

# $E2_3$ ( $\text{LiIO}_3$ ) Structure (*Obsolete*): ABC3\_hP10\_182\_c\_b\_g-001

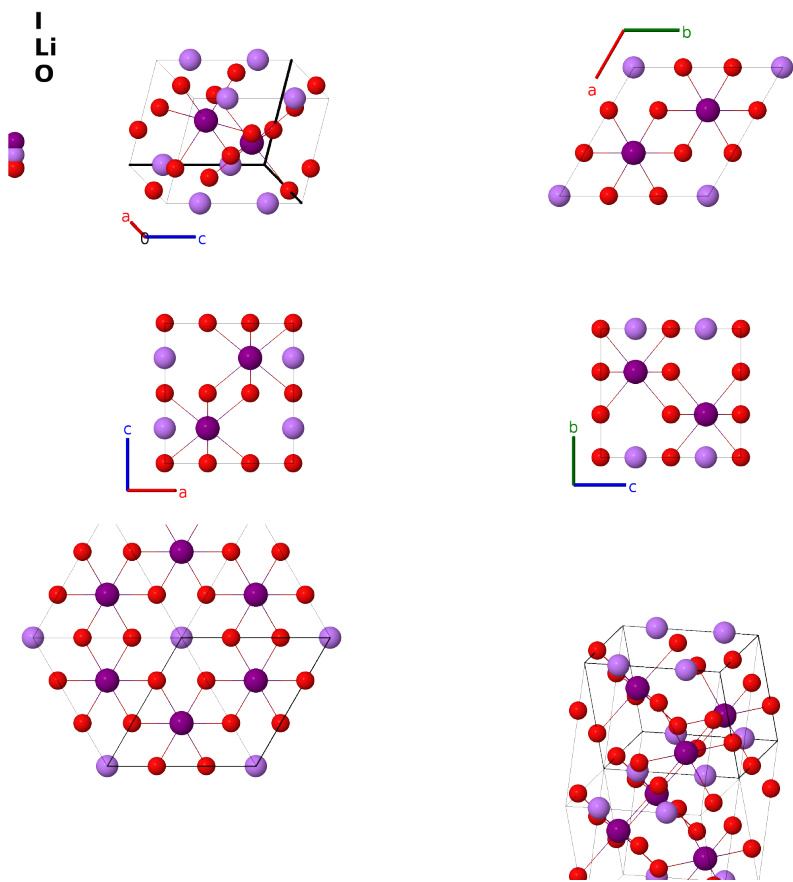
This structure previously used the label(s) **ABC3\_hP10\_182\_c\_b\_g**. Calls to these addresses will be redirected here.

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<https://aflow.org/prototype-encyclopedia/6B26>

[https://aflow.org/prototype-encyclopedia/ABC3\\_hP10\\_182\\_c\\_b\\_g-001](https://aflow.org/prototype-encyclopedia/ABC3_hP10_182_c_b_g-001)



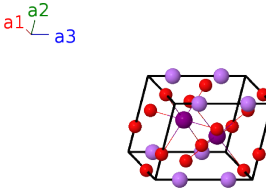
|                                    |                         |
|------------------------------------|-------------------------|
| Prototype                          | $\text{ILiO}_3$         |
| AFLOW prototype label              | ABC3_hP10_182_c_b_g-001 |
| <i>Strukturbericht</i> designation | $E2_3$                  |
| ICSD                               | 20012                   |
| CCDC                               | 1597858                 |
| Pearson symbol                     | hP10                    |
| Space group number                 | 182                     |

Space group symbol  $P6_322$

AFLOW prototype command `aflow --proto=ABC3_hP10_182_c_b_g-001`  
`--params=a, c/a, x3`

- $\text{LiIO}_3$  is known to exist in three forms:
- $\alpha\text{-LiIO}_3$ , stable below 470K:
  - (Zachariasen, 1931) originally determined that the structure of  $\alpha\text{-LiIO}_3$  was in space group  $P6_322$  #182, which (Hermann, 1937) designated *Strukturbericht E*<sub>23</sub>. (this structure)
  - (Rosenzweig, 1966) subsequently determined that this structure was incorrect because of the small sample size, and determined that the true structure was in space group  $P6_3$  #173.
- $\beta\text{-LiIO}_3$ , stable from 573K up to the melting point at 708K.
- $\gamma\text{-LiIO}_3$ , stable between the  $\alpha$ - and  $\beta$ -phases, with an orthorhombic structure in space group  $Pna2_1$  #33.
- The ICSD entry is from (Butolin, 1975). If we can obtain a copy we will report on their research into this structure.

