII WYCKOFF.R.W.G., CRYSTAL STRUCTURES 1,(1963) PR 02 PT+2 VI PT+N VI 44 VI BUORLING-C.O., ARKIV KEMI.WINERAL.GEOL, 158,2(1942) K2 PT (0 H)6 HOEKSTRAIH.R. AND S.SIGGEL, INORG.CHEM, 7:141(1968) TL2 PT2 OT MELLOR.D.P. AND N.C.SIGFUMENSON, AUSTR.J.PES. \$4406(1951) K2 PT F6 SHANNON.R.D., SOLID STATE COMM. IN PRESS, R3 V5 V (PT R2) PU+3 VI Smannon,R.D. and C.T.PREWITT, TO BE PUBLISHED, R3 V5 V (LA F3) PU+4 VI WYCKOFF,R.W.G., CRYSTAL STRUCTURES 2,(1964) BA PU 03 PU+4 VIII WYCKOFF,R.W.G., CRYSTAL STRUCTURES 1,(1963) PU 02 RA+2 VI RB+1 VI HOPPER, AND H.SABROWSKY, Z.ANORG,ALLGEN.CHEM. 339,144(1965) RB SC 02 PLYUKHN.V.V. AND N.V.BELOV. SOVIET PHYS.-CRYST. 6.685(1962) RB RE2 F5 WYCKOFF,R.W.G., CRYSTAL STRUCTURES 1,(1963) RB F R8+1 VII CORBRIDGE+0.E.C., ACTA CRYST, 9,308(1956) R8 P 03 CRUICKSHANK-D.W.J., ACTA CRYST, 17,681(1969) (R8 P 03) R8+1 VIII ANDERSSON'S. AND A.D.WADSLEY, ACTA CRYST. 15,194(1962) R82 TI6 013 BURNSJ.J.M. AND W.R.BUSING. INORG.CHEW. 4.1510(1965) R8 LI F2 CRUICKSMANK'D.W.J., ACTA CRYST. 17,641(1964) (R8 P 03) R8+1 XII Ellingerffh. And W.H.Zachariasen, J.Phys.Chem. 58,405(1954) R8 AM 02 C 03 Hoard-Jl. And V.Blair. J.Am.Chem.Soc. 57,1985(1935) R4 B F4 RE+4 VI SHANNON, AND C.T.PREWITT, TO BE PUBLISHED. R3 VS V (RUTILE) RE+5 VI DONOHUE,P.C. ET AL., INORG.CHEM. 4,1152(1965) CD2 RE2 07 RE+6 VI WYCKOFF+R.W.G., CRYSTAL STRUCTURES 2,(1964) RE 03 RE+7 IV KREBS+8. ET AL., CHEM.COMM. (1968)263 RE2 07 MORROW.J.C., ACTA CRYST, 13:443(1968) K RE 04 RE+7 VI KREB5+8. ET AL., CHEM.COMM. (1968)263 RE2 07 RH-3 VI HEPRORTH-M.A, ET AL., ACTA CRYST, 10,63(1957) RH F3 Shannon, R.D., Indrg.chem. 6:1478(1967) R3 V5 V (PEROVSKITE) Shannon, R.D., And C.T., PREVITT. TO RF FUBLISHED. R3 V5 V (CORUNDW) RH+4 VI SHANNON, AND C.T.PREWITT, TO BE PUBLISHED. R3 VS V (RH 02) RU+3 VI Hepworth+M.A. Et al., Acta Cryst, 10,63(1957) RU F3 Shannon.r.D. And C.T.PREWITT, TO BE PUBLISHED. R3 VS V (PEROVSKITE) RU+N VI Cotionafia, and J.T.Maguf. Indrg.chem, 5.517(1955) RU 0.2 Donohue, P.C. et Al., Indrg.chem, 4.306(1955) BA PV 0.3 Simanohur, B. And C.T.Premitt, To be published. R3 VS V (Rutile) Smannohur.D. And C.T.Premitt, To be published. R3 VS V (Rutile) 5 -2 VI 5 +4 VI S +6 IV BANRIW-H., ACTA CRYST. 17,865(1964) MG 5 04.4 H2 0 BANRIW-H., ACTA CRYST. 17,1167(1964) MG 5 04.7 H2 0 BANRIW-H., ACTA CRYST. 17,1151(1964) FFG 094.7 H2 0 CROMERDET. FT AL., ACTA CRYST. 21,351(1966) CC AL (S 04)2,12 H2 0 GRAEBER.E.J. ET AL., AM,HINERALOGIST 50,1929(1965) K FF (S 04)2

