The existence of the radiating system 5^6 suggested the possibility of a radiating (3,4)-connected net formed of hexagons which would fill the gap between the plane 5-gon net and the 7-gon, 8-gon, and 9-gon periodic nets. There is in fact a net of this kind (Fig. 15) which consists of an infinite set of concentric tetrahedra linked alternately by lines joining vertices and midpoints of edges. Apart from the central group of 10 3-connected points each successive tetrahedral shell contains 6 3-connected and 4 4-connected points so that for the infinite system $c_3: c_4=3:2$. This net, $R_{3,4}$, completes the family of fundamental nets of Table 2.

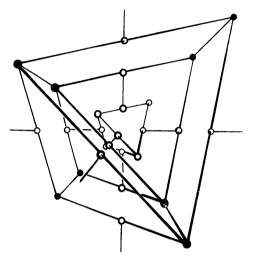


Fig. 15. The radiating net $\{6,\frac{3}{4}\}$.

The relation of the radiating systems and the surface tessellations of part VII to other 4-connected systems is shown below.

Regular and uniform 4-connected systems

y=2 These include the following:

n=3 octahedron, $\{3,4\}$

n=4 plane net, $\{4,4\}$

 $n \ge 5$ (a) infinite plane radiating nets,

(b) the infinite three-dimensional surface tessellations of part VII, {5,4}, {6,4}, and {7,4}; others have $2\langle y\rangle 3$. (The y values for these nets were not given, and {7,4} was not illustrated; it is the reciprocal of Fig. 22(c) of part VII).

v=3

n=3 finite radiating 36 [Fig. 13, (a) and (b)]

n=4 finite radiating 46 [Fig. 13(c)]

n=5 infinite radiating 56 (Fig. 14)

v = 6

n=6 diamond net, 6^6 or (6.4).

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A Ternary Alloy with PbCl₂-type Structure: TiNiSi(E)*

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Beck and coworkers have found E phases in several ternary systems of transition elements with either silicon or germanium at the composition 1:1:1. The crystal structure of TiNiSi(E) has been determined and refined by least squares with (limited) three-dimensional single-crystal data to a final R value of 0.086 (excluding 002 due to apparent extinction, and all non-observed reflexions). The lattice parameters for the primitive orthorhombic cell are:

$$a_0 = 6.1484 \pm 0.0012$$
, $b_0 = 7.0173 \pm 0.0014$, $c_0 = 3.6698 \pm 0.0007$ Å.

The E phase is isotypic with $PbCl_2(C23)$, space group Pnam. All near-neighbor distances are within 0.06 Å of the following average values: Ti-Ti 3.18, Ti-Ni 2.83, Ti-Si 2.61, Ni-Ni 2.67, Ni-Si 2.33 Å The numbers of near-neighbors are compared with those in Co_2Si , $\theta - Ni_2Si$ and U_3Si_2 .

Introduction

The E phase was first identified by Westbrook, DiCerbo & Peat (1958) in the titanium-nickel-silicon

* Sponsored by Army Research Office (Durham). Computations were done in large part at the M. I. T. Computation Center.

system at the composition TiNiSi. Subsequently Spiegel, Bardos & Beck (1963) concluded from the powder X-ray diffraction diagrams that twenty-one additional phases in other ternary systems of transition elements with either silicon or germanium are isomorphous with TiNiSi(E) and they indexed the powder patterns on large orthorhombic cells. They found