### Hexagonal primitive vectors

$$\begin{aligned}
 \mathbf{a}_1 &= \frac{1}{2}a \hat{\mathbf{x}} - \frac{\sqrt{3}}{2}a \hat{\mathbf{y}} \\
 \mathbf{a}_2 &= \frac{1}{2}a \hat{\mathbf{x}} + \frac{\sqrt{3}}{2}a \hat{\mathbf{y}} \\
 \mathbf{a}_3 &= c \hat{\mathbf{z}}
 \end{aligned}$$


### Basis vectors

|                   | Lattice coordinates                                                                |     | Cartesian coordinates                                                                                         | Wyckoff position | Atom type |
|-------------------|------------------------------------------------------------------------------------|-----|---------------------------------------------------------------------------------------------------------------|------------------|-----------|
| $\mathbf{B}_1$    | $= \frac{1}{4} \mathbf{a}_3$                                                       | $=$ | $\frac{1}{4}c \hat{\mathbf{z}}$                                                                               | (2b)             | Li I      |
| $\mathbf{B}_2$    | $= \frac{3}{4} \mathbf{a}_3$                                                       | $=$ | $\frac{3}{4}c \hat{\mathbf{z}}$                                                                               | (2b)             | Li I      |
| $\mathbf{B}_3$    | $= \frac{1}{3} \mathbf{a}_1 + \frac{2}{3} \mathbf{a}_2 + \frac{1}{4} \mathbf{a}_3$ | $=$ | $\frac{1}{2}a \hat{\mathbf{x}} + \frac{\sqrt{3}}{6}a \hat{\mathbf{y}} + \frac{1}{4}c \hat{\mathbf{z}}$        | (2c)             | I I       |
| $\mathbf{B}_4$    | $= \frac{2}{3} \mathbf{a}_1 + \frac{1}{3} \mathbf{a}_2 + \frac{3}{4} \mathbf{a}_3$ | $=$ | $\frac{1}{2}a \hat{\mathbf{x}} - \frac{\sqrt{3}}{6}a \hat{\mathbf{y}} + \frac{3}{4}c \hat{\mathbf{z}}$        | (2c)             | I I       |
| $\mathbf{B}_5$    | $= x_3 \mathbf{a}_1$                                                               | $=$ | $\frac{1}{2}ax_3 \hat{\mathbf{x}} - \frac{\sqrt{3}}{2}ax_3 \hat{\mathbf{y}}$                                  | (6g)             | O I       |
| $\mathbf{B}_6$    | $= x_3 \mathbf{a}_2$                                                               | $=$ | $\frac{1}{2}ax_3 \hat{\mathbf{x}} + \frac{\sqrt{3}}{2}ax_3 \hat{\mathbf{y}}$                                  | (6g)             | O I       |
| $\mathbf{B}_7$    | $= -x_3 \mathbf{a}_1 - x_3 \mathbf{a}_2$                                           | $=$ | $-ax_3 \hat{\mathbf{x}}$                                                                                      | (6g)             | O I       |
| $\mathbf{B}_8$    | $= -x_3 \mathbf{a}_1 + \frac{1}{2} \mathbf{a}_3$                                   | $=$ | $-\frac{1}{2}ax_3 \hat{\mathbf{x}} + \frac{\sqrt{3}}{2}ax_3 \hat{\mathbf{y}} + \frac{1}{2}c \hat{\mathbf{z}}$ | (6g)             | O I       |
| $\mathbf{B}_9$    | $= -x_3 \mathbf{a}_2 + \frac{1}{2} \mathbf{a}_3$                                   | $=$ | $-\frac{1}{2}ax_3 \hat{\mathbf{x}} - \frac{\sqrt{3}}{2}ax_3 \hat{\mathbf{y}} + \frac{1}{2}c \hat{\mathbf{z}}$ | (6g)             | O I       |
| $\mathbf{B}_{10}$ | $= x_3 \mathbf{a}_1 + x_3 \mathbf{a}_2 + \frac{1}{2} \mathbf{a}_3$                 | $=$ | $ax_3 \hat{\mathbf{x}} + \frac{1}{2}c \hat{\mathbf{z}}$                                                       | (6g)             | O I       |

### References

- [1] W. H. Zachariasen and F. A. Barta, *Crystal Structure of Lithium Iodate*, Phys. Rev. **37**, 1626–1630 (1931), doi:10.1103/PhysRev.37.1626.
- [2] C. Hermann, O. Lohrmann, and H. Philipp, eds., *Strukturbericht Band II 1928-1932* (Akademische Verlagsgesellschaft M. B. H., Leipzig, 1937).
- [3] S. A. Butolin, L. F. Belova, R. N. Samoylova, O. M. Kotenko, I. M. Dokuchaeva, and N. M. Ivanova, *Optical and physico-chemical properties of  $\alpha\text{LiIO}_3$  monocrystal*, Izvestiya Akademii Nauk SSSR, Neorganicheskie Materialy **11**, 862–865 (1975).

**Found in**

- [1] A. Rosenzweig and B. Morosin, *A reinvestigation of the crystal structure of  $\text{LiIO}_3$* , Acta Cryst. **20**, 758–761 (1966), doi:10.1107/S0365110X66001804.

**First cited in**

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