GRAEBER, E, J. ET AL., AM. MINERALOGIST 50.1929(1965) MG (N H4)2 (S 04)2.6 H2 0 HONNE, E, SOVIET PHYS.-CRYST, 7.559(1963) CA S 04 KOXKOROS, P.A. AND P.J.RENTZEPERIS, Z.KRIST, 119,224(1963) HG S 04 LARSONIA.C. ACTA CRYST, 187,171655) LIZ S 04.H2 0 0'CONNOR.B.H. AND D.H.DALE, ACTA CRYST, 21.705(1966) NI S 04.6 H2 0 RENTZEPERISP.J., Z.KRIST, 117,431(1962) S NS 04 SAML, K., BEITA, MINERAL, PETROG, 9,111(1963) BA 5 04 SAML, K., BEITA, MINERAL, PETROG, 9,111(1963) PB S 04 ZALKIN, AE TA L., ACTA CRYST, 15,1219(1962) CO S 04.6 M2 0 5 46 VT SB+3 IVPY BYSTROW-A. AND K.WILHELMI, ARKIV KEMI 3,373(1953) CS SR2 F7 Rođers.d. And A.C.SKAPSKI, proc.chem.Soc. 12,400(1964) SB2 04 SKAPSKI,A.C. AND D.ROGERS, CHEM.COMM. (1965)611 SR NB 04 S8+3 V 8YSTROM,A, EY AL., ARKIV KEMI 4,175(1952) K S8 F4 8YSTROM,A. EY AL., APKIV KEMI 6,77(1953) NA S8 F4 58+3 VI 50-5 11 S0-5 11 AUBIVILLIUS-B., ARKIV KEMI 3,153(1951) BI 59 04 AUBIVILLIUS-B., ARKIV KEMI 3,153(1951) BI 59 04 AUBIVILLIUS-B., ARKIV KEMI 25,505(1966) K3 5H5 014 BUDRS,1,4, ACTA (CATST. 15,1098(1962) L1 58 F6 BUTSTROM.A. ET AL., ARKIV KEMI-MIERAL,6EOL. 158,HW(1942) X 582 06 MAGNELIA., ARKIV KEMI-MINERAL,6EOL. 158,HW(1942) X 582 06 ROGERS-D. AND A.C.SKAPSKI. PROC.(HEM. SOC. 12:400 (1964) B 592 06 ROGERS-D. AND A.C.SKAPSKI. PROC.(HEM. SOC. 12:400 (1964) B 592 06 SCHREVELIUS-N., ARKIV KEMI-MINERAL,6FOL. 158,H7(1943) TL 58 F6 SPIEGELBERGEP., ARKIV KEMI-MINERAL,6FOL. 158,H7(1943) TL 58 F6 SPIEGELBERGEP., ARKIV KEMI-MINERAL,6FOL. 158,H7(1943) TL 58 F6 ZEMANN-J., NEUES JAHRB,MINERAL,NONATSH, 67,(1959) NA 59 (0 H)6 2C-RAWNJL, HEUES JORNOUTHICHAELHOUND, 3, 0711017 HAT DATA SCA3 VI SCA3 VI SCA3 VI SRUENCHARKLD, N.J., ACTA CREST, 15,441(1962) SC2 SI2 07 GELLER-S, ET AL.- Z.KRIST, 124,136(1967) SC2 03 HOPPE/R, ET AL.- Z.KRIST, 124,136(1967) SC2 03 HOPPE/R, ET AL.- Z.KRIST, 124,136(1967) SC 203 HOPPE/R, ET AL.- Z.ANDRG.ALLGEM.CHEM, 339,130(1965) LI SC 02 HOPPE/R, ET AL.- Z.ANDRG.ALLGEM.CHEM, 339,130(1965) LI SC 02 HOPPE/R, ET AL.- Z.ANDRG.ALLGEM.CHEM, 339,130(1965) SC 52 04 JACK.K.H. AND V.GUTMANN. ACTA CRYST. 4,246(1951) SC F3 HILLIGAN.H.O. AND J.L.MATEE, J.PHYS.CHEM. 602/37(1956) SC 0 0 H HULLER-BUSCHBAUM.H., H.G.SCHNERING, Z.ANDRG.ALLGEW.CHEM, 336,295(1965) CA SC2 04 NOMACKI.N., Z.KRIST. 1012/371393) SC F3 ROTHIR.S, AND S.J.SCHNEIDER. J.RES.NATL,AUR.STD. 64A.309(1960) R3 VS V (MN2 03) SCHADERT:K. AND A.SETTZ. Z.ANDRG.ALLGEV.CHEM. 2562619401) SC (D H)3 SHANNON.R.D., INORG.CHEM. 6,1474(1967) R3 VS V (MN2 03) TC+7 VI TE-2 VT -SC+3 VIII MOONEY:R.C.L., ACTA CRYST, 9,677(1956) SC P 0% MILLIGAN;H.O. AND L.M.VERNON, J.PHYS,CHEM. 56,145(1952) SC V 0% SHANNON;R.O. AND C.T.PREWITT, TO BF PUBLISHED. R V5 A (PYROCHLORE) TE+4 VI TE+6 V1 SE-2 VI 5E+4 VT TH+4 VIII SE+6 IV FUESS:M, AND G.WILL; Z.ANORG.ALLGEM.CHEM. 358:125(1968) M SE 04 NARAT-SZABO:I. AND G.ARGAY. ACTA CHIM.ACAD.5CI.HUNG. 39.A5(1963) NA2 SE 04 SEAR VI SE46 VI SIAN IV SIAN IV SIAN IV SIAN IV SIAN IV SIAN INSTITUTION SINCELLER, ACTA CRYST. 11/437(1958) GPOSSULARITE BROWNB,E, AND SIM,BALLEY, ACTA CRYST. 17/1391(1964) K AL ST3 08 BURNHAM,C.W., Z.KRIST. 118/337(1963) AL2 ST 05 CRUICKSHANK,D.W.J., ACTA CRYST. 17/65(1964) ST-05 GIBBSSG,V. AND J.V.SWITH. AM MINERALOGIST 50,222(1965) MG3 AL2 ST3 012 GLASSER.LS.D. AND J.V.SWITH. AM MINERALOGIST 50,222(1965) MG3 AL2 ST3 012 GLASSER.LS.D. AND J.V.SWITH. AM MINERALOGIST 50,222(1965) LIEBAUF. ACTA CRYST. 14,355(1961) NA2 ST2 05 LIEBAUF. ACTA CRYST. 14,355(1961) NA2 ST2 05 HIEBAUF. ACTA CRYST. 14,355(1961) NA2 ST2 05 HIEBAUF. ACTA CRYST. 14,355(1961) NA2 ST2 05 HIEBAUF. ACTA CRYST. 110/117(1965) CA2 ST 04 MINITAD.K. ET AL.. ACTA CRYST. 18.787(1965) CA2 ST 04 SMITH-J,V. AND SIM,BALEY. ACTA CRYST. 16,801(1965) ST-0 HILLIAMS,F.P. AND H.O.HEGAM. ACTA CRYST. 16,801(1965) ST-0 HILTEST 0, ACTA CRYST. 110/117(1965) CA2 ST 04 SMITH-J,V. AND SIM,BALEY. ACTA CRYST. 16,801(1965) AL2 ST 04 SMITH-J,V. AND SIM,BALEY. ACTA CRYST. 17.882(1964) ALDITES ZEMANNIA. AND J.ZEMANN. ACTA CRYST. 14/835(1961) MG3 AL2 ST3 012 HILLIAMS,F.P. AND M.O.H. ACTA CRYST. 14/835(1961) MG3 AL2 ST3 012 HILLIAMS,F.P. AND M.O.H. ACTA CRYST. 14/835(1961) MG3 AL2 ST3 012 HILLIAMS,F.P. AND M.O.H. ACTA CRYST. 14/835(1961) MG3 AL2 ST3 012 HILLIAMS,F.P. AND M.O.H. ACTA CRYST. 14/835(1961) MG3 AL2 ST3 012 HILLIAMS,F.P. AND M.O.H. ACTA CRYST. 14/835(1961) MG3 AL2 ST3 012 HILLIAMS,F.P. AND M.O.H. ACTA CRYST. 14/835(1961) MG3 AL2 ST3 012 HILLIAMS,F.P. AND M.O.H. ACTA CRYST. 14/835(1961) MG3 AL2 ST3 012 HILLIAMS,F.P. AND M.O.H. ACTA CRYST. 14/835(1961) MG3 AL2 ST3 012 HILLIAMS,F.P. AND M.O.H. ACTA CRYST. 14/835(1961) MG3 AL2 ST3 012 HILLIAMS,F.P. AND M.O.H. ACTA CRYST. 14/835(1961) MG3 AL2 ST3 012 HILLIAMS,F.P. AND M.O.H. ACTA CRYST. 14/835(1961) MG3 AL2 ST3 012 HILLIAMS,F.P. AND M.O.H. ACTA CRYST. 14/835(1961) MG3 AL2 ST3 012 HILLIAMS,F.P. AND M.O.H. ACTA CRYST. 14/835(1961) MG3 AL2 ST3 012 HILLIAMS,F.P. AND M.O.H. ACTA CRYST. 14/835(19 TI+2 VI TI+3 VI 14 V1 DEADMORE:D.L. AND W.F.BRADLEY, ACTA CRYST. 15,186(1962) K3 SI F7 HAMILTON.W.C., ACTA CRYST. 15,353(1962) FC SI F6.6 H2 0 HAMRDJ.L. AND H.B.WILTAMS, J.AW.CEME.SOC. 64,653(1962) (N H4)3 SI F7 STANKOJJA, AND I.C.PAUL. INDRG.CHEM. 64,863(1967) (CO (N H4)3 SI F7 STANKOJJA, AND I.C.PAUL. INDRG.CHEM. 64,864(1967) (CO (N H4)3 SI F7 STANKOJ,A, AND N.J.BLCOV, DOKLARG.JNAN FOR SIGNATION (CO (N H4)3 SI F6 STISHOV,SN, AND N.J.BLCOV, DOKLARG.JNAN FC. GRYST J.201(1956) (N H4)2 SI F6 ZALKINA. ET AL., ACTA CRYST. 17,1060(1964) NA2 SI F6 5I+4 VI DEADM SN+3 VI TEMPLETON, D.H. AND C.H. DAUBEN, J.AM. CHEM. SOC. 76, 5237(1954) 5M2 03 SH+3 VIII BRISSEF., PH.D.THESISJOALHOUSIE UNIV,,HALIFAX,H.S.(1967) SH2 M2 07 EULERF, AND J.A.BRUCE, ACTA CRYST, 19,971(1965) SH3 FF2 FE3 012 SHANNON,R.D. AND C.T.PREWITT, TO BE PUBLISHED, R VS A (PYROCHLORE) 5N+2 V1 SN+2 VIII BRISSEFF., PH.D. THESIS. DALHOUSIE UNIV., HALIFAX.N.S. (1967) 5N2 M2 07 SN+4 VI BAUR,WH., ACTA CRYST, 9,515(1956) SN 02 BRISSE:F., PH.D.THESIS.DALHOUSIE UNIV.,HALIFAX,N,S.(1967) M2 SN2 07 Hoppe,R, AND W.DAHNE, NATURVISS, 49r258(1962) SN F4 LARSEN,FK.ET AL., ACTA (FUR,SCAND, 2:1:21(1967) NAB SN% GEID 030 (0 H)4 WYCKOFF,R.W.G., CRYSTAL STRUCTURES 2,(1964) M SN 03 SR+2 VI BARTL:, AND W.SCHUCKMANN, NEUES JAHDB.MINERAL, MONATSH. 1966;253 SR2 82 05 GLASSER:L.S.O. AND F.P. GLASSER, ACTA CRYST. 18.435(1965) SR3 51 05 RIETVELDIH.M., ACTA CRYST. 20.506(1966) SR3 06 WYCKDFF.M.W.G., CRYSTAL STRUCTURES 1.1(963) SR 0 SR+2 VII Bartl-H, And W.Schuckmann, Neues Jahrb.Hineral, Monatch, 1966,253 Sr2 82 05 Schwertno:H.G. And R.Hoppe, Z.Anorg.Allgem.Chem, 312,87(1961) Sr ZN 02 SR+2 VIII BARNIGHAUSEN.H. AND J.WEIDLEIN. ACTA CRYST. 22,252(1967) 5R (0 H)2,H2 0 SMITH.H.G., ACTA CRYST. 6.604(1953) 5R (0 H)2.6 H2 0 VANMERBERGUN., ACTA VENI 14,17(1959) 5R 02.6 H2 0 VINMERBERGUN.; ARVIV KENI 26,149(1967) 5R CR2 07 WILMELSIK.K.A., ARVIV KENI 26,157(1967) 5R CR2 07 WILMELSIK.A., CRYSTAL STRUCTURES 1,(1963) 5R F2 U +4 VI

