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. A. SEMILETOV, T. V. HITAR, Kristallogativa Some Proposals for Transformation Mechanisms of the Li₂ZrF₆, Trirutile and Na₂SiF₆ Structure Types: Simple Cation Rearrangements

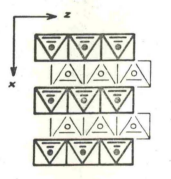
JEAN GALY AND STEN ANDERSSON

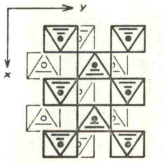
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Under high pressure, rutile transforms into a structure of the α -PbO₂ type. In a previous paper (1) we proposed a simple mechanism for this transformation which is illustrated in Fig. 1. We now suggest similar cation displacements to interconvert the Li₂ZrF₆, trirutile, Na₂SiF₆ and columbite structure types.

When Li_2NbOF_5 was reported (2) as being isostructural with Li_2ZrF_6 (3), a picture was made showing its relation to the rutile structure. In Figs. 2a and b, the structures of trirutile and Li_2NbOF_5 are compared; Fig. 3 shows how the Li_2ZrF_6 structure type can transform into the trirutile type if 50% of the cations are shifted in the way





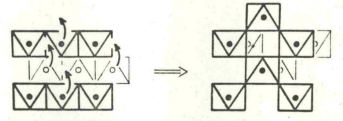
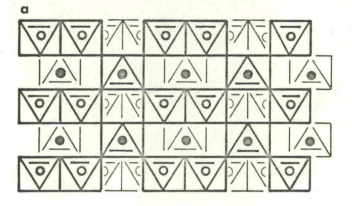


FIG. 1. The rutile structure (upper left) is compared with the α -PbO₂ structure type (upper right). Below is shown a possible mechanism for the rutile $\rightarrow \alpha$ -PbO₂ transformation. Cation movements are indicated with arrows.

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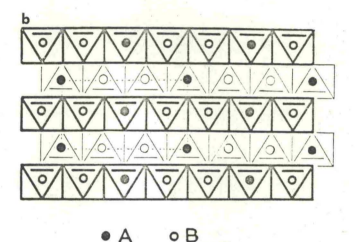


FIG. 2. In (a) the Li_2ZrF_6 structure type (Li_2NbOF_5) is given. In (b) the trirutile structure of the composition AB₂X₆. In both the structures, A corresponds to filled circles, and open circles are the B-atoms.

indicated by the curved arrows, and without disorder. To our knowledge, no such transformation has been reported for the compounds Li_2NbOF_5 and Li_2ZrF_6 . However, α -Li_2SnF₆ of the Li_2ZrF_6 type has been found to transform to a structure of the trirutile type, designated β -Li_2SnF₆ at 510°C (4).

Another interesting transformation was reported recently (5, 6). At about 400°C, α -Li₂GeF₆, of the trirutile type, transforms to β -Li₂GeF₆ which has the Na₂SiF₆ type structure (7). Figure 4 shows the Na₂SiF₆ structure projected along the *c*-axis, and Fig. 5 the trirutile structure in an analogous projection. The geometrical relation between these structures can be shown by considering the Na₂SiF₆ type. The cations in the octahedra drawn with heavy lines are at $z = \frac{1}{2}$, those in the lighter octahedra at z = 0. If the metal atoms at z = 0 in *m*-planes move to $z = \frac{1}{2}$, and those at $z = \frac{1}{2}$ in *n*-planes to z = 0, the trirutile structure is obtained. This involves half the cations passing through octahedral faces along half the *c*-axis; the anion lattice remains intact. If the A and B atoms keep their relative order during the transformation, the ordered Na₂SiF, structure type may, in the reverse way, be geometrically derived directly from the ordered Trirutile type. That β -Li₂SnF₆ and β -Li₂GeF₆ are formed as ordered compounds, with, respectively, the trirutile and Na₂SiF₆ structures, indicates that the geometrical proposals described here may very well be correct mechanisms. FIG. type.

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All these compounds have approximately the same (hexagonally close packed) anion arrangement. At very high pressures the preferred cation arrangement seems to be like that in the α -PbO₂

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STRUCTURAL TRANSFORMATION MECHANISMS

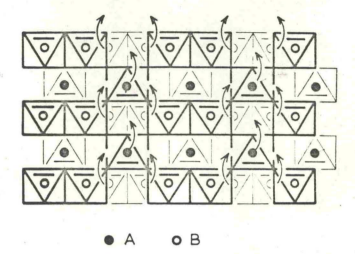


FIG. 3. If cations are shifted in the way described by arrows, the trirutile structure is formed from the Li_2ZrF_6 structure type.

structure type. Amongst the ternary compounds of general formula AB_2X_6 a similar arrangement of (ordered) cations occurs in columbite FeNb₂O₆. By analogy with the rutile $\rightleftharpoons \alpha$ -PbO₂ transformation,

we might therefore expect the AB_2X_6 compounds discussed above to transform at high enough pressures to a ternary analog of α -PbO₂, possibly with the cations ordered as in FeNb₂O₆.

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approximately the d) anion arrangete preferred cation hat in the α -PbO:

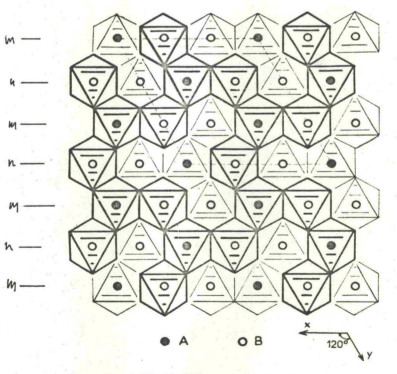


FIG. 4. The Na₂SiF₆ structure type.

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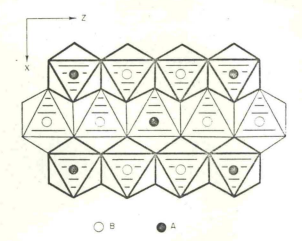


FIG. 5. The trirutile structure type.

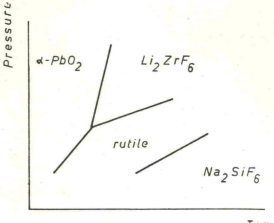
It would be interesting to study the effects of applied pressure in the high temperature experiments referred to above.

In the meantime a tentative pressure-temperature phase diagram is given in Fig. 6. Its construction is based on the observations that

- (a) α -LiSnF₆ transforms to β -Li₂SnF₆,
- (b) α -Li₂GeF₆ transforms to β -LiGeF₆ at high temperatures and
- (c) rutile transforms to a structure of the α -PbO₂ type at high pressure.

Acknowledgment

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FIG. 6. Tentative temperature-pressure phase diagram for the various MX_2 structure types discussed.

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