Table 1. X-ray powder photographic data for TiNiSi

	1.		•	٠.	A ruy	ponue	i pitot	одгирт	c uuii	u j	U	1.	11 (15)		
WVL#10BS	THETA OBS*	н *	ĸ	L	THETA CAL	DELTA THETA	1CAL***	WVL 1085	THETA OBS	н	ĸ	L	THE TA CAL	DEL TA THE TA	ICAL
3 M	9.648	1	1	0	9.596	0.052	1266	2 22.		56	1	1	52.069	0.170	70
		0	2	0	12.692 13.713		238 1	3 W(BR)	52.239	٠,	6	2	52.339 52.549	-0.100	296 156
		2	2	0	14.523		8 139			0	7	1	52.861		55 4
3 M	15.633	1	1	1	14.653 15.556	0.078	1092	3 VW	53.780	(4	ó	3	53.563 53.649	0.130	2
з vs	19.210	2 (2	1	0	15.889 19.094	0.116	66 2769			1 2	7	1	53.801 53.916	-0.022 -0.137	230 2
		(ı	2	1	19.197	0.013	4904			2	7	2	53.984		10
3 W 3 VS	19.486 20.252	2	2	0	19.475 20.186	0.011 0.065	835 4090	3 VW	54.244	{6 4	2	3	54.231 54.376	0.013 -0.132	196 83
3 MS	20.653	1	3	ō	20.647	0.006	1590	3 W	55.198	j2	6	2	55.174	0.025	151
3 VVW 3 VVW	21.902									\6 5	3	2	55.216 55.436	-0.018	69 23
3 5	23.100	10	3	1	23.006	0.095	2667	3 VW 3 VW	55.877 56.675	(5	5	0	55.919 56.460	-0.042 0.145	145 38
3 W	23.201	2	2	0	23.070 23.207	0.030 -0.006	1895 870	, ,,	20.077)4	2	3	56.598	0.007	66
3 M 3 W	24.230	1	3	1	24.233	-0.002 -0.033	1126 842			20	7	1	56.687 56.718	-0.087 -0.117	45 123
3 S	24.833	0	ó	2	24.843	-0.010	2415	3 W	57.170	0	ņ	4	57.167	0.002	289
3 M	25.787	3	?	0	25.824 26.067	-0.037	820 123			1		3	57.632 57.892		0 19
3 MW	26.339	3	1	ī	26.399	-0.060	611			4		2	58.599 58.623		11 53
3 VW	27.230	1	1	0	26.872 27.191	0.039	153 335	1 M	58.814	13	4	3	58.726	0.088	385
3 VW	27.669	2	3	1	27.668	0.001	164			13	7	0	58.798 58.862	0.016 -0.048	43 11
		3	2	2	28.302 28.922		49 68			5	5	1	59.188	-0.048	10
		2	0	2	29.293 29.368		1 33			6	9	2	59.422 60.080		37 0
3 VW(BR)	30.029	[3	3	ő	30.007	0.022	71	I MW	60.233	Jó	2	4	60.202	0.031	2
		{2 4	1	2	30.096 30.101	-0.067 -0.072	19			16	1	2	60.216	0.017	150 13
		1	4	1	30.194	0.0.2	6			6	4	9	60.517		9
		`2 4	4	0	30 • 394 30 • 892		49 68	1 VW	60.796	{2 5	ě	3	60.698 60.757	0.088	49 328
3 M	32 • 402	2	2	2	32.435	-0.033	313			2	2	4	61.181	-	0
		4	3	1	32.849 32.939		19			0	2	9	61.257 61.417		1 57
3 MS	33.255	14	2	0	33.199 33.217	0.057	420			3	7	1	61.730		0
		12	3	1 2	33.264	0.038	6 651			7	1	ō	62.102		19
3 VW	33.670	4	1	1	33.692	-0.022	269			1	7	2	62.186		33
3 W 3 MS	34.272 35.051	1	1	2	34.292 35.065	-0.020 -0.015	236 925			6	2	2	62.683		ō
3 M	36.017	3 (4	2	0	35.340 35.905	0.111	11 165	1 W	63.498	{5 6	1	3	63.499 63.566	-0.001 -0.068	327 2
,	10.017	{∘	5	1	36.021	-0.004	324	2 VW	63.875	(5	1	3	63.787	0.087	164
		(2	3	2	36 • 151 36 • 879	-0.135	44C	1 VW	64.691	{S	2	1	63.855 64.586	0.019 0.105	1 70