SR+2 X CLARX,J,R., AM,MINERALOGIST 109,1509(1964) 5R 86 09 (0 H)2.3 H2 0 FERRARI,A, ET AL., ACTA CRYST, 21+68(1966) 5R (MN 0417,3 H2 0 ZACHAPITASEN.W.H., ACTA CRYST, 1:263(1948) 5R3 (P 04)2 SR+2 XII Donmay.j.D.H. Et Al., Crystal data determinative tables(1963) SR TI 03 Garsked. And D.R.PEACOR, Z.KRIST, 121.204(1965) SR 5 nu Zachariasen.w.H., Acta Cryst. 1,263(1948) SR3 (P 04)2 TA+3 VI HEPWORTH+M+A. ET AL., ACTA CRYST, 10,63(1957) TA F3 Jahnberg-L, Acta Chem.Scand. 17,2548(1963) CA TA2 06 TA+4 VI SHANNON,R.D. AND C.T.PREWITT, TO BE PUBLISHED R3 VS V (RUTILE) TA+5 VI ARRAHAMS/S.C. AND J.L.BERNSTEIN, J.PHYS.CHEM.SOLIDS 28,1685(1967) LJ TA 03 ABRAHAMS/S.C. ET AL., J.PHYS.CHEM.SOLIDS 28,1685(1967) LJ TA 03 BODE-H. AND M.DOHREN, ACTA (RYST, 11,80(1958) K TA F6 BRISSE/F, PH.O.,THESIS/DALHOUSIC UNIU-MALIFAXINIS,(1967) CD2 TA2 07 VULSDENJE, ACTA CRTST. 4,373(1951) K TA 03 WYCKOFF, W.G.G. GRTSTAL STRUCTURES 2,(1964) K TA 02 WYCKOFF, R.W.G.G. GRTSTAL STRUCTURES 2,(1964) K TA 03 TA+5 VIII Hoard+J.L. ET AL., J.AM.CMEM.SOC. 76.3820(1954) NA3 TA FR TB+3 VI TEMPLETON,D.H. AND C.H.DAUBEN, J.AM.CHEM.SOC, 76,5237(1954) 782 03 TB+3 VIII SHANNON,R.D. AND C.T.PREWITT, TO BE PUBLISHED. R V5 A (PYROCHLORE) TB+4 VI SHANNON,R.D. AND C.T.PREWITT, UNPUBLISHED DATA. R3 V5 V (L12 SN 03) TB+4 VIII WYCKOFF,R.W.G., CRYSTAL STRUCTURES 1,(1963) TB 02 TC+4 VI SHANNON,R.D. AND C.T.PREWITT, TO BE PUBLISHED. R VS & (PYROCHLORE) TE+4 III HANKE+K,, NATURWISS. 54,199(1967) ZN TF 03 TH+4 VI RUH:R, AND A.D.WADSLEY, ACTA CRYST. 21,974(1966) TH TIP 06 H44 YIII Lundrefn,g., Arkiv kemi 2,535(1950) Th (0 H)2 5 04 Lundrefn,g. And L.g.Sillen, Ariv kemi 1,277(1949) Th (0 H)2 CR 04 H2 0 Wyckoff, M.g.S., Crystal Structures 1,1963) Th 02 TH+4 IX ZACHARIASEN,W.H., ACTA CRYST. 2,390(1949) NA TH2 F9 1-3 VI Abrahams, S.C., Phys.Rev, 130,2230(1963) T12 03 Bertaut, E.F. and P.BLUM, ACTA CRYST, 9,121(1956) CA T12 04 Newnaka, R.E. and T.M.Dehaan, 2, XRIST, 117-235(1963) T12 03 Shannon, R.D., TNORG, CHEM, 6,1474(1967) R3 V5 V (PEROVSKITE) SigeglyS, ACTA CRYST, 9,684(1956) T1 F3 V5 V (PEROVSKITE) SigeglyS, ACTA CRYST, 9,684(1956) T1 F3 TI+4 V ANDERSSON-S. AND A.D.WADSLEY, ACTA CHEM.SCAND. 15:663(1961) K2 TI2 05 GUILLEN.M. AND E.F.BERTAUT. COMPT.REND.SER.A.B 262P.962(1966) LI2 TI 05 MOORE:P.B. AND J.LOUISNATHAN, SCIENCE 156:1361(1967) BA2 (TI 0; 572 07 RUDGEFF.C. AND ALCUIDARTHAND SILEY ACTA CRYST. 14,1245(1961) NA2 TI3 07 TI+W VI ANDERSSON.S. AND A.D.WADSLEY, ACTA CRYST. 15,194(1962) NA2 TI3 07 ANDERSSON.S. AND A.D.WADSLEY, ACTA CRYST. 15,194(1962) NA2 TI6 013 BRISSEFF.PH.D.,THESIS.OALHOUSE UNIV.HALIFAX.N.S.(1967) M2 TI2 07 EVANS.H.T.JB.A. ACTA CRYST. 4,377(1951) BA TI 05 FISCHER.JL ET AL., ACTA CRYST. 22,338(1967) CU TI F6.4 M2 0 KNOP.D. ET AL., CAN.JCHEM. 43,2812(1965) ER2 TI2 07 RUM.R. NO A.D.WADSLEY, ACTA CRYST. 21,374(1966) TH TI2 06 SHIRANE'G, ET AL., J.PHYS,50C.JAPAN 14,1352 (1959) M TI 03 SIEGEL.S., ACTA CRYST. 5,635(1952) K 2T IF 6 TEMPLETON.D.H. AND C.H.DAUDEN. J.CHEM.PHYS. 32,1515(1960) RA TI4 09 WEISSAR, ET AL., ACTA CRYST. 20,534(1966) NA3 H TI F8 WYCKOFF.R.W.G., CRYSTAL STRUCTURES 2,(1964) SR TI 03 TL+1 VI SHANNON, R.D. AND C.T. PREWITT, UNPUBLISHED DATA. R VS & (NA 58 F6) TL+1 VIII BROSSET.C., Z.ANORG.ALLGEM.CHEM. 239,301(1938) TL AL F4 SMANNON.R.D. AND C.T.PREWITT, UNPUBLISHED DATA. R3 VS V (SCHEELITE) TL+1 XII Schrewelius,n., arkiv kemi mineral geol, 169,87(1983) TL SR F6 Shannon,r.d. and c.t.premitt, umpublished data. R3 VS V (TL S8 F6) -3 ¥1 Hoppejr, and M.J.Rowrborn, Z.Anorg.Allgew.chem. 329,199(1964) NA TL 02 Johanssonig., Acta Chen.Scandd, 13,925(1959) TL 0 H 5 04 (H2 0)2.5 Mooney:r.C.L., Acta Cryst, 9,113(1956) TL 0 04 Papamantellos;f., Z.RTIST, 126,143(1968) TL2 03 Roth,r.S., S.J.Schweider, J.RtS.Natt.Burs.Std. 644,309(1960) R3 V5 V (MN2 03) TL+3 VI HOPPE,R, TL+3 VIII HOEKSTRA,H.R. AND S.SIEGEL: INORG.CHEM. 7.141(1968) TL? PT? 07 TH+3 VI TEMPLETON,D.H. AND C.H.DAUBEN, J.AM.CHEN.SOC. 76,5237(1954) TM2 03 TH+3 VIII SHANNON, R.D. AND C.T. PREWITT, TO BE PUBLISHED. R VS & (PYROCHLORE) U 43 VI SMANNON.R.D. AND C.T.PREWITT, TO BE PUBLISHED. R3 VS V (LA F3) U +4 VII 24CHARIASEN, W.H., ACTA CRYST, 7,792(1954) K3 U F7

EFFECTIVE IONIC RADII IN OXIDES AND FLUORIDES

Table 1(b) (cont.)

U +4 VIII BRUNTONIG., ACTA CRYST. 21-814(1966) LI U F5 BRUNTONIG., J.INORG.NUCL.CHEW. 29,1651(1967) L14 U F8 DURIFIA., ACTA CRYST. 9,533(1986) U GE 04 LARSOWIAC. ET AL., ACTA CRYST. 17.555(1964) U F4 WTCAGFF.R.M.G., CRYSTAL STAUCTURES 14(1963) U 06 ZACHARIASENTW.H.H. ACTA CRYST. 2309(1949) NA2 U F6 +3 VI BARTAM,S.F., INORG,CHEN, 5,749(1965) Y6 U 012.LU5 U 012 BERTAUT,F. AND JIMARESCHAL, COMPT,REND, 257,867(19651) Y AL 03 FERT.A., BULL,SOC,FRANC.MINERAL,CRIST, 85,267(19621) Y2 03 HARRISILA, ND H.LIYAKEL,ACTA CRYST, 22,354(1967) Y2 05 MULLER-BUSCHBAUMIH, Z,ANORG,ALLGEM,CHEM, 357,138(1968R) SR Y2 09 PATON,M.S. AND E.N.,MASLEN, ACTA CRYST, 19,307(1965) Y2 03 SHANNON,R.C. AND E.N., PREWITT, TO BE PUBLISHED. R VS A (PYROCHLORE) U +4 1X ZACHARIASEN, W.H., ACTA CRYST, 2,390(1949) U2 F9 SHARADATICS THE GALL AND GALL U +5 VI Rošenzweig,A. And D.T.CROMER, Acta Cryst. 23,865(1967) CS U F6 Zachariasen,W.H., Acta Cryst. 2,390(1949) K2 U F6 Zachariasen,W.H., Acta Cryst. 2,390(1949) Naz U F6 U +5 VII LOOPSTRA,B.O., ACTA CRYST, 17,651(1964) U3 08 ZACHARIASEN,W.H., ACTA CRYST, 2,296(1949) U F5 U +6 II Zachariasen.w.H., Acta Cryst, 7,783(1954) K3 U 02 F5 Y +3 IX FRICKE+R. AND W.DURRWACHTER+ Z.ANORG.ALLGEM.CHEM. 259:305(1949) Y (0 H)3 U +6 IV Rossim, et al., am.mineralogist 49,1603(1964) CU (U 02 P 04)2.8 H2 0 YB+3 YI FERT,A., BULL.SOC.FRANC.MINERAL.CRIST. 85,267(1962) YB2 03 TEMPLETON.D.H. AND C.H.DAUBEN. J.AM.CHEM.SOC. 76,5237(1954) YB2 03 U +6 VI BARTANS,F., INORG.CHEM. 5,719(1966) LUG U 012 DEBETS:P.C., ACTA CRYST, 21:589(1966) U 03 RIETVELD:H.H., ACTA CRYST, 20:508(1966) H3 U 06 SIEGELS:N, AND HR., NGEXSTRA, ACTA CRYST, B2:967(1968) CU U 04 SIEARMS:H., ACTA CRYST, 23:263(1967) PB3 U 06 WYCKOFF.R.M.G., CRYSTA, SIRGUERS 2,(1964) U 03 ZACHARIASENIW.H., ACTA CRYST, 7,788(1954) H6 U2 04 YB+3 VIII Eulerf, and J.A.Bruce, acta cryst, 19,971(1965) YB3 M5 D17 Milligan, W.O. And L.W.YERNON, J.Phys.Chew, 56,145(1952) YB V O% Shannon, R.O. And C.T.PREWITT, TO BE PUBLISHED. R VS A (PYROCHLORE) ZN+2 IV ANSELG.6.B. AND L.KATT. ACTA CRYST. 21.442(1966) ZN2 MO3 08 ANSEVO.C. J.PHYS.CHEM.50LIDS 20,114(1983) ZN3 FD 0812 COMEYIG.8. AND R.W.GWYKOFF.ZKATST. 86:41933) ZN1 KO H12 PREWITT.C.T. ET AL., Z.KRIST. 1324.15(1967) PB ZN SI 04 RENYZEPERISIP.J., Z.KRIST. 13117(1963) ZN2 MN 10 H12 SI 04 SCHNERING.H.G. AND R.HOPPE. Z.ANDRG.ALLER.CHEM. 312:87(1961) SR ZN 02 U +6 VII DEBETSIP.C.+ ACTA CRYST. 21+589(1966) U 03 LOOPSTRA.BLO., ACTA CRYST. 17+651(1964) U3 08 SIEGEL+S. ET AL.+ ACTA CRYST. 20+292(1966) U 03 SCHNERING.H.G. ET AL., Z.ANORG.ALLGEM.CHEM. 305.281119601 RA ZN 02 SMITHP.ET AL., Z.KRIST. 119,37511964) ZNW 0 (R 02)6 STITSBERGER.U., ACTA CHYST. 33.19711960) RA ZN 02 STEPHENSIJ.S. AND C.CALVO, CAN.J.CHEM. 45,230311967) ZN3 (P 04)2 V +2 VI MONTGOMERYIH, ET AL., ACTA CRYST, 22,775(1967) V (N Hu)2 (S 04)2,6 H2 O WYCKOFF.R.W.G., CRYSTAL STAUCTURES 1,1363) V O V -3 VI HEPWORTHIMIA, ET AL., ACTA CRYST, 10,63(1957) V F3 NEWNAMIRE, AND Y.H.DEHAAN, Z.KRIST, 117,625(1962) V2 03 SHANNOH, RO., INORGO.CHEM, 6.1974(1967) R3 VS V (PEPOVSVITE) WYCKOFF, R.W.G., CRYSTAL STRUCTURES 2,(1964) CE V 03 ZN+2 ¥ 42 4 ABRANANG,5.C., J.CHEW,PHYS. 66,2052(1967) ZN M0 04 COCCOG, ET AL., Z.KRIST. 123,321(1966) ZN2 (0 H) P 04 MANKEK, NATUWNISS. Soij99(1967) ZN TE 03 MONTGOMERY,H. AND E.C.LINGAFELTER, ACTA CRYST. 16,746(1963) ZN (C5 H7 02)2 H² 0 PLIETHYK. AND G.SANGER, Z.KRIST, 124,91(1967) ZN2 (U AS2 0A STEPHENSJJS. AND C.CALVO CANJLCHEM, 45,2303(1967) ZN3 (P 04)2 V + 4 VI ANDERSSONIG, ACTA CHEM,SCAND, 10,623(1956) V 02 XIERREGAARDIP, AMD J.LONGO, UMPUBLESHED DATA, V 02 STANNON,A.D, AMD CJ.PREMITT, TO BE PUBLESHED, R3 VS V (RUTILE) WTCKOFF,R.W.G., CRYSTAL STRUCTURES 2,(1964) CA V 03 2N=2 VI ABRAYANS,S.C., J.CHEM, PHYS, 46,2052(1967) 2N MO 04 ANSELL-65.B. AND L.KATZ. ACTA CRYST, 21,482(1966) 2N2 MO3 0A BATES,C.H. ET AL.. SCIENCE 137,993(1962) 2N 0 BAURRIKH.A. ACTA CRYST, 11,488(1958) 2N F2 BYSTROM.A. ET AL.. ARKIY KEMI:MIMERAL.GEOL, 1584(1942) 2N 582 06 CALVOC., CAN.J.CHEM. 457(197(1965) 2N3 (P 04)2 CALVOC., J.PHYS.CHEM. 457(197(1965) 2N3 (P 04)2 CALVOC., J.PHYS.CHEM.SOLIDS 24,141(1963) 2N3 (P 04)2 CALVOC., J.PHYS.CHEM.SOLIDS 24,142(1963) 2N3 (P 04)2 CALVOC., J.PHYS.CHEM.SOLIDS 24,144(1963) 2N3 (P /** IV AU-P.K.L. 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ET AL., INORG.CHEM. 5/2131(1966) V O 10 C H3)3 EVAN5H.T.JR., AND S.BLOCK, INORG.CHEM. 5/1808(1966) K V3 OB EVAN5H.T.JR., AND S.BLOCK, INORG.CHEM. 5/1808(1966) C V3 OB SMALLOWA.G., ET AL., ACTA (RTST: 21.397(1966) C X3 V10 O20,17 H2 O ZR+4 VII LUNDGREN,G., ARKIV KEMI 13,59(1958) ZR4 [0 H]6 [CR 04)5,2 H2 0 SMITH-D_K. AND H.M.NEMKIRK, ACTA CRYST, 18,983(1965) ZR 02 N +& VI Shannon,R.D. and C.T.PREWITT, TO BE PUBLISHED. R3 VS V (PUTILE) R+4 VIII BODELH, AND G.TEUFER, ACTA CRYST, 9+929(1956) K2 ZR F6 BUBRS,J,H, ET AL,, ACTA CRYST, B24:230(1968) NAT ZR6 F31 KRSTANOVICTI,R., ACTA CRYST, 11:461(964) ZR (1 03)4 LARSONHA,C, AND J.T. CROMER, ACTA CRYST, 14:128(1964) ZR (1 03)4 SERREJ,R. AND J.H. BUNRS, J.CHEM.PWYS, 41:3578(1964) LIG RE F4 ZR F8 SINWERJ,J. AND J.T. CROMER, ACTA CRYST, 12:719(1959) ZR (5 04)2,4 H2 0 TEUFERG, ACTA CRYST, 15:5187(1962) ZR (5 04)2,4 H2 0 Υ. 46 IV ARRAHAMS'S.C. AND J.L.BERNSTEIN; J.CHEM.PMYS. 45;2745(1966) SC2 (V D0)3 BURBANK:R.D., ACTA CRYST. 18:88(1965) CA W O4 TEMPLETON-DH. AND A.ZALKIN, ACTA GRYST. 16:762(1963) FU2 (V O4)3 ZACHARTASENWIH, AND DH.TEMPLETON. J.CHEW.PMYS. 40350(1964) L12 W OU ZALKINA: AND D.H.TEMPLETON. J.CHEW.PMYS. 40350(1964) CA W O4 ₩ +6 VI KEESINGAARDIP., ACTA CRYST, 10,209(1957) NI ₩ 06 KEESINGAARDIP., ACTA CHEM,SCAND, 12,1715(1958) ₩ 0 ₽2 07 LOOPSTRA.B.O. AND P.BOLDRINI, ACTA CRYST, 21,158(1966) ₩ 03 UCKUN.D., Z.KRIST, 124-192(1967) FE ₩ 06

