Dixenite, $Cu^{1+}Mn_{14}^{2+}Fe^{3+}(OH)_6(As^{3+}O_3)_5(Si^{4+}O_4)_2(As^{5+}O_4)$: metallic $[As_4^{3+}Cu^{1+}]$ clusters in an oxide matrix

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Abstract

The crystal structure of dixenite was analyzed using a crystal from the type and sole locality at Långban, Sweden. The end-member formula $Cu^{1+}Mn_{14}^{2+}Fe^{3+}(OH)_6(As^{3+}O_3)_5(Si^{4+}O_4)_2(As^{5+}O_4)$ is proposed. Dixenite is rhombohedral, a=8.233(4), c=37.499(1)Å, space group R3, Z=3. Twenty eight atoms occur in the asymmetric unit including two disordered Cu^{1+} cations. R=0.064 for 2507 independent reflections.

The structure is related to but distinct from that of hematolite, $(Mn^{2+},Mg,Al)_{15}$ $(OH)_{23}(AsO_3)(AsO_4)_2$. Three kinds of anionic radicals occur: $(As^{3+}O_3)$ trigonal pyramids; and $(Si^{4+}O_4)$ and $(As^{5+}O_4)$ tetrahedra. Three of the five nonequivalent layers along [001] are similar in hematolite and dixenite. One layer in dixenite, however, contains a disordered cluster, idealized as $(Cu^{1+}As_4^{3+})$ where a tetrahedron of As^{3+} ions surrounds a Cu^{1+} ion. All lone pair electrons from As^{3+} point into the central cavity which houses $Cu^{1+}(d^{10})$ and this cluster is believed to be stabilized by the 18-electron rule where $Cu^{1+}As_4^{3+}$ forms a closed argon core.

Introduction

Dixenite is a rare mineral, originally described by Flink (1920) from the mineralogically complex Fe-Mn oxide ore deposit in Långban, Sweden. The mineral was long problematical: Wickman (1951) proposed the formula (Mn,Fe,Cu,As³+)₂₀(Si,As⁵+)₃ (O,OH)₃₂, Wuensch (1960) presented a relationship to the complex arsenosilicate mcgovernite, and Moore and Araki (1978) proposed Mn¹¹+ Mn⁴+ (OH)₈(AsO₃)₆(SiO₄)₂ and a model for the structure derived from hematolite, (Mn²+,Mg,Al)₁₅(OH)₂₃ (AsO₃) (AsO₄)₂ to which it shares similarities in cell parameters and space group.

We studied dixenite's structure in hopes of gathering more clues about the structure of mcgovernite, and discovered several unusual features, including incorrectness of the proposed structure of Moore and Araki (1978), the presence of [As₄³⁺Cu¹⁺] metal clusters, the occurrence of As₅³⁺O₃ trigonal pyramids, and solid solution between As₄₊ and Si⁴⁺ in tetrahedral oxygen coordination.

Experimental details

On the basis of a relationship to hematolite, kraisslite and mcgovernite (Moore and Ito, 1978) we

suspected that platy deep red-brown crystals of dixenite from the only recorded locality at Långban, Sweden may in fact consist of more than one structure or polytype. Our dixenite sample selected for this structure study was NMNH No. C-6440. We also examined Nos. B-20579, 94920, 94935 and R-5755 (all in the U.S. National Museum of Natural History) by X-ray study and found all of them to be identical. We thank Mr. John S. White, Jr. for permission to select fragments of these specimens. The crystal selected was a deep red plate measuring 0.18 mm $||a_1 \times 0.25 \text{ mm}||a_2 \times 0.06$ mm ||c. With $\mu = 132.1 \text{ cm}^{-1} \text{ (Mo}K\alpha$), seven divisions by the Gaussian integral method (Burnham, 1966) led to significant absorption corrections, ranging from 0.148 for low angle (00*l*) reflections to 0.458.

Cell data were obtained from calibrated precession photographs (Mo $K\alpha$ radiation) and yielded a=8.233(4), c=37.499(1)Å, Laue symmetry $\overline{3}$. Intensities were collected on a PAILRED semi-automated diffractometer with the a_2 -axis || rotation and with graphite monochromatized Mo $K\alpha$ radiation. Background counting time on each side of the peak was 20 sec, scan speed 1° min⁻¹, scan width 4.0° to 4.8°. Angular coverage maximum was $\sin\theta/\lambda=0.80$, the

Table 1. Dixenite: atomic coordinate parameters[†]

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Atom	Population	æ	y	z
M(1)	1.0Mn ²⁺	0	0	0
M(2)	$0.90(2)Mn^{2+} + 0.10(2)Mg^{2+}$	1/3	2/3	0.00622(9
M(3)	1.0Fe ³⁺	0	0	0,25749(7
M(4)	1.0Mn ²⁺	0,0408(1)	0.2617(1)	0.06782(6
И(5)	1 OMn*	0.4158(2)	0.3359(2)	0.12987(6
M(6)	1.0Mn ²⁺	0.1089(1)	0.3976(1)	0.19230(6
M(7)	1.0Mn ²⁺	0.4226(1)	0.3154(1)	0.26133(6
Cu(1)	0.651(9)Cu1+ + 0.3490	1/3	2/3	0.31292(9
Cu(2)	0.192(9)Cu ¹⁺ + 0.808	2/3	1/3	0.0030(3)
T(1)	0.86(1)Si4+ + 0.14As5+	2/3	1/3	0.18792(9
T(2)	0.60(1)Si*+ + 0.40As5+	0	0	0,14620(7
T(3)	$0.24(1)Si^{4+} + 0.76As^{5+}$	1/3	2/3	0.11357(6
As(1)	1.0As 3+	2/3	1/3	0.06992(6
As(2)	1.0As 3*	1/3	2/3	0.25062(6
As (3)	1.0As ³⁺	0.08854(9)	0.37369(9)	0.31589(6
0(1)		0	0	0.1019(2)
0(2)		1/3	2/3	0.1584(3)
0(3)		2/3	1/3	0.2311(2)
0(4)		0.0899(7)	0.4444(7)	0.0212(2)
0(5)		0.4698(7)	0.1659(7)	0.0949(1)
0(6)		0.2837(7)	0.4545(7)	0.0988(2)
0(7)		0.1655(7)	0.2081(6)	0.1617(1)
0(8)		0.5194(7)	0.1277(7)	0.1707(1)
0(9)		0.3690(7)	0.5010(7)	0.2277(1)
0(10)		0.1468(7)	0.2272(7)	0.2905(1)
0(11)		0.4608(6)	0.1185(7)	0.2956(1)
OH(1·)		0.2140(7)	0.1791(7)	0.0375(2)
OH(2)		0.2352(7)	0.0814(7)	0.2271(1)