		1	5	1	36+963		ō			{7	2	0	64.689 65.280	0.002	61 0
3 MS	37.222	}2 {3	5	2	37 • 144 37 • 243	0.078	312 498			5	Ē	ć	65.354		0
2 46	37.925	0	4	2	37.441 37.980	-0.055	80 1188	1 VW	65.544	{1 2	?	2	65.573 65.691	-0.029 -0.057	164
3 MS 3 VW	38 • 378	3 1	4	2	38.371	0.008	205			1	9	1	65.714		17
		4	3	1	39 • 48 1 39 • 528		51 21	1 VW 1 VW	65.933 66.250	2 (5	Р 2	3	65.947 66.191	-0.013 0.069	86 50
3 VW	39.684	12	5	1	39.741	~0.057	112			(3	5	3	66.339	-0.080	102
		10 [1	1	3	39.769 40.683	-0.086 0.156	0 45			4	7	3	66.451 66.538		13 46
3 VW	40.839	{3	3	2	40.780 40.862	0.059	55 5			1	6	3	66.888		115
		2	4	2	41.123	-0.024	42			13	1	2	67.185 67.875	0.155	108 280
		0	6	0	41.234 41.560		0 50	1 M(BR)	6P.030	14	£ 2	2	68.115 68.132	-0.085 -0.102	249
		5	2	0	41.628		103			6	5	0	68.545	-0.102	101
		4	4	0	41.738 41.821		64			5	÷	?	68 • 847 68 • 879		69 1
3 M	42.094	15	1 6	0	42.085	0.009	700			2	3	4	69.388		145
3 M	42.797	12	0	3	42.141 42.707	0.090	218 201	1 MW	69.549	17	8	1	69.509 69.550	0.040 -0.001	113 309
3 M	43.350	l1 2	2	3	42.771 43.400	0.026 -0.051	37C 365	2 VW	69.902	j7	,	ē	69.896	0.007	57
3 M	43.601	4	2	2	43.642	-0.041	359	1 MW	70.302	12	2	1	69.938 70.225	-0.035 0.077	155 426
3 MW	44.254	{5 3	2	1	44.164 44.274	0.089 -0.020	76 169	2 VW	70.617	4 2	7	1	70.627	-0.010	213
3 MW	44 70-	(4	4	1	44.355	-0.102	21	1 VW(BR)	71.982	3	5	4	70.907 71.027	0.055	41 223
o MW	44.705	{i	6	2 1	44.653 44.674	0.053	217 188			5	?	3	71.306 71.340		3 <i>2</i> 39
		(2	6	0	44.848	-0.143	96			3	7	2	72.327		127
3 MW	45.409	Jó	3	3	45.093 45.330	0.078	14 316	1 MW(BR)	72.562	6	5	0	72.568 72.779	-0.006	305 0
		12	2	3	45.476 45.635	-0.057	9g 10			7	4	4	72.878		92
3 VW	46.312	1	3	3	46.231	0.081	158			6	4	2	73•730 75•112		27 31
3 W	47.417	12	5	2	47 • 10 2 47 • 358	0.059	6 324	1 W	76.757	10	8	2	76.730 76.923	0.027 -0.166	225 165
		12	6	1	47.379	0.038	60			3	3	4	77.690	00100	66
		3	3	1	47.625 47.910		31 105			4	9	4	77.887 77.948		11
		6	5	0	48.056 48.789		11	1 W	70 505	7	1	2	78.073		75
		. 2	3	3	48.945		31 34		78.503	2 6	0	3	78•523 78•681	-0.020	39 37
3 VW	49.376	{3	6	0	49.369 49.488	0.007 -0.112	0 118			7	4	0	78.729 78.911		74
3 W		. 5	1	2	49.596		25			4	:	4	79.698		59
3 W	49.878	15	2	3	49.958 50.005	-0.080 -0.126	258 12	1 w	80.548	6	6	3	80.277 80.636	0.012	7 147
3 VW	50.581	4	4	1	50.615 51.117	-0.033	149	1 MW	82.169	5	4.	3	82.208	-0.039	379
3 VVW	51.284	[1	7	0	51.183	0.101	2 26	2 MW	83.376	5	1	3	81.549 83.335	0.040	73 190
		6	6	1	51.358 51.412	-0.074 -0.128	33		•						
		6	2	0	51.692		ō								
		5	2	2	51.813 51.926		134								
		3	6	1 2	51.948 52.010		8 89								
_	_	•	•		25 0110	_									