* This is not intended to be an exhaustive list of references for each ion. For example, there are many references in the literature to certain ions in the usual coordinations such as ${}^{VI}Al^{3+}$ and some of these have been omitted to save space. For those ions and coordinations which occur rarely such as ${}^{VC}r^{5+}$ or ${}^{V}Zn^{2+}$, the list is apt to be more complete.

 Table 2. Anion radii for different coordination numbers

	02-		F -
CN	Radius, Å	CN	Radius, Å
п	1·35 (1·349±5)	II	1.285(1.285+9)
Ш	1·36 (1·357 ± 10)	III	1.29(1.300+12)
IV	$1.38(1.378\pm 4)$	IV	1.31
VI	$1.40(1.396\pm 9)$	VI	1.33
VIII	1.42		

In the next step appropriate anion radii were subtracted from average distances in cation-anion polyhedra. Approximately 1000 average interatomic distances in oxides and fluorides were taken from more than 700 structure determinations. Values of radii for the trivalent rare-earth ions determined in this way agreed quite well with those of Templeton & Dauben (1954). Since our method was similar to theirs, the Templeton & Dauben values, quoted to three decimal places, were used without revision. Since anion coordination is frequently not discussed in structure papers, the following system (see also Holser, 1959) was used to determine the proper values for a compound ${}^{P}A_{p}{}^{Q}B_{q}{}^{R}X_{r}$: pP+qQ=rR, where P and Q are the CN's of the cations and R is the CN of the anion. Table 1(b) lists, by ion according to CN and spin, (where applicable) all the references from which data were taken to construct Table 1. With most references, the source of data is indicated by a chemical formula, compound type, or radius plot (see below). Structure determinations from 1930 through early 1968 were covered in the literature search. In general, many qualitative judgements were made as to the weight given to a particular interatomic distance. More recent structure determinations not involving film methods, those reporting low R values, and those made on well-characterized crystals were given the most weight. The set of values thus obtained was considered to be a first approximation.

The next step was to plot ionic volume, r^3 , vs. unit cell volume, V, for more than 60 isotypic series. According to assumption 6 these plots should be linear. Fig. 1 shows the plot for the rutile MO₂ structure (Shannon, 1968). This example is chosen because the rutile structure covers a wide range of cation radii,



Fig. 1. Cell volume vs. r³ for compounds with the rutile structure. Circles represent data from structure refinements, and the horizontal lines are for compositions where only the cell volumes are known (after Shannon, 1968).

0.39–0.78 Å, and unit-cell volumes are accurately known. The radii were adjusted to provide linearity and consistency with crystal structure data. The radii were further adjusted to be consistent with both crystallographic data and unit-cell volume data over those structures containing common ions. A severe limitation, however, was found to be the accuracy of many reported cell dimensions. Numerous examples which first appeared to deviate from linearity were found to fall in line with other data after cell dimensions were redetermined with least-squares refinement using Guinier camera data.

No structure data are available for many ions. In such cases, a reliable estimated value can often be obtained from plots of radii vs. cell volume (Fig. 1), radii vs. oxidation state, and radii vs. electron configuration (Figs. 3 & 4). Examples of radii determined by radii-volume plots are Ni³⁺ and Co³⁺ LS [perovskite (Shannon, 1967)], Rh⁴⁺, Pt⁴⁺, Re⁴⁺, Os⁴⁺, W⁴⁺, and Ta⁴⁺ [rutile (Rogers, Shannon, Sleight & Gillson, 1969)] (Fig. 1), and Rh³⁺ [perovskites (Shannon, 1967) and corundum (Prewitt, Shannon, Rogers & Sleight, to be published)]. The accuracy of these values depends on the accuracy of the cell dimensions and on the linearity of the plot.

Plots of radii vs. CN (Fig.2) and radii vs. oxidation state are generally found to be regular. This is consistent with the equations given by Zachariasen (1931) and Pauling (1927). Consequently, when the values of radii as determined by structural data appeared to be inconsistent with these regular curves, the sources of data were investigated; if these appeared questionable, the value of the radius was made consistent with the radii plots and marked in Table 1(a) with an E, e.g. Nb³⁺.

The variation in radii of the first-row transition metal ions with the number of e_g and t_{2g} electrons has been discussed qualitatively by van Santen & van Wieringen (1952), calculated theoretically by Hush & Pryce (1958), and plotted experimentally by Liehr (1960), Knox (1961), Blasse (1965), and Pearson (1968). Similar plots were made for our data, Figs. 3 and 4. By use of these plots, hypothetical values of radii were determined for the low spin ions, Mn^{2+} , Ni^{3+} , Fe^{3+} , Mn^{3+} and Co^{3+} . Low-spin radii for Fe^{2+} and Co^{2+} were taken from radii vs. volume plots for the pyrite structure; these radii were consistent with the above electron configuration plots.

As a final step, the anion radii with CN II, III, IV, VI, and VIII were recalculated using interatomic distances from the simple oxide structures and the final refined set of cation radii. These values, given in parentheses in Table 2 with their estimated standard deviations do not differ significantly from the original values. It was not possible to calculate average values for $^{VIII}O^{2-}$ because of the lack of examples. No calculations for $^{IV}F^-$ or $^{VI}F^-$ were made because of a problem with highly symmetric structures discussed in another section.

Results

Table 1(*a*) presents three sets of radii. The first is that of Ahrens (A–IR). The second column contains crystal radii as defined by Fumi & Tosi (CR). This column was derived from column three but it is based on $r(^{VI}F^{-})=1.19$ Å and an assumed difference between six-coordinated oxygen and fluorine radii of 0.07 Å as was found for the ionic radii. The third column contains the radii obtained in this work for various coordination numbers and spin states. Except for the radii of S^{2-} , Se^{2-} , and Te^{2-} , which are included in column one and marked with *P*, Pauling's radii are omitted because Ahrens' radii are taken from revised values of Pauling. Each row in Table 1 (*a*) contains the ion, the electron configuration (EC), coordination number (Roman nume-





ral), spin state (HS or LS), three values of ionic radii, and finally a symbol giving some indication of the reliability (?, E, R, *, or no symbol). The values indicated by a question mark are considered doubtful because of one or more reasons: uncertainty as to oxidation state,



Fig. 3. 'Effective' ionic radius vs. number of 3d electrons for divalent transition-metal ions.