†Estimated standard errors refer to the last digit.

highest level k = 11. A total of 4989 reflections was covered including (hkl), (hkl), (h+l), h, l) and (h+k), h, l). Unobserved reflections with $I_0 < 2\sigma(I)$ accounted for 397 (8%) of the total reflections.

Equivalent reflection pairs, such as (hkl) and (h+k,h,l) were found to have equivalent intensities within error of observation after absorption correction and were therefore averaged. The Bijvoet pairs were preserved in the data set owing to pronounced acentricity, as determined with an N(z) test on general reflections (Howells $et\ al.$, 1950). The space group is therefore R3. A total of 2507 independent reflections were used in the ensuing study.

Solution and refinement of the structure

The dixenite model proposed by Moore and Araki (1978) and derived from the structure of hematolite, a basic manganese arsenite-arsenate which has similar cell parameters, was first tested and found to be incorrect. The structure was solved piecemeal, with stepwise approach by Fourier methods starting with atomic positions which satisfied the most prominent vectors of a Patterson synthesis. The problem of structure analysis proved to be exceedingly complicated, the results of which require an extensive revision of dixenite's proposed formula. Much like magnussonite (Moore and Araki, 1979a), a cluster of As³⁺ cations appeared, the core of which afforded two electron density maxima. Like magnussonite, the distances between

As3+ and these maxima were short (<2.7Å) and we anticipated "metal-metal" bonding. Unlike magnussonite, the As3+ defined a tetrahedral array, not an octahedral array. We assumed a similar mechanism operated in dixenite, i.e., the 18-electron rule and observed that sufficient Cu1+ was present to account for this residual density. Counting valence electrons there exist $2\times4(As^{3+}) + 10(d^{10} \text{ in } Cu^{1+})$ binding the cluster, defining a closed argon core. At this stage, we suspected that Cu¹⁺ reported in some magnussonite analyses may in fact occur in the center of the As₆³⁺ octahedron in that structure but no sensible electron "rule" could be extracted from this model. Solid solutions and partly occupied sites required application of mixed scattering curves $[xSi^{4+} + (1-x)As^{5+}]$ and ordered vacancies $[yCu^{1+} +$ $(1-v)\square$, where \square = vacancy]. Several cycles of atomic coordinate parameter, site population and anisotropic thermal vibration parameter refinement led to R = 0.064, and $R_w = 0.065$ for 2507 independent reflections where

$$R = \frac{||F_0| - |F_c||}{\sum |F_0|} \text{ and } R_w = \left[\frac{\sum_{w} (|F_0| - |F_c|)^2}{\sum_{w} |F_0|^2} \right]^{1/2}$$

with $w = \sigma^{-2}$ of F_0 . Refinement minimized $w(F_0 - F_c)^2$. Programs used in this study have been listed

Table 2. Dixenite: anisotropic thermal vibration parameters[†]

_	Atom	β11	B22	Saa	β12	β13	β _{2 3}	_
	М(1)	46(2)	46	12(1)	23	0	0	
	M(2)	48 (4)	48	41(2)	24	0	0	
	M(3)	28(2)	28	14(1)	14	0	0	
	М(4) М(5)	41(2) 45(2)	51(2) 61(2)	16(1) 18(1)	24(1) 29(1)	-0(0) -1(0)	-2(0) -4(0)	
	И(6)	42(2)	42(2)	18(1)	22(1)	-2(0)	-1(0)	
	M(7)	35(2)	35(2)	15(1)	19(1)	-1(0)	0(0)	
	m(/)	33(2)	33(2)	13(1)	19(1)	-1(0)	0(0)	
	Cu(1)	29(4)	29	19(2)	15	0	0	
	Cu(2)	38(14)	38	30(7)	19	0	0	
				(.,				
	T(1)	32(5)	32	9(2)	16	0	0	
	T(2)	30(3)	30	10(1)	15	0	0	
	T(3)	29(2)	29	8(1)	15	0	0	
	As(1)	38(2)	38	9(1)	19	0	0	
	As (2)	37(2)	37	10(1)	18	0	0	
	As(3)	35(1)	42(1)	12(1)	23(1)	-0(0)	-0(0)	
	N3(3)	33(1)	72(1)	12(1)	23(1)	-0(0)	-0(0)	
	0(1)	64(14)	64	14(5)	32	0	0	
	0(2)	60(14)	60	18(6)	30	0	0	
	0(3)	35(11)	35	10(5)	18	0	0	
	0(4)	72(9)	55(9)	14(3)	28(7)	-1(1)	1(1)	
	0(5)	34(7)	32(7)	19(3)	12(6)	-2(1)	-3(1)	
	0(6)	45(8)	39(7)	23(3)	30(7)	-4(1)	-2(1)	
	0(7)	40(7)	26(7)	18(3)	17(6)	0(1)	-2(1)	
	0(8)	37(7)	45(8)	11(3)	11(6)	2(1)	-1(1)	
	0(9)	50(8)	50(8)	15(3)	34(7)	-2(1)	0(1)	
	0(10)	49(8)	39(7)	19(3)	25(6)	0(1)	-2(1)	
	0(11)	27(7)	34(7)	14(3)	1(6)	1(1)	2(1)	
	OH(1)	44(8)	47(8)	17(3)	19(7)	-2(1)	-2(1)	
	OH(2)	50(8)	40(7)	14(3)	25(6)	-1(1)	-1(1)	
	(-)	(-)	(,)	- (-)	(-)	-(-)	- \-/	

[†]Coefficients in the expression exp- $[\beta_{11}h^2 + \beta_{22}k^2 + \beta_{33}l^2 + 2\beta_{12}hk + 2\beta_{13}hl + 2\beta_{23}kl]$. Estimated standard errors refer to the last digit. The coefficient β_{33} is $\times 10^5$, the others each $\times 10^5$.