^{*} Numbers given in first column: $3 = \text{Cu } K\alpha$, $2 = \text{Cu } K\alpha_2$, $1 = \text{Cu } K\alpha_1$.

** THETA OBS is the observed value for the Bragg angle θ in degrees minus an absorption correction of the form $57\cdot29578$. $K\lambda$. ($\cos\theta_r/\sin\theta_r+\cos\theta_r/\theta_r$). ($1/4d\theta$), in which θ_r is the Bragg angle in radians, d is the interplanar spacing, and K is an adjustable parameter (see text).

**** ICAL is the calculated powder intensity (without correction for absorption) divided by 100 and multiplied by 1, $\frac{1}{3}$, or $\frac{2}{3}$ in the respective cases α , α_2 , α_1 .

that all these E-silicides and germanides are characterized by a rather restricted range of homogeneity around the 1:1:1 composition.

Our studies based on tiny single-crystal fragments isolated from a crushed sample of TiNiSi(E), kindly sent to us by Professor Paul A. Beck (University of Illinois), did not confirm the large orthorhombic cell found by Spiegel et al. (1963). Approximate values for the cell dimensions were obtained from precession photographs and were refined by a least-squares fit of the lines observed on a powder diffraction diagram taken by the Straumanis method with Cu Ka radiation in a Philips powder camera with 57.3 mm radius. A least-squares program LSCELP was written in FORTRAN for the IBM 7094 computer, similar to the ALGOL program described by Evans, Appleman & Handwerker (1963). An absorption parameter K was introduced in conjunction with the absorption function of Nelson & Riley (1945). An orthorhombic cell with the following cell dimensions was obtained for TiNiSi(E) on least-squares refinement:

$$a_0 = 6.1484 \pm 0.0012$$
, $b_0 = 7.0173 \pm 0.0014$,
 $c_0 = 3.6698 + 0.0007$ Å; $K = (0.92 + 0.24) \times 10^{-3}$.

The limits given are standard deviations. Table 1 is the output of LSCELP and shows the indexing of the powder diagram and the agreement obtained for the observed and calculated theta values.

The systematic absences indicate that the space group is *Pnam* or *Pna*. The observed density (Spiegel et al., 1963) is 5.66 g.cm⁻³ and the calculated number of formula units per cell is 4.00. The ratios of the cell dimensions, the number of atoms per cell and the space group suggest that the structure of the E phase is related to the PbCl₂ structure (C23 type). Trial positional parameters were derived from the PbCl₂ structure, assuming Si in the Pb position and Ti and Ni in the two Cl positions so that Ti would have a larger number of neighbors than Ni. (In Ti- or Ni-containing Laves phases Ti occupies the CN16 position and Ni the CN12 position.) Intensities calculated for this structure were in reasonable agreement with the observed powder diffraction intensities.

Refinement of the structure

The structure was refined on the basis of structure amplitudes, obtained from the visual estimation of intensities recorded with Mo $K\alpha$ radiation by the equi-inclination Weissenberg method. Two irregular crystal flakes were used, one with largest dimension about 0.02 mm for the recording of layers (hk0)—

(hk2) and one with largest dimension about 0.03 mm for layers (h0l)-(h2l). Although exposures up to 200 hours were used only 107 independent reflexions were strong enough to be observed.

The refinement was carried out by several cycles of full-matrix least-squares analysis on the IBM 7094 computer with the program ORFLS (Busing & Levy, 1962). The structure factors were calculated with the scattering factors tabulated in *International Tables for X-ray Crystallography* (1962). No absorption, extinction or dispersion corrections were applied. The standard errors in the structure amplitudes, on which the weights were based, were set equal to 4.0 for all reflexions with $|F_o| \le 40$ and to $\frac{1}{10} |F_o|$ for $|F_o| > 40$. Ten parameters were refined: an overall scale factor and for each atom an isotropic temperature factor and two positional parameters.