Fig. 4. 'Effective' ionic radius vs. number of 3d electrons for trivalent transition-metal ions.



Fig. 5. Si-O distance vs. oxygen coordination. The dashed line represents the variation of Si-O distances reported by Smith & Bailey (1963).

CN, or composition of the compounds studied, or deviation from radii vs. CN, or radii vs. valence plots. The values estimated from radii vs. CN, radii vs. oxidation state, or radii vs. electron configuration plots are followed by the letter E. Values of radii taken from radii vs. volume plots are marked by R. Finally, those values for which at least 5 structural determinations resulted in radii differing by no more than ± 0.01 Å are marked with an asterisk.

Although many different types of coordination exist for a given CN, for simplicity we have chosen to differentiate only the common types of four coordination: square planar (IVSQ), square pyramidal (IVPY), and tetrahedral (IV).

Table 3 compares observed and calculated interatomic distances in more than 100 compounds with the use of radii from Zachariasen, Pauling, Ahrens, and Table 1(a). The first three columns in Table 3 contain calculated interatomic distances of Zachariasen, Pauling, and Ahrens corrected for cation coordination using the correction factors listed by Pauling (1960) and Zachariasen (1931). The agreement between observed and calculated distances is seen to be much improved in most cases where CN of the cation and/or anion is different from VI.

When the distances calculated from Table 1(a) differ significantly from the observed distances, a reason for the discrepancy can usually be found. One of these reasons is that interatomic distances in highly symmetric structures are generally shorter than expected and another is that it is often difficult to establish exact coordination numbers for some ions. These effects are discussed further in later sections of this paper. Other reasons are that some radii are based on insufficient data and/or that a structure solution and refinement may be, to some degree, in error. In addition there are undoubtedly some unexplained departures from additivity which are reflected in deviations of the order of ± 0.01 Å.

Discussion

Change of anion radius with CN

Although the dependence of interatomic distance on CN was noted by Goldschmidt (1927) and the idea of increases of cation radii with CN has been mentioned by several investigators (Pauling, 1927; Zachariasen, 1931; Geller, 1957; Sasvári, 1960), the possibility that anion radii may also depend upon CN appears to have been considered by only a few authors (Slaughter, 1966; Jeffrey & Slaughter, 1963). The effect of CN on anion radii is not great, but it is noticeable when comparing interatomic distances in structures containing a series of M-O distances where the CN of O^{2–} varies (Tables 3 and 4). This is probably the principal cause of the variations in Si–O distances noted by Smith & Bailey (1963) in their survey of silicate structures.*

^{*} This effect was recently discovered independently by Gibbs & Brown (1968).

Table 3. Comparison of observed and calculated interatomic distances in oxides and fluorides

			RC	alc							RC	alc			
Compound	Distance	Zach.	Pauling	Ahrens	S & P	RODB	Year	Compound	Distance	Zach.	Pauling	Ahrens	<u>S & P</u>	RObs 1	<u>fear</u>
AgFeO	IIAg-IV0		2,36	2,40	2.05	2,06	68	KP	VIK-VIF	2.66	2.69	2,66	2.71	2,673	63
ALSIO	IVAL-III	1.78	1.78	1.81	1.75	1.77	63	KMnO _{li}	VIII K-III O	2.90	2.83	2.82	2,86	2.89	67
AlPO.	IVAL-II	1.78	1.78	1.81	1.74	1.74	66	L1 WO	IV _{L1-} III ₀	1.99	1,84	1.91	1.95	1.96	61
A1.0_	VIAL-IVO	1.90	1.90	1.91	1.91	1.91	62	LIA10	IVLI_IVO	1.99	1.84	1,91	1.97	2.00	65
2 3	IVAL IVO	1.78	1.78	1.81	1.77	1.76	65	LiGaOo	IV _{L1} IV _O	1.99	1.84	1.91	1.97	1.98	65
L(A)0	VIALVIO	1 00	1 00	1 01	1 03	1 00	66	14 St 0	IV III	1 99	1.84	1.91	1.95	1.94	61
Panto2	VIA IV	1.90	1.90	1.91	1.01	1.90	63	201205	VI, VI	0.10	2.00	2 08	2 14	2.12	66
AT204	VIA II	1,90	1.90	1.91	1.91	1,914	03	L1A102	VI, VI	2,12	2,00	2.00	2.14	214	65
A1F3	IV. II.	1.((1.00	1.04	1.015	1.00	51	LIGE02	VI. VI.	2.12	2.00	2.00	2.14	2,14 0,008	63
As205*5/3H20	AS- U III III		1.70	1.75	1,005	1.09	00		VI. II	2,01	1.90	2.01	2.01	2.000	60
CdB407	<u></u> В0 ТТТ ІТ	1.35	1.38	1.40	1.38	1.37	66	L14 ^{UF8}	L1- F VI TII	2,01	1.96	5.01	2.025	2.00	67
CaB204		1.35	1.38	1.40	1.37	1.37	63	L1UF5	VT VT	2.01	1.96	2.01	2.04	2,055	66
L1B02	ту тт	1,42	1.47	1.50	1.50	1.47	65	MgO	Mg- O VT TV	2.10	2.05	2,06	2.12	2,105	63
Y-HBO2	1°B-110	1.42	1.47	1.50	1.47	1.47	63	мg504 4H20	'14g-1'0	2.10	2.05	2.06	2.10	2.076	62
KB508	1 ⁰ B-1110	1,42	1.47	1.40	1,48	1.46	65	MgF2	Mg-TTF	1.98	5.01	1.99	5.05	1.992	56
BaO	Ba-VIO	2.76	2.75	2.74	2.76	2.76	63	MgHPO4 · 3H20	Mg-1110	2,10	2.05	2.06	2,08	2.084	67
BaGe03	VIII Ba-IVO	2.84	2.82	2.80	2,80	2.785	62	MnMo04	VI HS Mn-III		2.20	2.20	2,18	2.164	65
Bar	VIII Ba-IVF	2,68	2.79	2.74	2.73	2.69	63	Mg6Mn08	VI Mn-VO		1.94	2,20	1,93	1,928	54
Ba(OH), 8H20	VIIIBa-IVO	2.84	2,82	2,80	2.80	2.74	64	DyMn ₂ 05	VIMn-IVO		1.94	2.20	1,92	1.92	67
BaGe,,O	X Ba-IIIO	2,90	2.88	2.84	2,88	2,96	66	MoO2	VIMO-IIIO		2.06	2.10	2,01	2.011	67
BaSO _{li}	XII _{Ba-} IV _O	2,96	2.94	2,89	2,98	2,95	63	ZnoMooOg	VI MO-IIIO		2,06	2,10	2.02	2.02	66
Baa(PO _k)a	XII Ba-VO	2.96	2.94	2.89	2.99	3.01	48	MnMoO,	IV MO-IIIO	1.82	1.94	1,92	1.78	1.763	65
Y Be0.	IIIIV	1.55	1.47	1.51	1.55	1.55	67	PhMoQ.	IV III	1.82	1.94	1.92	1.78	1.772	65
*20004 Bao	IV	1.63	3 67	1.51	1.60	2.50		NoR	VI VI	0.21	0.21	0.20	0.25	0 31	63
Deo IA Deb	IV_ III	1.05	1.57	1.01	1.05	1.049	- 04 cc	Nar Na Giri	VI III	2.51	2.31	2.30	0.20	0.30	61
¹ 2 ^{ber} 4	VI_ III.	1.55	1.55	1.54	1.57	1,554	. 00	Ma2S1F6	VI. IV.	2.34	2.31	2.30	c.jc	2.Je	64 67
^{B1} 4 ^{S1} 3 ⁰ 12	VT TT			2,36	2.38	2,385	66	NAHCO3	TNA-TO	2,44	2,35	2.37	2,40	2.44	65
^{B1} 2 ^{Ge0} 5	B1-TTO			2.36	2.38	2.35	64	Nb205	VT TTT	2,00	2,10	2.09	1,99	1.99	64
Ca25104	VIII TIT	2.50	2.47	2.48	2.50	2.515	64	Gando 4	VT TV	2,00	2,10	2.09	2.00	2.031	65
CaWO4	VIII Ca-1110	2.50	2.47	2.48	2.48	2,461	65	LINDO3	VIND-IVO	2.00	2.10	2.09	2.02	2.00	66
CaB204	VIIICa-1110	2,50	2,47	2.48	2.48	2,505	63	N1P2	VIN1-IIIF		2.08	2,02	2.00	2.00	58
CaF2	VIIICa-IVF	2.34	2.43	2.40	2.43	2.37	63	niso ₄ •6h ₂ 0	VIN1-1110	~ •	2.12	2,09	2.06	2.06	66
CaSO	Ca-1110	2.50	2.48	2.48	2.48	2.46	63	Ba2NIF6	VINI-VF		2.08	2,02	5.05	2.01	67
caso	IV _{cd-} II ₀		5.58	2.28	2.19	2,168	63	Pb02	VI _{Pb-} III _O		2.24	2.24	2.135	2.16	47
CdO	VICd-VIO		2.37	2.37	2.35	2,348	63	Ru02	VI _{Ru-} III _O		2.03	2.07	1.98	1.971	66
Cd ₂ V207	VI _{Cd-} III _O		2,37	2,37	2,31	2,31	67	LisbF	VI _{Sb-} II _F		1.98	1.95	1,895	1.88	62
CePO,	VIII _{Ce-} III ₀		2.57	2.53	2.50	2,50	50	Sb_0,	VI _{Sb-} II ₀		2,02	2.02	1,96	1.97	64
CoF	VI HSCO-IIIF		2,10	2.05	2.035	2.04	58	KaSbr0.	VI _{Sb-} IVo		2,02	2.02	1.99	1,984	66
CoMoO,	VI HSCO-IIIO		2.14	2.12	2.095	2.08	65	5 5 14 Se_0	VI _{Sc-} IVo	2.16	2.21	2.21	2.11	2.11	67
4 CoQ	VI HSCO_VIO		2.14	2.12	2.135	2.133	63	2-3 Sc. (WOw).	VISC_II	2.16	2.21	2.21	2.08	2,063	66
Cr 0	VI IV		2.00	2.03	1.005	1.00	62	2(4/3 CaSc 0.	VISC V	2.16	2,21	2.21	2.12	2.12	65
°-2°3	VI II		2.00	2.02	1 065	1.07	65	Sall 204	VICII	2 02	2 17	2 14	2 015	2 01	51
015 12 CmP	VICLII		2.09	1.06	1.905	1.00	60	SoP0	VIII	2.02	2 28	2 20	2.01	0.03	56
⁰¹¹ 3.	IV_ III		2.09	1.90	1.90	1.90	60	BCF04	IV., II.	1 69	1 21	2 70	1 61	1 607	50
100104	IV_ III	1.05	1,02	1,02	1.00	1.05	60	S102	IV, III,	1.03		1.70	1.01	1.007	24
110r308	II. IV.	1.05	1,02	1.02	1.00	1.05	ογ·	KAIS1308	IV. IV.	1.03	1.11	1.(2	1.02	1.01	64
CUNEO 2	VIII IV.		5.01	5.10	1.04	1.03	00	Ca25104	VI. III.	1.03	1./1	1.72	1.04	1.044	05
^{Dy} 3 ^{Fe} 5 ⁰ 12	Dy- 0			5.38	2,40	2,40	65	S102	S1- 0	1.73	1.81	1.65	1.75	1.768	60
Eu304	VIEu-VIO		2,43	2.38	2.35	2.340	66	K ₃ SiF7	VIS1-IVF	1.60	1.77	1.75	1.71	1.70	62
FeO	VI _{Fe-} VIO		2,16	2.14	2,17	2,155	63	Na ₂ S1F ₆	VI _{S1-} III _F	1,60	1.77	1.75	1.70	1.695	64
FeWO ₁₁	VI _{Fe-} III _O		2,16	2.14	2.13	2.104	67	(NHA) SIF6	VI _{S1-} II _F	1.60	1.77	1.75	1,685	1.674	56
Fe_SiO	VI _{Fe-} IVo		2.16	2.14	2.15	2.172	65	Sn0	VI Sn-III O		2,11	2,11	2.05	2.053	56
FeGe (OH)	VI _{Fe-} III ₀		2,16	2.14	2.12	2.13	61	SnFh	VI _{Sn-} II _F		2.07	2.04	1.975	2.14	62
Y_Fe_0	IV _{Fe-} IV ₀		1,94	1.94	1.88	1,88	62	SrF ₂	VIII _{Sr-} IV _F	2.54	2,56	2.52	2.56	2,51	63
Ne Fe O	IV _{Fe-} IV ₀		1,94	1.94	1,88	1,867	67	Tipos	VI _{T1-} IVO		2,16	2.16	2.05	2.040	62
reo	VIIV		റവം	ອດພ	9 095	2 030	66	- <u>-</u>	VIII		2 12	2 09	1 055	1.07	56
20	VI. II.		2.04	2.04	2.02)	2.030		1143	VI_ III	2.00	0.09	0.00		+• 31	
3	VI III		2,00	1,97	1.93	1.92	57	T102	VI IV	1.96	2.08	2.08	1.905	1.959	50
a-Ga. 0	VIGa_IV		2.02	2.02	7.90	2,00	, 05 65	Na-T1-0-	VITI VO	1.90	2.08	2,08	1.905	1.99	61
	VIGa_IV		2.02	2.02	1.080	1,005	5 65	V-0-	vi, iv		2.14	2,14	2.02	2,01	62
-3-5-12 Gal	VI _{Ca} II		1 09	1 05	1 005	1 20	, U) E0	*2~3 No.	VI III	_	5 M	2 02	1 05	1 097	52
BaGe. 0	VIGe_III		1.03	1.03	1,000	1,877	- 66 - 66	1~2 Cd-V-0-	IV_III	1 77	1.80	1,80	1.715	1.717	در 67
Na. Ge 0	vige_III		1,02	1.03	1,010	1,000	69	2-2-7	VI III			2.02	1.94	1.04	67
49-20 To 0-	VITTIN		2.01	2.01	2.17	JU(66	¥0-	VILII			2.02	1.02	1.027	RR.
	vITv		2.01	2.01	2.18	2.10	67	"°3 ¥-0-	៴៲៓៓៲៷	2 32	2 33	2,32		2.97	62
InPO, *2H.0	VI IN IIO		2,21	2,21	2.14	2,16	61	*2*3 ZnF	VIZn-III		2.10	2.07	2.045	2.03	58
4 2 Ca_IrO.	VI IV		2.04	2.08	2.01	2.01	66	2 Z=0	VIII2 IV	2.91	2:26	2.26	2.22	2.26	62
24							~~	2							