Table 3. Dixenite: parameters for the ellipsoids of vibration[†]

Atom	i	μ_i	θ_{ia}	θ_{ib}	810	Beq(Ų)	Atom	i	μį	θ_{ia}	θ_{ib}	θ_{ic}	$\operatorname{Beq}(\mathring{\mathbb{A}}^2)$
М(1)	1 2 3	0.093(4) 0.109(4) 0.109(4)		90 determined determined		0.85(3)	As (3)	1 2 3	0.089(1) 0.092(1) 0.105(1)	10(90) 84(20) 98(4)	126(70) 105(14) 140(8)	82(33) 165(8) 77(4)	0.72(1)
М(2)	1 2 3	0.112(6) 0.112(6) 0.171(4)		determined determined 90		1.43(7)	0(1)	1 2 3	0.101(19) 0.128(19) 0.128(19)		90 determine determine		1.13(18)
М(3)	1 2 3	0.086(5) 0.086(5) 0.101(4)		determined determined 90		0.66(3)	0(2)	1 2 3	0.114(19) 0.124(19) 0.124(19)		90 determine determine		1.15(17)
M(4)	1 2 3	0.092(2) 0.102(2) 0.125(2)	120(12) 146(14) 76(4)	52(8) 90(7) 142(5)	41(11) 106(7) 54(4)	0.91(1)	0(3)	1 2 3	0.085(20) 0.095(20) 0.095(20)		90 determine determine		0.67(14)
И(5)	1 2 3	0.097(2) 0.104(2) 0.139(2)	103(14) 166(90) 85(3)	53(10) 72(5) 137(3)	38(8) 97(14) 53(2)	1.04(2)	0(4)	1 2 3	0.097(11) 0.120(9) 0.143(8)	81(10) 93(17) 170(90)	104(19) 146(35) 59(40)	14(14) 100(19) 80(18)	1.16(7)
м(6)	1 2 3	0.097(2) 0.100(2) 0.120(2)	55(30) 41(90) 108(4)	77(24) 160(90) 106(5)	56(7) 96(29) 35(5)	0.89(2)	0(5)	1 2 3	0.077(12) 0.105(9) 0.123(9)	71(17) 19(90) 89(23)	55(19) 136(90) 114(22)	64(10) 99(42) 28(12)	0.84(7)
M(7)	1 2 3	0.087(2) 0.095(2) 0.107(2)	33(31) 102(10) 121(6)	128(21) 138(27) 75(8)	60(14) 97(7) 31(9)	0.74(1)	0(6)	1 2 3	0.076(12) 0.099(10) 0.141(9)	31(90) 107(18) 115(7)	144(78) 125(27) 95(8)	75(26) 119(10) 33(11)	0.93(7)
Cu(1)	1 2 3	0.087(8) 0.087(8) 0.117(6)		determined determined 90		0.76(7)	0(7)	1 2 3	0.073(12) 0.101(9) 0.118(9)	108(22) 26(44) 73(26)	22(63) 97(20) 111(13)	71(17) 101(27) 23(22)	0.78(6)
Cu(2)	1 2 3	0.101(24) 0.101(24) 0.146(18)		determined determined 90		1.10(24)	0(8)	1 2 3	0.074(13) 0.097(10) 0.133(9)	123(11) 112(19) 42(36)	81(14) 122(15) 146(45)	35(23) 113(17) 66(20)	0.86(7)
T(1)	1 2 3	0.081(7) 0.091(9) 0.091(9)		90 determined determined		0.61(7)	0(9)	1 2 3	0.077(12) 0.111(10) 0.122(9)	41(33) 75(28) 127(21)	132(28) 133(22) 107(28)	56(27) 137(35) 67(26)	0.88(7)
T(2)	1 2 3	0.083(5) 0.087(6) 0.087(6)		90 determined determined		0.58(5)	0(10)	1 2 3	0.083(11) 0.112(9) 0.125(9)	115(21) 30(48) 75(32)	30(40) 90(21) 120(15)	60(20) 91(29) 30(28)	0.93(7)
T(3)	2 3	0.077(4) 0.086(5) 0.086(5)		90 determined determined		0.55(4)	0(11)	1 2 3	0.068(13) 0.102(10) 0.127(9)	122(17) 113(19) 139(69)	115(11) 83(18) 26(90)	72(11) 156(15) 75(34)	0.82(6)
As(1)	1 2 3	0.079(3) 0.098(3) 0.098(3)		90 determined determined		0.68(2)	OH(1)	1 2 3	0.091(11) 0.114(9) 0.125(9)	73(19) 20(90) 79(41)	62(18) 122(42) 135(49)	49(17) 110(47) 47(22)	0.97(7)
As(2)	1 2 3	0.083(3) 0.097(3) 0.097(3)		90 determined determined		0.67(2)	OH(2)	1 2 3	0.090(11) 0.103(10) 0.114(9)	107(30) 121(45) 143(60)	41 (52) 49 (47) 95 (24)	50(45) 137(49) 77(26)	0.84(7)

 $[\]dot{t}_i = ith$ principal axis, $\mu_i = rms$ amplitude, $\theta_{i\mathcal{Q}_i}$, $\theta_{i\mathcal{D}_i}$, $\theta_{i\mathcal{C}_i} = angles$ (deg.) between the *i*th principal axis and the cell axes α_1 , α_2 , and α_i . The equivalent isotropic thermal vibration parameter, Beq, is also listed. Estimated standard errors in parentheses refer to the last digit.

earlier (Moore and Araki, 1976). Scattering curves for Mn^{2+} , Cu^{2+} , Mg^{2+} , As^{0+} , Si^{4+} and O^{1-} were obtained from Cromer and Mann (1968) and anomalous dispersion corrections, f'', for all metals from Cromer and Liberman (1970).