The refinement proceeded very rapidly from an initial R value of 0.399 to a final value of 0.086. The non-observed reflexions were excluded from refinement and from the R index; the 002 reflexion was also excluded from the last refinement cycle because of evidence of extinction. An analysis of R as a function of l did not show any trend inconsistent with confinement of all atoms to mirror planes as would be required by space group *Pnam* (D_{2h}^{16}) . No structure factors calculated for the non-observed reflexions in the observed range of the reciprocal lattice exceed significantly the estimated values of F_{\min} , except for the reflexions 120 and 462, which were therefore included in the last refinement cycle. The R value including the non-observed reflexions with $|F_o|$ = $\frac{1}{2}$ F_{\min} and including 002 is 0·191, the large increase probably resulting from the weakness of the photographs and the high values for F_{\min} .

In order to determine whether the three kinds of atom were placed correctly in the three positions one least-squares cycle was run in which only the positional parameters and the 'atom multipliers' (occupancy factors) were varied. This resulted in no significant changes in the parameters. The final parameters are given in Table 2 and the observed and calculated structure factors in Table 3.

Discussion

The interatomic distances listed in Table 4 were calculated with the IBM 7094 FORTRAN program DISTAN, written by one of us (D.P.S.). This program takes the coordinates of the atoms in the asymmetric unit, calculates the symmetry equivalent positions and searches all adjacent cells for distances below a specified limit.

Table 2. Atomic parameters for TiNiSi

Atom	CN	x	$\sigma(x)$. 104	у	$\sigma(y)$. 10^4	z	В	$\sigma(B)$
Ti	15	0.0212	10	0.1803	11	$\frac{1}{4}$	0.12	0.13
Ni	12	0.1420	8	0.5609	8	1	0.37	0.12
Si	9	0.7651	18	0.6229	18	1	0.24	0.19