Fig. 5 shows how the average Si-O distance would vary with average oxygen coordination in silicates although most silicates fall in the range ^{II}O–^{IV}O as indicated by the dashed line which is taken from the Smith & Bailey work.

The change in anion radii with CN is much smaller in magnitude $[r(^{VI}O^{2-})-r(^{II}O^{2-})=0.05 \text{ Å}]$ than the change in radii with cation CN $[r(^{VIII}Cd^{2+})-r(^{IV}Cd^{2+})]$ = 0.26 Å; $r(^{VIII}Mg^{2+}) - r(^{IV}Mg^{2+}) = 0.41$ Å]. One explanation for this effect involves electronic polarizabilities of cations and anions. Intuitively, one might expect the change in radius of an ion with CN to be roughly proportional to its electronic polarizability. A comparison of cation and anion polarizabilities from various sources made by Tessman, Kahn & Shockley (1953) shows generally that polarizabilities of

 Table 4. Comparison of observed and calculated interatomic distances in oxides and fluorides with varying coordination

Distance	CN of 02-	Compound	Robs.	R _{calc.}	Distance	CN of 02-	Compound	Robs.	Rcalc.
IVA1-0	11	LIAISILO	1.720	1.74	VI Sc-O	11	Sc2 (W04)3	2.063	2.08
		AlPon	1.739	n		III	Y-SCOOH	2,10	2.09
	III	KAlSizOn (microcline)	1.741	1.75			Sc(OH)2	2.08	tt
		NaAlSipOn (low albite)	1.737	11		IV	Scoog	2.11	2.11
		AloSiO _c (sillimanite)	1.770	17		v	Case_Oh	2,122	2.12
	IV	LiAlo	1.761	1.77		VI	KScOo	2,12	2.13
VI.		N 12 (20:) - 2011 0					Lisco	2.11	81
A1-0	11	NAA1 (SO4)2*12H20	1.001	1.00			RbScO2	2.12	tı
	~ ~ ~	CBA1(SO4)2-12H20	1.002	1 00	IV		ELO (questa)	1 607	1 610
	111	Al ₂ Si0 ₅ (kyanite)	1.905	1.09	51-0	11	SiO_2 (quartz)	1.605	"
	IV	Mg3 ^{A12^{S1}3⁰12}	1.885	1,91			SiO ₂ (10w cristobalite)	1.009	
		^{A1} 2 ⁰ 3	1.91	1.91				1.013	"
	VI	LIA102	1.90	1.93			Al25105 (Sillmanite)	1.015	r
								1.600	1 600
VICd-0	II	3caso4.8H20	2.30	2,30		111	50251207	1.610	"
	III	, cd ₂ v ₂ 07	2,31	2.31			275104	1.608	
	VI	.060	2.348	2.35			Al Sto (andolugita)	1 600	n
VIco2+-0	II	CoSOJI	2.07	2.085			72 ND(04) - 510	1 626	v
	III	Сомоод	2,082	2.095			AlBest O. OH	1 625	
		Co(NH2)2(SO4)2.6H20	2,094	8			Net1St-00	1.618	11
	VI	Co0	2.133	2.135			NA284-00	1 61 2	n
VI. 3+ -			1 070	2.065			KAIDI308	1 631	"
-Cr ³ -0	11	cr5012	1,972	1.905			L1251205	1.630	II.
	111	KCF3012	1.97	1.9/5			Ca2Man51309	1.050	
		HCF02	1.919	1 005			CaMnSi206	1,623	
	14	CF203	1.99	1.999		τv	~-Ca-SiQ1	1.644	1.640
V1Fe ²⁺ -0	III	FeW04	2.104	2.13		11	1-0022204 Mg-S10	1.625	
		FeGe(OH) ₆	2.14	10			MgFeS10.	1.638	
	IV	FeSiF6'6H20	2.140	2.15			Fe-S10.	1.634	11
		Fe2S104	2.172	11			-2^{4} Cs_Al_c(S10)	1.649	н
	VI	FeO	2.155	2,170			Mg_AloS1o010	1.635	н
VIGa ³⁺ -0	III	Gando _{ll}	1.997	1.98			103-20-3-12		
		B-Ga203	2.00	n	VT				
	IV	a-Ga203	2.00	2.00	*\$1-F	II	(NH ₄) ₂ 51F ₆	1.674	1.685
	VI	LiGa02	2.00	2.02			[CO(NH3)5C1]S1F6	1.654	
IV.	**	600	1 730	1.75		III	Na ₂ 51F6	1.095	1.700
66-0	.11	Becer 0-	1 740	1.76		IV	^k 3 ^{81F} 7	1.70	1.710
	111	No. 0:000 . 6H 0	1.769	"			(NH4)3 ^{31F7}	1.71	
		Na-SniGarolog(OH)	1.752	H	VI _{T1} 44-0	11	Na2T16013	1.957	1.955
	717	Pageo	1 785	1.78			Rb2T16013	1.983	Ħ
	14	Badeo3	11,00			III	ThT1206	1.970	1,965
VIN12+-0	III	N1S04• бн ₂ 0	2,063	2.06			T102	1.959	ti -
		niwo4	2.08	n		IV	NITIO3	1.99	1.985
	VI	NIO	2.084	2.08			MnTiO3	1.99	91
IV_5+_0	II	Mopo _{ll}	1.528	1.52			BaTi409	1.98	12
		ALPO4	1.515	11		v	NB2TI307	1.992	1,995
		P205	1.54	11					
	III	Na2H2P4012	1.542	1.53					
		NaPO ₂ NH ₂	1.528	n					

anions are larger than those of most cations, in distinct contrast to the relative change in anion and cation radii with CN. However, Ruffa (1964) has shown in TiO_2 and other compounds with high Madelung potentials that the polarizability of the cation may be considerably greater than that of the anion. Thus, the possibility still exists that there may be some correlation between change in radius with CN and polarizability.