Reasonable errors in bond distances (±0.005Å for metal-oxygen distances), sensible equivalent isotropic thermal parameters (<1.4Å), good agreement with observed specific gravity and chemical plausibility support our findings on this unusual structure and demonstrate yet again that crystal structure analysis may be required to establish a meaningful chemical formula.

Atomic coordinate parameters are given in Table 1, anisotropic thermal vibration parameters in Table 2, thermal vibration ellipsoids and equivalent iso-

tropic thermal parameters in Table 3, structure factor tables in Table 4, bond distances and angles in Table 5, select chemical analyses in Table 6 and bond length-bond strength relations in Table 7.

Discussion of the structure

Table 6 presents cell contents based on the structure analysis and on the Johansson analysis in Wickman (1951). The agreement is excellent and demonstrates the importance of structure study in ascribing formal charges. From the structure study,

¹ To obtain a copy of Table 4, order Document AM-81-181 from the Business Office, Mineralogical Society of America, 2000 Florida Avenue, N.W., Washington, D.C. 20009. Please remit \$1.00 in advance for the microfiche.

Table 5. Dixenite: bond distances and angles[†]

C	u(1)		As	3(1)		
3 Cu(1) (3) -As(3) (9) 1 Cu(1) (3) -As(2) (3)	2.240(1) Å 2.336(3)		3 As(1)-0(5)	1.779(5)		
average	2.336(3)		3 0(5)-0(5) (1)	2.619(8)**	94.8(2)	
		Angle (°)				
3 As(2) ⁽³⁾ -As(3) ⁽³⁾	3,316(1)	92.85(7)	As	s(2)		
$3 \text{ As}(3)^{(3)} - \text{As}(3)^{(4)}$	3.875(1)	119.75(1)	3 As(2)-O(9)	1.755(5)		
average	3.595	106.3		1,755(5)		
			3 0(9)-0(9) (1)	2,650(8)**	98,0(2)	
C	lu(2)					
3 Cu(2)-As(3) (3)	2.365(4)		As	s(3)		
1 Cu(2)-As(1)	2.510(10)		As(3)-O(4) ⁽⁸⁾	1.721(5)		
average	2.401		-0(11) ⁽¹⁾	1.762(5)		
(1)(1)			-0(10)	1.779(5)		
3 As(3) ⁽³⁾ -As(3) ⁽⁴⁾ 3 As(1)-As(3) ⁽³⁾	3.875(1)	110.0(2)	average	1.754		
3 As(1)-As(3)	3.967 3.921	108.9(3) 109.5				
average	3.741	109.3	0(4) (8) ~0(10)	2 647/71++	09 1(7)	
			0(4) ~ ~ 0(10) (1) (1)	2.643(7)** 2.785(7)	98.1(3) 103.8(2)	
3	1(1)		0(4) (8) -0(11) (1)	2.765(7)	105.6(2)	
3 M(1)-OH(1)	2.157(5) Å		average	2.740	102.8	
3 M(1)-O(11) (3)	2.137(5) A 2.235(5)					
average	2.196					
				T(1)		
3 OH(1)-OH(1) (1)	2.832(8)*	82.0(2)	1 T(1)-0(3)	1.620(9)		
3 0(11) (3) -0(11) (4)	2.998(8)*	84.2(2)	3 T(1)-0(8)	1.641(5)		
3 OH(1)-O(11) (4)	3,151(7)	91.7(2)	average	1.636		
3 OH(1)-O(11) (5)	3.432(7)	102.7(2)				
average	3.103	90.1	3 0(8)-0(8) (1)	2.614(9)	105.5(2)	
-			3 0(3)-0(8)	2.722(9)	113.2(2)	
1	M(2)		average	2.668	109.4	
3 M(2)-0(4) 3 M(2)-0(10) ⁽³⁾	2.001(5) 2.464(6)			T(2)		
average	2.232		1 7/2) 0/1)	1 662(0)		
2,700			1 T(2)-0(1) 3 T(2)-0(7)	1.662(9)		
6.3			average	1.667		
3 0(4)-0(10) (4)	2.643(7)**	71.7(2)				
3 0(10) ⁽³⁾ -0(10) ⁽⁴⁾ 3 0(4)-0(4) ⁽¹⁾	2.842(9)*	70.4(2)	(1)			
3 0(4)-0(4) (5) 3 0(4)-0(10) (5)	3.327(9) 3.403(8)	112.4(2)	3 0(7)-0(7) (1)	2.712(8)	108.6(2)	
3 0(4)-0(10) average	3.403(8)	98.8(2) 88.3	3 0(1)-0(7)	2.734(9)	110.3(2)	
2102460	J. 054	00.5	average	2.723	109.4	
				T(3)		
1	M(3)		3 T(3)-O(6)	1.675(5)		
3 M(3)-OH(2)	2.048(5)		1 T(3)-0(2)	1.681(10)		
3 M(3)-0(10)	2,056(5)		average	1.676		
average	2.052					
			3 0(2)-0(6)	2.738(9)	109.3(2)	
3 0(10)-0(10) (1)	2.842(9)*	87.4(2)	3 0(6)-0(6) (1)	2.738(8)	109.6(2)	
3 O(10) -OH(2) (2)	2.901(7)	90.0(2)	average	2.738	109.4	
3 OH(2)-0(10)	2.915(7)*	90.5(2)				
3 OH(2)-OH(2) ⁽¹⁾	2.946(9)	92.0(2)				
	2,901	90.0				

†Estimated standard errors in parentheses refer to the last digit. The equivalent positions (referred to Table 1) are designated as superscripts and are (1) = -y, x-y, z; (2) = y-x, -x, z; (3) = (1/3 2/3 2/3) + (x, y, z); (4) = (1/3 2/3 2/3) + (-y, x-y, z); (5) = (1/3 2/3 2/3) + (y-x, -x, z).