Table 3. Observed and calculated structure factors for TiNiSi

H K FOBS	FCAL	H K FOBS FCAL
L=0 2 0 *13 4 0 *19 6 0 38 8 0 48 1 1 31 2 1 15 3 1 92 4 1 29	5 9 34 49 30 ~11 ~95	1 7 47 -45 2 7 *35 -19 3 7 *37 -2 4 7 51 -54 5 7 *41 20 1 8 *38 12 2 8 51 -47 3 8 *40 7
L=0 2 0 *13 4 0 *19 4 0 *3 8 0 48 8 0 48 8 1 1 31 2 1 15 5 1 22 6 1 1 48 7 1 22 6 1 1 48 7 1 22 6 1 2 ** 83 7 2 32 7 1 3 69 7 2 37 7 2 37 7 2 37 7 2 37 7 3 3 3 29 4 3 3 3 3 29 4 3 3 3 3 29 4 3 3 3 3 29 6 3 3 3 3 29 6 3 3 3 3 29 6 3 3 3 3 29 6 3 3 3 3 29 6 3 3 3 3 29 6 3 3 3 3 3 29 6 3 3 3 3 3 29 6 3 3 3 3 3 29 6 3 3 3 3 3 29 6 3 3 3 3 3 29 6 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3	5 9 34 49 30	L=2 0 0 E111 -165 2 0 17 -3 4 0 22 -8 6 0 33 -29 8 0 40 -44 1 1 24 -22 2 1 15 -25 5 1 20 15 6 1 39 -41 7 1 24 -17 0 2 19 -19 1 2 40 40 3 2 55 58 4 2 55 55 5 2 35 -34 6 3 39 -41 7 1 34 -17 0 2 19 -19 1 2 40 40 3 2 55 58 4 2 55 55 5 2 35 -34 6 3 30 -14 6 3 30 -33 1 4 36 -38 3 3 24 -20 7 2 35 30 -14 6 3 39 -21 7 2 35 30 -14 6 3 39 -27 7 2 35 30 1 4 36 -8 7 3 4 -8 7 4 36 -8 7 4 5 8 31 1 4 36 -18 3 4 26 -18 3 4 26 -18 3 4 26 -18 3 4 36 -6 3 6 4 36 -6 3 6 4 6 7 3 6 8 31 -0 4 6 8 36 -7 5 6 8 36 -7 5 6 8 36 -7 6 6 8 36 -7 7 8 33 11 7 8 33 -11 7 8 33 -11 7 8 33 -11 7 8 33 -11 7 8 33 -11 7 8 36 -7 8 6 8 36 -7 8 6 8 36 -7 8 6 8 36 -7 8 6 8 36 -7 8 6 8 36 -7 8 7 8 31 -11 8 7 8 33 -11 8 7 8 33 -11 8 7 8 33 -11 8 7 8 33 -11 8 7 8 33 -11 8 8 35 5
		2 4 26 -18 3 4 26 9 4 4 36 28 5 4 54 -60 6 4 34 -12 1 5 48 54 3 5 29 7 4 5 31 -11 5 5 33 22 6 5 36 -7 0 6 47 51 2 6 36 -7 1 6 47 51 2 6 36 -36 -7 1 6 47 51 2 6 36 -36 -7 3 6 431 -0 4 6 433 -0 4 6 433 -0 5 4 36 -36 36 -7 1 7 432 18 3 7 434 -28 0 8 47 45 -28
L=1 24 0 *20 4 0 *20 4 0 *20 5 8 0 1 *10 1 1 *38 3 1 48 3 1 48 5 1 72 5 1 72 7 1 *23 1 2 2 50 3 4 2 37 7 2 48 6 7 3 *36 6 3 *36 6 3 *36 6 4 4 *31 7 6 4 *31 7 6 4 *31 7 6 4 *31 7 6 6 *34 7 7 7 8 8 8 8 9 9 9 9 9 9 9 9 9 9 9 9 9	93 7 7-24 61 133-85 -85 -44-38 -25-60 -88 45-16-25 -16-22 154-20 -21 15-24 -10 17-15 12-5 5 16-25 16-26 11-27 17-37 18-3	2 0 51 -58 4 0 *26 -7 6 0 *30 16 8 0 51 -52 0 1 *18 -2 1 1 26 -19 2 1 54 55 3 1 35 -31 4 1 28 27 5 1 51 56 6 1 *24 21 1 2 53 -55 2 2 33 -29 3 2 *21 10 6 2 28 23 5 2 *24 -20 6 2 32 33
2 3 23 3 3 *26 4 3 *29 5 3 *36 7 3 *39 1 4 *25 2 4 *26 3 4 *31 5 4 *27 6 4 *37 7 5 *28 1 5 *28	-24 10 -19 17 15 12 -5 -91 -13 -37 5 16 64 0	L=4 0 0 82 97 2 0 *27 1 4 0 *30 9 1 1 *22 12 2 1 *22 -4 3 1 46 -67 4 1 *24 14 5 1 *26 -11 6 1 33 31 0 2 *23 9 1 2 *23 -25 3 2 37 -38 4 2 23 33 -37
2 5 39 4 5 49 5 5 *36 6 5 *39 1 6 *32 2 6 *32 3 6 *34 4 6 40 5 6 *38 6 6 *41 0 7 38	-38 -36 11 43 40 -23 -8 25 -3 36	L=5 2 0 35 34 0 1 *26 -1 1 1 *26 9 2 1 37 -36 3 1 *27 18 4 1 *28 -20 5 1 38 -38 L=6 0 0 56 -61

* Non-observed, value listed under FOBS is F_{min}.
** Non-observed, but included in least-squares cycle.
E Left out of least cycle because of extinction.

Table 4. Observed interatomic distances for TiNiSi
The standard deviations in the distances average about 0.012 Å.
The distances are given for the atoms listed in Table 2, going around each atom clockwise in Fig. 1(a) starting at 12 o'clock.
Only distances smaller than 3.3 Å are listed.