An alternative explanation for this effect depends on the anion-anion repulsive forces described by Pauling (1928). In most of the oxides for which reliable radii have been determined, the cation/anion radius ratio, q, is considerably less than 1. When this condition holds, anion-anion forces can be large. As ρ becomes small and as the anion CN diminishes from VI to II. the nearest neighbor anion interactions become more important. The increase in anion-anion forces prevents the anion from deforming as much as might be expected from polarizability considerations. The cations in such structures, however, cannot approach each other closely and cation-cation repulsion is probably less than anionanion forces. Consequently, the cation is easily able to expand into the void between the anions - thus shortening considerably the cation-anion distance.

Absolute values of radii

The question of absolute values for anion and cation radii is still not resolved. Although it has been possible to devise sets of radii which are almost additive, the values of anionic radii have been determined by approximate methods. This question of the relative sizes of anion and cation radii is very important because it determines the physical picture we have of ionic solids. Traditionally this has been a picture of anion-closepacked solids.

Values of anion radii have been chosen from: (1) molar refractivities of salts in solution (Wasastjerna, 1923); (2) anion-anion distances in close-packed solids (Bragg & West, 1927); (3) Born repulsive coefficients determined from isothermal compressibilities and thermal expansion coefficients (Fumi & Tosi, 1964) and (4) electron density distribution in alkali halides (Witte & Wölfel, 1955; Schoknecht, 1957; and Meisalo & Inkinen, 1967). The traditional sets of radii were based on (1) and (2). The first values of ionic radii were determined by Wasastjerna who used the ratio of the cube root of the molar refractivities of the free ions to determine $r_c:r_a$ in the alkali halides. He arrived at $r(VIF^{-}) = 1.33$ Å and $r(VIO^{2-}) = 1.32$ Å. Bragg & West noted that observed O-O distances in silicates were roughly 2.7 Å and concluded that $r(O^{2-}) = 1.35$ Å. Used with available sets of interatomic distances in common oxides, these values led to small $r_c:r_a$ and resulted in the concept of anion close-packed oxides. Many solid state phenomena such as diffusion, mechanical deformation, kinetics of transformations, and solid state reactions have been interpreted on this basis.

More recent evidence, however, indicates that the ratio $r_c: r_a$ may be considerably larger than indicated by

the traditional sets of radii. Comparative calculations of wave functions of free ions and ions in a crystal by Petrashen, Abarenkov & Kristofel (1960) showed a tightening of electron clouds in anions and a relaxation of cation electrons in solids. Analysis of electron density distributions in NaCl (Witte & Wölfel, 1955), LiF (Krug, Witte & Wölfel, 1955) and KBr (Meisalo & Inkinen, 1967), indicated anion radii and cation radii smaller and larger respectively than traditional ionic radii. Fumi & Tosi on the basis of calculations of Born repulsive parameters from compressibility and thermal expansion data present convincing evidence that $r_c:r_a$ is larger than in the traditional sets of radii. They found $r(VIF^{-}) = 1.15 - 1.20$ Å with an estimated error of 0.05 Å. Assuming a constant difference of $r(VIO^{2-}) - r(VIF^{-}) =$ 0.07 Å, then $r(^{VI}O^{2-}) = 1.22 - 1.27$ Å. These values are more in agreement with the shortest nonbonding O-O distances in many oxides. As shown in Table 5 these range from 2.15 Å in the NaNO₃ structure to greater than 3 Å in the antifluorite structures. Perhaps the most surprising distance in this group is that of the 2.16 Å shared edges of the SiO₆ octahedra in the rutile form of SiO₂ (stishovite). Although extremely high pressure must be used to synthesize stishovite, it is difficult to reconcile this short O-O distance with an $r(^{III}O^{2-}) = 1.36$ Å, especially since Fig. 1 shows there is nothing unusual about the cell volume of stishovite. Furthermore, with smaller anions one is not forced to assume negative radii for IIIC4+ in the carbonate ion and ^{III}N⁵⁺ in the nitrate ion. Although there is intrinsically no objection to negative radii so long as one does not assume a hard sphere model, it is esthetically more pleasing to have positive radii. However, even with Fumi & Tosi's crystal radii it is not possible to achieve a complete set of positive radii. The O-H and F-H distances in hydrogen-bonded compounds result in negative radii of -0.04 and -0.24 Å for πH^+ and \bar{H}^+ . Although the value for $r(^{II}H^+)$ falls within the estimated error of the crystal radii and would still allow a completely positive set of radii, this is not possible for ^IH⁺. One is forced to the interpretation that an ^IH⁺-O bond results in a highly deformed oxygen ion in which the proton penetrates the electron cloud of the oxygen ion. On the basis of the above considerations, the

 Table 5. Shortest oxygen-oxygen distances in some simple oxides

Compound	Structure type	0-0
NaNO ₃	calcite	2·15 Å
SiO ₂	rutile	2.16
CaCO ₃	calcite	2.22
Al ₂ SiO ₅	andalusite	2.25
$B_2O_3(II)$	$B_2O_3(II)$	2.36
S ₃ O ₉	S ₃ O ₉	2.37
$B_2O_3(I)$	$B_2O_3(I)$	2.38
Al ₂ O ₃	corundum	2.52
SiO ₂	quartz	2.60
BeO	wurtzite	2.70
MgO	rocksalt	2.98
Li ₂ O	antifluorite	3.27

Fumi & Tosi crystal radii (CR) are probably more satisfactory than the traditional sets of radii.

Pauling (1928) noted a correlation between radius ratios and the relative stability of certain structures. Values of ϱ for structures with tetrahedral cation coordination were generally of the order of 0.2–0.4 whereas values of ϱ for octahedral cation coordination were of the order of 0.4–0.7. There were exceptions, but in general the radius ratio was found to provide some indication of the type of prevalent coordination of anions about the cation in ionic crystals. Phillips & Williams (1965) noted that the critical radius ratios were not satisfactory in predicting CN changes but their analysis involved highly covalent rather than ionic crystals.

It is clear that any new set of radii should provide a correlation between radius ratios and CN at least as satisfactory as Pauling's table of ionic radii. Table 6 gives values of ρ calculated by means of an average radius of 1.38 Å for oxygen for numerous cations with CN III, IV, VI, and VIII. As long as ρ falls within ± 0.02 of the limits defined by Pauling (1960) the radius ratio is in boldface. It will be noted that values of ρ calculated from ionic radii (IR) do not fall within the expected limits for most of the small ions, e.g. VIA13+, VIAs5+, IIIB3+, IVB3+, IVS6+, IVSi4+, and VISi4+, whereas values of ρ calculated from crystal radii (CR) give poor agreement for ^{IV}Ga³⁺, ^{IV}Li⁺, ^{VI}Mn²⁺, ^{IV}Zn²⁺, and ^{IV}Mo⁶⁺. Since ions with strong covalent mixing (Ga³⁺ and Zn²⁺) prefer four coordination via sp³ hybridization, even though $\rho > 0.414$, it is felt that crystal radii predicts CN on the basis of radius ratio as well as effective ionic radii.

Perhaps the most important conclusion to be drawn from the above arguments is that specific values from this or any other set of radii should not be used too rigidly to derive a physical picture of ionic solids. Currently available information on electron density distributions in solids is not sufficient to establish absolute values of radii. Furthermore, if one assumes that the radius of an ion, say oxygen, is the distance from the oxygen nucleus to the minimum in electron density between the oxygen ion and a cation, this radius may vary depending on the type of cation, *i.e.* on the covalency of the bond. If, however, the crystal radii in Table I(a) are more accurate than the ionic radii, certain concepts of the structures of ionic solids are no longer valid. For example, oxides should not be thought of as close-packed arrangements of anion spheres whose interstices are partly filled with cations. Instead, the cation-anion bonds should be considered as the important factors determining crystal structure and closepacked anion arrangements as merely the result of the structure attaining a lowest-energy configuration.

Interatomic distances in highly symmetric oxides and fluorides

Agreement between observed and calculated distances from Table 1(a) over a wide variety of compounds with varying cation and anion coordination was shown to be quite good. However, the compounds with rocksalt, fluorite, and perovskite structures show rather poor agreement (Table 7). A possible source of error could be the cation radii. However, the value of $r(VIIICa^{2+})$ taken from more than 15 Ca–O distances which agree to within 0.02 Å results in $R_{cale} - R_{obs} =$ 0.06 Å for CaF₂. Therefore, it is felt that the lack of agreement of interatomic distances between symmetric and unsymmetric structures is significant. The effect appears to be greater for the fluorides than for oxides and greater for smaller cations than for larger cations. The smaller observed interatomic distances are qualitatively explained by minimization of anion-anion repulsive forces. The regular polyhedra result in the maximum distances between all cations and anions and,

Table 6. Radius ratios for oxides

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Ion	CN	IR	CR	Ion	CN	IR	CR
A13+	IV	0.28	0.42	Mg ²⁺	VI	0.52	0.69
	VI	0.38	0.54	Mn ²⁺	VI	0.59	0.79
As ⁵⁺	IV	0.24	0.38	M06+	IV	0.30	0.45
	VI	0.36	0.52		VI	0.43	0.60
B3+	III	0.02	0.12	Ni ²⁺	VI	0.51	0.66
	IV	0.09	0.21	P5+	IV	0.12	0.25
Be ²⁺	III	0.12	0.25	S6+	IV	0.09	0.21
	IV	0.20	0.33	Sc ³⁺	VI	0.53	0.70
Ca ²⁺	VIII	0.81	1.01	Si ⁴⁺	IV	0.19	0.32
Ce ⁴⁺	VIII	0.70	0.90		VI	0.30	0.43
Co ²⁺	VI	0.53	0.71	Sn ⁴⁺	VI	0.20	0.67
Cr ³⁺	VI	0.44	0.61	Ti ⁴⁺	VI	0.44	0.60
Fe ²⁺	VI	0.56	0.73	W6+	IV	0.30	0.44
Fe ³⁺	IV	0.36	0.52		VI	0.42	0.58
	VI	0.47	0.63	Y3+	VI	0.65	0.83
Ga ³⁺	IV	0.34	0.49		VIII	0.74	0.93
	VI	0.45	0.61	Zn ²⁺	IV	0.43	0.60
Ge ⁴⁺	IV	0.29	0.44		VI	0.54	0.71
	VI	0.39	0·55	Zr ⁴⁺	VI	0.52	0.69
In ³⁺	VI	0.57	0.75				
Li+	IV	0.43	0.59				
	VI	0.54	0.71				

therefore, the least repulsion interactions. The greater polarizabilities of the oxide ion and larger cations result in deformation of these ions into voids and consequent reductions in repulsive effects.