Table 5 (continued)

		-				
	М(4)			M(6)		
M(4)-0(5) (1)	2.152(5)		$M(6)-0(8)^{(1)}$	2.086(5)		
-OH(1) ⁽¹⁾	2.160(5)		-0(7)	2,166(5)		
-0(6)	2,165(5)		-OH(2) ⁽¹⁾	2.244(5)		
-OH(1)	2.178(5)		-0(9)(2)	2.262(5)		
-0(4)	2,207(5)		-0(9)	2.289(5)		
-0(1)	2.377(5)		-0(2)	2.415(5)		
average	2,206		average	2.244		
OH(1)-OH(1)(1)	2.832(8)*	81.5(3)	0(9)-0(9)(2)	2.650(8)**	71.2(3)	
O(4)-OH(1)	2.898(7)	82,7(2)	0(8) (1) -0(9) (2)	2.967(7)		
0(1)-OH(1)(1)	2.915(9)*	79.8(2)	0(2)-0(9)		86.0(2)	
O(1)-OH(1)	2.915(9)*	79.4(2)	0(2)-0(9) (2)	3.015(9)*	79.7(2)	
O(4)-OH(1)(1)	3.026(8)	87.7(2)	0(8) ⁽¹⁾ -0H(2) ⁽¹⁾	3.015(9)*	80.2(2)	
0(5) (1) -OH(1) (1)	3.048(7)	90.0(2)	O(7) -OH(2) (1)	3.031(7)	88.8(2)	
0(6)-OH(1)	3.072(7)	90.0(2)		3.071(7)	88.3(2)	
0(5)(1)-0(6)	3.130(7)	98.2(2)	0(7)-0(9)	3.272(7)	94.5(2)	
0(1)-0(6)			0(2)-0(7) 0(2)-0(8) ⁽¹⁾	3.307(5)	92.3(2)	
0(4)-0(6)	3.272(5) 3.298(7)	92.0(2)	0(2)-0(8) (1) 0(9)-0H(2) (1)	3.336(5)	95.4(2)	
0(4)-0(5)(1)		98.0(2)	0(9) -OH(2) (1) 0(9) (2) -OH(2) (1)	3.361(7)	95.7(2)	
0(1)-0(5)(1)	3.311(7)	98.9(2)	0(7)-0(8)(1)	3.405(7)	98,2(2)	
average	3.404(5) 3.093	97.3(2) 89.6	O(7)-O(8)	3.442(7) 3.156	108.1(2) 89.9	
	M(5)			M(7)		
M(5)-O(8) (2)	2.093(5)		M(7)-OH(2)	2,182(5)		
-0(5)	2.120(5)		-0(9)	2.188(5)		
-0(6)	2,133(5)		-0(11)	2.210(5)		
-0(7)	2.145(5)		-0(3)	2.245(5)		
			-0(11)(2)	2.284(5)		
-0(5)(2)	2.429(5)		-0(10)	2.285(5)		
			average	2.232		
-0(8)	2.730(5)			4.202		
average (inner four)	2.123					
average (inner five)	2.184		O(10)-OH(2)	2.915(7)*	81.4(2)	
average (six)	2,275		0(3)-0(11)	2.974(8)*	83.8(2)	
			0(3)-0(11)(2)	2,974(8)*	82.1(2)	
			0(11)-0(11)(2)	2,998(8)*	83.7(3)	
Inner four anions			O(9)-OH(2)	3.053(7)	88.6(2)	
0(6)-0(7)	2.939(7)	04.045	O(3)-OH(2)	3.091(5)	88.6(2)	
0(7)-0(8)(2)	3,120(7)	86.8(2) 98.1(2)	O(11)-OH(2)	3.094(7)	89.6(2)	
0(5)-0(6)	3.410(7)	106.6(2)	0(10)-0(11)	3.132(7)	88,3(2)	
0(5)-0(8)(2)	3.623(7)	118.6(2)	0(9)-0(10)	3.138(7)	89.1(2)	
0(5)-0(7)	3.677(7)	119.2(2)	0(3)-0(9)	3.359(5)	98.5(2)	
0(6)-0(8)(2)	3.724(7)	123.6(2)	0(9)-0(11)(2)	3.388(7)	98.5(2)	
average	3.416	108.8	0(10)-0(11) (2)	3.673(7)	107.0(2)	
-			average	3.149	89.9	

we accepted Cu^{1+} , Mn^{2+} , As^{3+} and As^{5+} as formal charges, then recalculated Johansson's analysis to accommodate these differences which resulted in a negligible amount of Mn^{3+} . From the structure study, we obtain $Mn^{2+}_{42.13}Mg^{2+}_{0.30}Fe^{3+}_{2.57}As^{3+}_{15.00}$ $Cu^{1+}_{2.53}Si^{4+}_{5.10}As^{5+}_{3.90}$ (OH)₁₈O₈₁ in the cell. This gives $\rho(\text{calc}) = 4.375 \text{ g cm}^{-3}$ in excellent agreement with the specific gravity of 4.36 reported in Wickman (1951). From this exceedingly complex formula a

unit formula is proposed for an ideal "end-member" composition (vide supra):

 $Cu^{1+}Mn_{14}^{2+}Fe^{3+}(OH)_6$

$$(As^{3+}O_3)_5(Si^{4+}O_4)_2(As^{5+}O_4), Z = 3.$$

This formula disguises the peculiar aspects of dixenite's crystal chemistry: first Si⁴⁺ and As⁵⁺ mix over their sites, and second the Johansson analysis