	Ti	Ni	Si
Towards atoms in the same plane	Si 2.603 Ti 3.226 Ni 2.772 Ti 3.226 Ni 2.881	Ti 2.881 Si 2.344 Si 2.358 Ti 2.772	Ni 2·358 Ni 2·344 Ti 2·603
Towards atoms in neighboring planes	Ti 3·136 Ni 2·891 Si 2·646 Ni 2·770 Si 2·574	Si 2·314 Ti 2·891 Ti 2·770 Ni 2·673	Ni 2·314 Ti 2·646 Ti 2·574

Fig. 1(a) gives the projection of the structure down the z axis. All atoms are situated in the mirror planes at $z=\frac{1}{4}$ and $\frac{3}{4}$ perpendicular to the z axis. The Si atom has a total of 9 near neighbors: 6 atoms (4 Ti + 2 Ni) at the corners of a triangular right prism (prism axis parallel to z) and 3 more atoms (2 Ni+1 Ti) outside the rectangular prism faces in the same mirror plane as the center atom. The shortest Si-Si distances are 3.554 Å. This type of 9-coordination for Si is very commonly found for the metalloid atom in silicides, borides, phosphides and carbides (Aronsson, 1960).

The Ni atom has a total of 12 near neighbors: 8 atoms (4 Ti + 2 Ni + 2 Si) at the corners of a quadrangular right prism (prism axis parallel to z) and 4 more atoms (2 Ti+2 Si) outside the side prism faces in the same mirror plane as the center atom. The twelvefold coordination around the Ni atom can be derived from a body-centered cubic arrangement (CN14) by elongating the cube in the direction of the z axis, so that the two Ni-Ni distances along the z axis become 3.670 Å, which is too long to be considered as a bonding distance. By contrast, in the frequently occurring CuAl₂ structure (C16 type) the Cu is at the center of a square anti-prism, which leads to a contraction along the z axis and larger distances to the four atoms in the equatorial plane, thus resulting in an effective coordination number of 10 for the Cu atom.

The Ti atom has a total of 15 near neighbors: 10 atoms (2 Ti, 4 Ni and 4 Si) at the corners of a pentagonal right prism (prism axis parallel to z) and 5 more atoms (2 Ti, 2 Ni and 1 Si) outside the rectangular prism faces in the same mirror plane. Two more Ti atoms are on the z axis at a distance of 3.670 Åfrom the center atom, that is, 0.45 Å farther away than the Ti atoms in the equatorial plane. One more Si atom in the center plane is at a distance of 3.482 Å. In the well known icosahedral arrangement of atoms as found in Laves phases, β -W phases and phases related to the σ phase, 10 atoms form a five-sided antiprism, which allows a closer approach of the atoms on the z axis and moves the atoms in the equatorial plane outward, thus giving the center atom 12 near neighbors.

Several transition element intermetallic compounds containing Si, Ge, P, etc. have been assigned the

(anti-) PbCl₂-type (C23) structure (for references see Jellinek, 1959). Inspection of the cell dimensions of these compounds shows that most of them have a/b values in the range 0·67-0·73, namely: Rh₂Ge, δ-Ni₂Si, Co₂Si, Ru₂Si, Ir₂Si (Bhan & Schubert, 1960), Rh₂Si, Rh₂Ta (Giessen, Ibach & Grant, 1964), Rh₂Sn, Pd₂Sn, Pd₂Al, Pd₂Ga, Pd₂In. However TiNiSi (E), Co₂P, and Ru₂P (Rundquist, 1960) have a/b values in the range 0·84-0·88, which is within that reported for several other compounds with the PbCl₂-type or anti-PbCl₂-type structure, for instance ThS₂ (Zachariasen, 1949b), ThSe₂(D'Eye, 1953), Ca₂Si, Ca₂Ge,

CaH₂ (Bergsma & Loopstra, 1962), SnCl₂ (van den Berg, 1961), and PbCl₂ itself. The atomic parameters determined for representatives of these two groups are not very different but the smaller a/b value for the compounds in the first group causes the numbers of neighbors for the various atoms to be significantly different from those we have described for TiNiSi. (A comparison of the Co₂Si and Co₂P structures is given by Rundquist, 1960).