Table 7. Calculated and observed interatomic distances in highly symmetric oxides and fluorides

	ROCK	salt	
	R _{Calc} ^{M-O}	Robs ^{M-O}	∆R
NiO MgO CoO FeO MnO CdO CaO SrO BaO	2·10 2·12 2·135 2·17 2·22 2·35 2·40 2·56 2·76	2.084 2.105 2.133 2.155 2.222 2.348 2.405 2.58 2.762	$\begin{array}{c} 0.016\\ 0.015\\ 0.002\\ 0.015\\ -0.002\\ +0.002\\ -0.005\\ -0.02\\ -0.002\\ -0.002\end{array}$
LiF NaF AgF KF CsF	2·07 2·35 2·48 2·71 3·03	2·008 2·31 2·46 2·673 3·00	0.062 0.04 0.02 0.037 0.03
	Fluo	rite	
MO_2			
VIII-IV	2.25	2.24	0.01
UO_2 ThO ₂	2·33 2·38 2·44	2·34 2·37 2·42	0.01 0.01 0.02
<i>M</i> F ₂ VIII_IV			
CdF ₂	2.38	2.333	0.023
CaF ₂	2·43	2·37	0.06
SrF ₂	2.56	2.51	0.05
BaF ₂	2.73	2.69	0.04
MBO3 VI–VI orX II–V	Perov I	skite	
BaCeO ₃	2·25	2·198	0.052
CeCrO ₃	2·23 2·015	1.933	0.082
KNbO3	3.00	2.84	· 0·16
KNDO3 BaMoO3	2.04 2.05	2.011	0.029
SrMoO ₃	2.05	1.987	0.063
BaPbO ₃ BaSnO ₁	2.175	2.136	0.039
CaSnO ₃	2.09	1.96	0.13
SrSnO ₃ BaZrOa	2·09 2·12	2·016 2·096	0.074
SrZrO ₃	2.12	2.05	0.07
<i>MB</i> F ₃ VI-VI or XII-V	т		
KCoF1	2.065	2.035	0.030
RbCoF ₃	2.065	2.03	0.035
TICoF ₃	2·065	2·064	0.001
CsMnF ₃	3.25	3-13	0.12
KFeF ₃	2.10	2.06	0.04

	1 able 7	(cont.)	
	Rcale ^{M-O}	$R_{\rm Obs}^{\rm M-O}$	∆R
RbFeF ₃	2.10	2.087	0.013
KMgF ₃	2.05	1.986	0.064
KMnF ₃	2.15	2.095	0.022
KNiF ₃	2.03	2.006	0.024
CsPbF ₃	2.51	2.405	0.102
KZnF ₃	2.075	2.027	0.048
AgZnF ₃	2.075	1.99	0.082

 $\mathbf{T}_{1} \mathbf{1} \mathbf{1}_{2} \mathbf{7} (\dots \mathbf{1}_{k})$

Unusual and irregular coordination

When the distances in polyhedra are almost equal, it is easy to determine the coordination. In this case there is usually a clear cut-off point for which peripheral anions may be considered as bonded to the central atom. However, in many cases there is a gradual increase in near-neighbor distances up to very large interatomic distances for the large alkali and alkaline earth ions in network structures such as silicates; consequently, the CN of these ions is frequently uncertain. The curves of radii vs. CN reflect this uncertainty [Figs. 2(a)and 2(b)]. For the purposes of this paper a break of 0.02-0.05 Å was sought as the cut-off point in the interatomic distances. When there was no such break, the distances were tabulated as questionable values. It seems likely that the slopes of radii vs. CN plots should be regular. The deviation from regularity is particularly noticeable for Na⁺, Ag⁺, K⁺, Cs⁺, Pb²⁺, Sr²⁺, and Ba²⁺. The lack of regularity is attributed either to an insufficient number of examples of a particular coordination and/or to the difficulty in determining the CN.

Several other groups of ions exhibit irregular coordination which frequently does not result in consistent interatomic distances. Also, the radii derived from these distances do not fall on the regular curves of radii vs. CN. These are the ions with the following electron configurations:

> d^4 -Mn³⁺HS, Cr²⁺HS d^8 -Ag³⁺, Au³⁺, Pd²⁺, Pt²⁺ d^9 -Cu²⁺, Ag²⁺ d^{10} -Ag⁺, Au⁺, Hg²⁺, Cu⁺ s^2 -As³⁺, Sb³⁺, Bi³⁺, Pb²⁺, Te⁴⁺

A detailed discussion of the stereochemistry of d^4 , d^8 , d^9 , and d^{10} ions is given by Orgel (1960). Ions with d^8 configuration frequently occur in square planar coordination. This coordination is usually clearly recognized, e.g. PdO, AuF₃. The ions with d^4 and d^9 configurations have the familiar Jahn-Teller distortion exemplified by Cu²⁺. These ions are characterized by four ligands at one distance and two ligands at a different distance, either closer or further away. When the difference is slight, the coordination is pseudo-octahedral; when the difference is large, square-planar or two-coordinated structures result. In the cupric oxides and fluorides the distances of the fifth and sixth neighbors range from just slightly larger than the other four to distances almost outside the bonding sphere. In the case of cupric compounds we have calculated the average distance of all six neighbors for octahedral coordination and the four near neighbors for square-planar coordination and have listed the radii in Table 1 as both IVSQ and VI.

Ions with the d^{10} configuration have been observed with linear two coordination. Orgel has related this behavior to the small separation in energy of the d^{10} and $d^{9}s$ states and predicts linear coordination of Cu⁺, Ag⁺, Au⁺, and Hg²⁺. Two-coordinated radii are tabulated for Cu⁺, Ag⁺, and Hg²⁺.

Another group represents the ions with a lone pair of electrons: As^{3+} , Sb^{3+} , Bi^{3+} , Pb^{2+} , and Te^{4+} . The oxides formed with these ions are considered to be highly covalent. For example in PbO the lone pair acts as a ligand and results in a coordination of four neighbors in a plane on one side of the Pb²⁺ ion and the lone pair on the other side at the apex of the pyramid. This configuration has been designated in Table 1(*a*) as IVPY.

Summary

The goal of this work has been to provide radii which will be useful to anyone interested in the structures of oxides and fluorides. The set of empirical radii derived here is primarily intended to be used in calculating expected interatomic distances in oxide and fluoride crystal structures. Although the absolute values of radii were considered, one cannot reliably assign specific numbers to specific ions. Certain effects such as anionanion repulsion, irregular coordination, and metalmetal bonding cause deviations from predicted interatomic distances wich have been largely ignored because these effects are difficult to include in a simple table of radii. These 'high-order' effects probably are of greater importance in structures with larger anions than in those containing oxide or fluoride ions and must be left to future work.

Note added in proof: – Prof. B. Reuter, who has just completed a refinement of the structure of the ordered spinel, MgV₂O₄, has suggested a value of $r(^{IV}Mg^{2+}) = 0.58$ Å. Because the original value of 0.49 Å came from a single structure refinement and because this new value is more consistent with Fig. 2(b), the new value is preferred.

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Stereochemistry and Crystal Structure of a Cobalt(III) Sulphito Complex: trans-Sulphiteisothiocyanate-bis(ethylenediamine)cobalt(III) Dihydrate

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The crystal structure of Co(en)₂SO₃NCS. 2H₂O has been determined from three-dimensional X-ray data and refined by the full-matrix least-squares method to a residual of 0.078. The compound crystallizes in the monoclinic system, space group $P2_1/c$ with four molecules in the unit cell of dimensions $a=9.13\pm0.02$, $b=6.59\pm0.02$, $c=22.88\pm0.06$ Å, $\beta=95.5\pm0.5^{\circ}$. The coordination around the metal atom is octahedral, the sulphito group being bonded through sulphur to the metal and in *trans* configuration. The Co-N(NCS) distance is not significantly different from the Co-N(en) distances. Some important dimensions are: Co-S, 2.203 ± 0.006 Å; mean Co-N(en), 1.962 ± 0.010 Å; mean S-O, 1.485 ± 0.012 Å; mean O-S-O, $110.3\pm0.7^{\circ}$; Co-N(NCS), 1.974 ± 0.018 Å.

Introduction

Recently, the infrared and visible absorption spectra of a series of sulphito-bis(ethylenediamine)cobalt(III) complexes were measured by Baldwin (1961). These spectra, together with measurements of conductivity in aqueous solution and a study of chemical reactions, were interpreted by Baldwin in terms of possible structures for these complexes. For Co(en)₂SO₃NCS.2H₂O it was postulated that the sulphito group is bonded through sulphur, the bond possibly involving $d\pi$ - $d\pi$ bonding, and that the compound had the *cis* configuration. Kinetic studies of sulphito complexes of Co¹¹¹ have also been made recently, indicating that the sulphito group has a marked *trans* labilizing effect (Tewari, Gaver, Wilcox & Wilmarth, 1967).

We undertook the structure determination of this compound in order to establish whether the postulated structure was correct, and, if so, to compare the dimensions of the sulphito group coordinated to Co^{III} , with the dimensions found for this group in PdSO₃(NH₃)₃ (Spinnler & Becka, 1967). Furthermore, we considered that a comparison between the different Co-N distances in this compound would be of interest in relation to the kinetics of reaction of sulphito complexes. In an earlier communication (Baggio & Becka, 1967) we described the main features of the structure as obtained from least-squares refinement with individual isotropic temperature factors. In this paper we give the details of the structure and structure determination with individual anisotropic temperature factors for Co, the atoms of SO₃, and the atoms of NCS.

Experimental

Preparation

 $Co(en)_2SO_3NCS.2H_2O$ was prepared by the method reported by Baldwin (1961), and crystals suitable for X-ray studies were obtained by recrystallization. The chemical and physical properties of the sample were in good agreement with those reported previously. Chemical analysis of a sample gave the following percentage composition for C, N and H: C, 17.37; N, 19.21; H, 5.75, the theoretical composition being: C, 16.59; N, 19.83; H, 5.66.

Crystal data

Name of substance: *trans*-sulphiteisothiocyanatebis(ethylenediamine)cobalt(III) dihydrate. Chemical formulae: structural, Co(en)₂SO₃NCS.2H₂O; alpha-

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