Table 6. Dixenite: chemical analysis and its interpretation

_							
_		1	2	3		4	5
	P2O5	0,02			P5+	-	-
	As205	-	7.74	5.96	As ^{s+}	3.90	3,90
	SiO ₂	5.31	5.30	6.23	Si*+	5.10	5.13
	As ₂ O ₃	32.16	25.64	25.65	As3+	15.00	14.97
	Mn ₂ O ₃	8.05	-	8	Mm 3+	-	0.66
	Fe ₂ O ₃	3.75	3.55	4.14	Fe ³⁺	2.57	2.73
	MgO	0.32	0.21	-	Mg 2 + Mn 2 +	0.30	0.46
	MnO	43.35	51,63	51.51	Mn ²⁺	42.13	40.73
	CaO	0.39	100	-	Ca ²⁺	-	0.41
	CuO	3.49	-	2	Cu ²⁺	-	. 70
	Cu ₂ O	_	3.13	3.71	Cu1+	2.53	2.55
	Na ₂ O	0.13	40	-			
	K ₂ O	0.14	2.1	Ĩ.			
	H ₂ O	2,80	2.80	2.80			
	Total	99,91	100.00	72.00 180.00	Σ Atoms Σ Charge	71.53 180.00	71.54 180.85
	Specific gravity Density (g cm ⁻³)	4.36	4.375				

¹ Johansson analysis in Wickman (1951).

suggests a slight deficit of cations which in Table 6 appears to result from less As⁵⁺ and more Si⁴⁺ and somewhat less Cu¹⁺ in his analysis. However, the table demonstrates very good agreement with the "end-member" formula.

The most interesting feature of the structure is a metallic cluster, ideally [As₄³⁺Cu¹⁺] where the Cu¹⁺ is coordinated by four As₃³⁺ at the vertices of a distorted tetrahedron, the lone-pair electrons pointing into the central Cu1+ cation. This feature was a surprise in the structure study but it is interesting to note that minerals coexisting with dixenite include native lead, Pb°; α-domeykite, α-Cu₃As and magnussonite, Mn₉²⁺Cl[As₆³⁺Mn¹⁺O₁₈]. Magnussonite's structure (Moore and Araki, 1979a) evidently consists of a metallic cluster [As₆³⁺Mn¹⁺] where the 18electron rule is also satisfied but the distribution of As³⁺ about Mn¹⁺ defines a distorted octahedron. However, in the more recent study on the related arsenite armangite (Moore and Araki, 1979b) we found a similar distribution of As₆³⁺ but no central metal, thus creating concern over the magnussonite study. However, the excellent convergence of the dixenite refinement, the role of Cu1+ and the appearance of a tetrahedral metallic cluster strongly implies that these "bits of metal" in an oxide matrix are real and that dixenite and coexisting magnussonite are two examples in natural systems which represent a transition between the ionic oxides and the sulfides, sulfosalts and alloy-like phases which contain strong metallic bonds. The end-member formula emphasizes that Cu¹⁺, and to a lesser extent Fe³⁺, are essential to the structure.

Dixenite is thus, like its relative mcgovernite, a basic arsenite-silicate-arsenate. Its name derived from the occurrence of two strangers—arsenite and silicate radicals—as originally proposed by Flink (1920). But it might better have been called "trixenite" owing to the presence of *three* radicals!

Although the structure cells of dixenite and hematolite are very similar, their contents and layer arrangements are quite different. Moore and Araki (1978) showed that hematolite is based on closest packing of oxygens and presented the five nonequivalent layers as Figures 2a-e and that the stacking sequence is · hhhch · . Using the same layer notation in the Figure la—e series in this paper it is seen that the m=0 layer in dixenite is quite unlike any layer in hematolite and includes the disordered [As₄³⁺Cu¹⁺] clusters. Even the layer itself is not close-packed as shown by the nonparallel orientation of the M(1) and M(2) octahedra. The m=1 layer is the same type as m=2 in hematolite. The dixenite m=2 layer has no correspondence with hematolite, consisting of T(2) and T(3) tetrahedra and very distorted M(5) polyhedra. The dixenite m=3 layer is similar to the m=1 layer in hematolite but with T(1)O₄ tetrahedra in place of hematolite's $As^{3+}O_3$ trigonal pyramids. The dixenite m=4 layer is similar to hematolite's m=1 layer but with As³⁺O₃ trigonal pyramids instead of (AsO₄) tetrahedra. This layer is interesting in that it is the same type of octahedral layer as found in Figure 1a (Horiuchi et al., 1979) while the m=3 layer is the same type as their Figure 1b as found in $2Mg_2SiO_4 \cdot 3Mg(OH)_2$. Welinite, Mn⁴⁺Mn₃²⁺SiO₇, the simplest of these structures, has an octahedral sheet like the m=4layer in dixenite. What is interesting about this layer is the occurrence of small octahedrally coordinated cations on the special 3-fold axial position: in welinite it is populated by Mn⁴⁺, in hematolite by mixed Al3+ and Fe3+, in dixenite principally by Fe³⁺(M(3)). For this reason, Fe³⁺ (or possibly Al³⁺) appears to be an important component in the structure.

Bond distances and their deviations

Table 5 lists individual bond distances and angles for the individual coordination polyhedra in dixenite's structure. The individual distances were arranged according to increasing values and shared polyhedral edges are starred. These shared edge

²Calculated weight percent from structure study and from column 4.

³For proposed end-member composition Cu¹⁺Mn₁₄Fe³⁺(AsO₃)₅(SiO₄)₂(AsO₄)(OH)₆.

 $^{^4\}mathrm{Cell}$ contents of cations computed from structure analysis. Total Fe $^{1+}$ was computed to exactly balance anion charge = 180 electrons.

 $^{^5}$ Cell contents of cations computed from Johansson analysis and converting Cu²++ Cu¹+:As³+ → As⁵+; As³+ → As⁵+: Mn³+ → Mn²+ in that order, the Cu¹+ and As⁵+ totals dictated from the structure study.