The coordination polyhedra for the atoms in these structures are very complicated and the atomic surroundings may be best compared by examining the

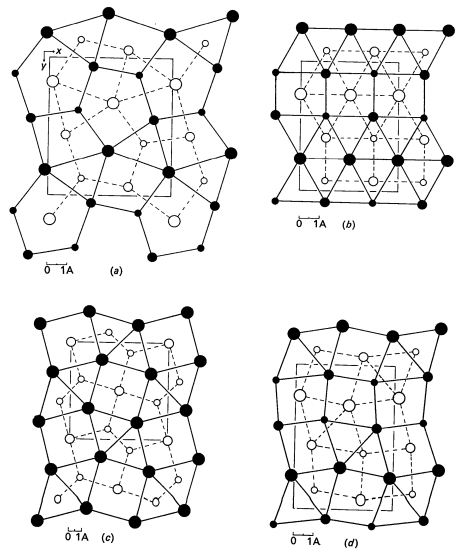


Fig. 1 (a). Projection of TiNiSi down the z axis. Solid-line net at $-\frac{1}{4}$ (and $\frac{3}{4}$), broken-line net at $\frac{1}{4}$. (b) Projection of θ -Ni₂Si (Ni₂In type) down hexagonal [110]. Solid-line net at $-\frac{1}{4}$ (and $\frac{3}{4}$) and broken-line net at $\frac{1}{4}$ of [110] lattice repeat. (c) Projection of U₃Si₂ down the (tetragonal) z axis. Solid-line net at $z=\frac{1}{2}$, broken-line net at z=0. (d) Projection of Co₂Si down the z axis. Nets as in (a). The atoms are to be identified as follows:

	Large circles	Intermediate circles	Small circles
(a)	Ti	Ni	Si
(b)	Ni(I)	Ni(II)	Si
(c)	U(ÌÌ)	U(Ì)	Si
(d)	Co(I)	Co(II)	Si

bonding networks lying in the mirror planes. The networks in Fig. 1 are drawn so that any distance between unconnected atoms is greater than the distance between any pair of connected atoms (Wells, 1954). Co₂Si, a representative of the first group, was first described by Geller (1955) as a distorted Ni₂In structure. A small distortion of the Ni₂In nets (which are drawn in Fig. 1(b) for θ -Ni₂Si; Toman, 1952) results in the Co₂Si structure with rather similar coordination polyhedra for the two transition metal atoms [Fig. 1(d)], but a much larger distortion is needed to accommodate the larger Ti atoms in the TiNiSi structure [Fig. 1(a)]. Going from the θ -Ni₂Si structure (Ni₂In type), through Co₂Si to the TiNiSi structure the numbers of near neighbors change as follows:

θ -Ni ₂ Si		C	o ₂ Si	TiNiSi		
Ni(I)	14	Co(I)	13(+1?)	Ti	15(+3?)	
Ni(II)	11	Co(II)	13	Ni	12	
Si(In)	11	Si	10(+5?)	Si	9(+1?)	

(The definition of 'neighbors' is, of course, to some degree arbitrary. The numbers in parentheses indicate the shading between neighbors and non-neighbors.)

The Co₂Si network is only a small distortion of the well known 3².4.3.4 network which is shown by solid lines in the diagram of U₃Si₂ [Fig. 1(c); Zachariasen, 1949(a)] and which also occurs in the CuAl₂ structure type (Frank & Kasper, 1959) and as the secondary layer in the σ phase. The U₃Si₂ structure has two kinds of planar nets, in alternating sequence: the first one is the 3².4.3.4 net already mentioned, which is formed by U(II) atoms, and the second one is a net consisting of pentagons formed by 3 Si and 2 U(I). The surroundings of the atoms in U₃Si₂ are very similar to those in TiNiSi which is built of two identical planar nets in different orientation, each consisting of triangles, quadrangles and pentagons. U(I) has surroundings similar to those of Ni with a practically nondistorted cube; U(II) corresponds to Ti and Si to Si.

The only other ternary compound described with the PbCl₂ structure is Pb(OH)Cl (Brasseur, 1940).

Professor P. A. Beck also kindly sent us a specimen of an alloy MnCoSi which we found to be indexable on an E-phase type cell with $a_0 = 5.8543 \pm 0.0016$, $b_0 = 6.8526 \pm 0.0017$, $c_0 = 3.6853 \pm 0.0013$ Å. The alloy presumably has the E-phase structure but no parameter refinement was carried out.

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