Table 7. Dixenite: electrostatic valence balance of cations and anions[†]

Coordinating	Cations
--------------	---------

	M(1)	M(2)	M(3)	M(4)	M(5)	M(6)	M(7)	T(1)	T(2)	T(3)	As(1)	As(2)	As(3)	Δρο
Bond Strength	2/6	2/6	3/6	2/6	2/4	2/6	2/6	4.25/4	4.5/4	4.75/4	3/3	3/3	3/3	1000
Anions											_		-	
0(1)				3			-		1	200000	0.000	22222		+0.125
				+0.17					-0.01					+0,125
				(x3)	+			****				****		
0(2)						3				1				+0.187
						+0.17				+0.01				+0.107
						(x3)								
0(3)	****					55516	3	1	*****					0.000
							+0.01	-0.02		*****	57075			+0.062
							$\frac{+0.01}{(x3)}$	-0.02						
0(4)		1		554/5										
~ (+)		-0.23		+0.00		20000	-		*****				1	-0.333
		-0.725		+0.00									-0.03	
		100000	.57555						*****					
0(5)				1	1		*****	****		*****	1			-0.167
		****	****	-0.05	-0.00		20000				+0.00	****		
			****								*****			
0(6)				1	1					1				-0.167
				-0.04	+0.01		****			-0.00				-0.107
											****	****		
0(7)		*****	*****		1	1			1			200000		-0.042
	*****		*****		+0.02	-0.08			+0.00					-0.042
			*****							*********				
0(8)			*****		1	1		1						
- (-)			*****		-0.03	~0.16		77				75777		-0.105
					-0.03			+0.01						
0(9)						0								
0(3)						2	1	50500		7.7777	*****	1	****	+0.000
						+0.02	-0.04					+0.00		
				****	****	+0.05								
0(10)		1	1		27222	****	1	*****	****	****	****	*****	1	+0.167
	****	+0.23	+0.00	****			+0.05	Seren.				meaner:	+0.03	
						****					****			
0(11)	1						2	****					1	+0.000
	+0.04					*****	-0.02						+0.01	.01000
			****				+0.05			*****		70000	777.02	
OH(1)	1			2		*****	07300000							. 0. 000
	-0.04			-0.05		*****		*****				*****		+0.000
				-0.03		22224		*****			*****	*****	*****	
OH(2)														
ULITED	PERSONAL PROPERTY.		-0.00		~~~~	1	1							+0.167
					*****	+0.00	-0.05							
		50000								****				

[†]A bond length deviation refers to the polyhedral average subtracted from the individual bond distance. The entries begin with the number of cations coordinating to an anion followed by the distance deviations. The Δp_0 = deviation of electrostatic bond strength sum from neutrality (p_0 = 2.00 e.s.u. for 0^{2-} , 1.00 for OH⁻). Bond length deviations which conform to Δp_0 are underlined.

distances usually occur toward the top of their appropriate list. It is worth noting that only the $[As^{3+}O_3]$ trigonal pyramids share edges with the larger polyhedra: all three individual trigonal pyramids involve some edge-sharing in contradistinction with the three $[T^{\geq 4}+O_4]$ polyhedra where no edges are shared. The same phenomenon occurs in hematolite (Moore and Araki, 1978); the $As^{3+}-O$ 1.75–1.78Å polyhedral averages are close to the 1.79Å average in hematolite. The T-O averages steadily

increase, from T(1)-O to T(3)-O, in accordance with increasing As⁵⁺ solution at these sites. The O-As³⁺-O' angles range from 95° to 103°, compared with 94° in hematolite and 96° in synadelphite (Moore, 1970). Two polyhedra presented problems but these are easily resolved if extensive ordering is assumed in the dixenite structure. The M(3)-O 2.05Å average, discussed earlier, is approximately an Fe³⁺-O distance. The M(5)-O averages show a range of distances: four distances below 2.15Å

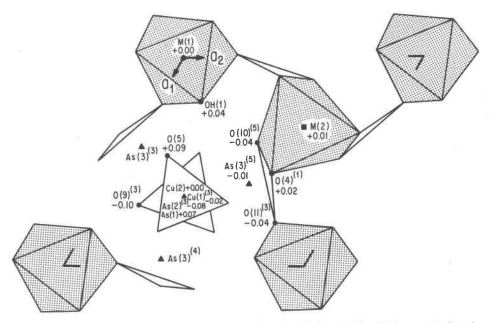


Fig. 1. Polyhedral diagrams of the five non-equivalent layers in dixenite. The layer is identified as z = 2m/30 where m is an integer. Heights are given as fractional coordinates.

Fig. 1a. The m=0 layer including the $As(1,2,3)O_3$ trigonal pyramids and the $M(2)O_6$ octahedron. This layer shows the region around the $Cu^{1+}As_3^{3+}$ cluster. Hematolite has no such layer.

(average 2.12Å), one distance at 2.43Å and a remaining distance at 2.73Å. Such multiple "coordination spheres" have been noted earlier for Mn²⁺ in an oxide environment. Arsenoclasite, Mn₅²⁺

(OH)₄(AsO₄)₂, has polyhedra involving tetrahedral (2.13Å average), trigonal bipyramidal (2.19Å average) and octahedral (2.22Å average) coordinations (Moore and Molin-Case, 1971). In the present

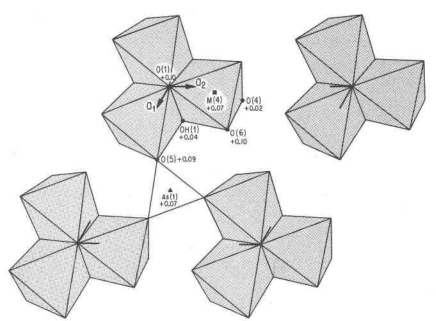


Fig. 1b. The m=1 layer showing the M(4)O₆ octahedra and the As(1)O₃ trigonal pyramid. This resembles the m=2 layer in hematolite.

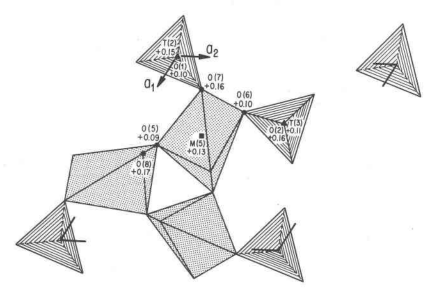


Fig. 1c. The m=2 layer showing the M(5)O₄ tetrahedron (here with additional O(5)⁽²⁾), the T(2)O₄ and T(3)O₄ tetrahedra. Hematolite has no such layer.

study, we have chosen tetrahedral coordination of oxygens about Mn(5) and used this for the valence balance calculations in Table 7.

The [Cu¹⁺As₄³⁺] tetrahedral cluster is very interesting as discussed earlier. This cluster (Figs. 1a, 2) involving a central Cu⁺¹ cation has no counterpart among the sulfosalt or arsenide structures. Besides, in dixenite the Cu atomic positions are not fully occupied but are split into two non-equivalent sites.

Mean distances (Table 5) are Cu(1)-As(2,3) 2.26 and Cu(2)-As(1,3) 2.40Å. Perhaps the lautite structure contains a good model of such a cluster since Cu¹⁺ is tetrahedrally coordinated by sulfur and arsenic to form a [CuAsS₃] cluster. Craig and Stephenson (1965) report a Cu-As 2.42Å distance which is close to the average for the Cu(2)As₄³⁺ cluster in dixenite. The short Cu(1)As₄³⁺ average distance is not so easily explained. Perhaps it is a

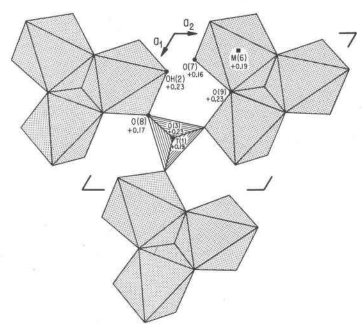


Fig. 1d. The m=3 layer showing the $M(6)O_6$ octahedra and the $T(1)O_4$ tetrahedron. This resembles the m=2 layer in hematolite.

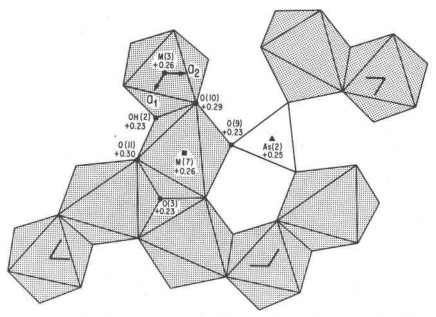


Fig. 1e. The m=4 layer showing the M(3)O₆ and M(7)O₆ octahedra, and the As(2)O₃ trigonal pyramid. This resembles the m=1 layer in hematolite.

consequence of the cluster disorder or even the possible presence of a different metal. This latter possibility is difficult to rationalize since there is no other site which Cu could occupy, either as Cu¹⁺ (owing to its large size) or Cu²⁺ (owing to pronounced Jahn-Teller distortion).

Calculated weight percentages for the structure analytical results and for the proposed end-member formula Cu¹⁺Mn₁₄²⁺Fe³⁺(OH)₆(AsO₃)₅(SiO₄)₂(AsO₄) show generally good agreement with Johansson's analysis of the mineral in Table 6, bearing in mind that some solid solution exists between Si⁴⁺ and

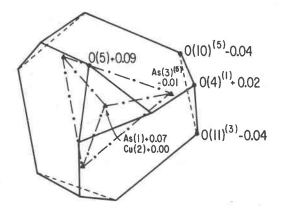


Fig. 2. The oxygen coordination polyhedron about the Cu¹⁺As₄³⁺ cluster in dixenite. This polyhedron is a distorted truncated tetrahedron.

Table 8. Dixenite: calculated and observed powder patterns[†]

(calc)	d(calc)	hkl	I(obs)	d(obs)
7	12.500	003	30	12.5
5	6.996	101	16	6.99
9	6.250	006	35	6.22
21	4.112	110	90	4.10
29	3.906	113	50	3.90
44	3.435	116	40	3.42
16	3.318	10.10	45	3.31
36	2.965	027	80	2.96
100	2.927	119	100	2.92
24	2,835	208	50	2.83
15	2.685	211	40	2,68
11	2.664	122	30	2.66
5	2.587	214	20	2.58
6	2.582	02.10		
7	2.533	125	16	2.53
7	2.507	01.14		
6	2.500	00.15		
25	2.488	11.12	40b	2.49
22	2.405	217	55	2,40
21	2.374	300	80	2.37
16	2.334	128	45	2.33
5	1.972	131	25	1.967
6	1.968	21.13		
5	1.820	318	12b	1.819
14	1.768	21.16	30	1.764
9	1.721	30.15	20	1.719
12	1.706	12.17	25	1.703
5	1.638	11.21	12	1.635
6	1.563	327	16	1.560
19	1.554	410	55	1.551
6 5	1.543	238	25Ъ	1.541

[†]The calculated data are from the refined structure, based on CuK_Ω radiation. These results are compared with ASTM Powder File No. 19-426. Only calculations with I(calc) \geq 5 are listed. Agreement is good, excepting preferred orientation effects in the experimental study.

As⁵⁺. Performing the appropriate valence conversions suggested by the structure study brings Johansson's analysis into very good agreement indeed (columns 4 and 5). It is gratifying to see that very little Mn³⁺ evidently occurs in the structure, conforming to the rather reduced state of the species.

When a complex structure is well-refined it is desirable to calculate a powder pattern and compare it with existing powder data as given in Table 8. One advantage is the correct assignment of the Miller indices and its advantage over experimentally determined powder patterns which usually exhibit some preferred orientation and absorption effects.

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