

ReSi₂ Structure:

AB2_oI6_71_a_e-002

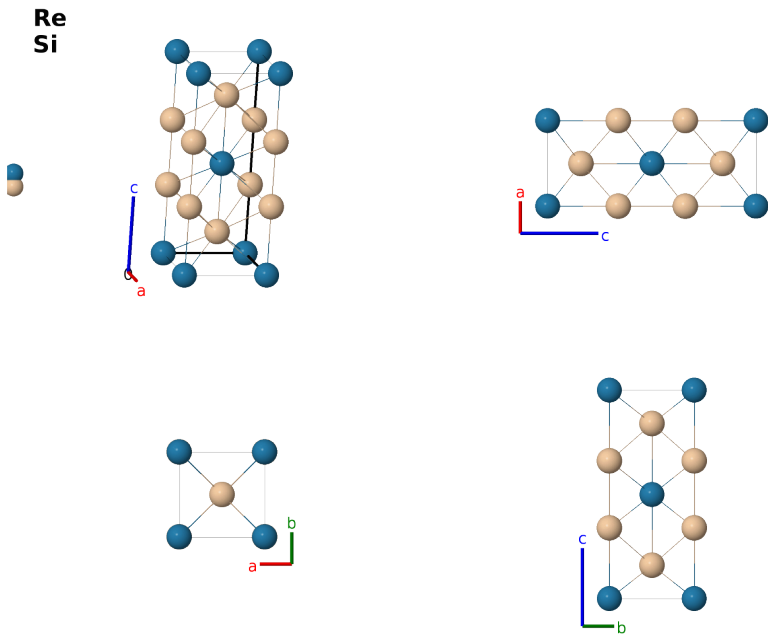
This structure previously used the label(s) **AB2_oI6_71_a_i**. Calls to these addresses will be redirected here.

Cite this page as:

- H. Eckert, S. Divilov, M. J. Mehl, D. Hicks, A. C. Zettel, M. Esters, X. Campilongo, and S. Curtarolo, *The AFLOW Library of Crystallographic Prototypes: Part 4*, Comput. Mater. Sci. **240**, 112988 (2024), doi: 10.1016/j.commatsci.2024.112988.
- D. Hicks, M. J. Mehl, M. Esters, C. Oses, O. Levy, G. L. W. Hart, C. Toher, and S. Curtarolo, *The AFLOW Library of Crystallographic Prototypes: Part 3*, Comput. Mater. Sci. **199**, 110450 (2021), doi: 10.1016/j.commatsci.2021.110450.

<https://aflow.org/prototype-encyclopedia/9FBX>

https://aflow.org/prototype-encyclopedia/AB2_oI6_71_a_e-002

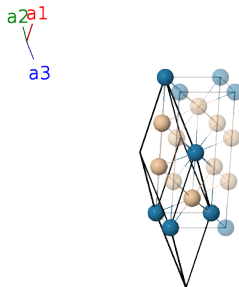


Prototype	ReSi ₂
AFLOW prototype label	AB2_oI6_71_a_e-002
ICSD	38274
CCDC	1608942
Pearson symbol	oI6
Space group number	71
Space group symbol	<i>Immm</i>
AFLOW prototype command	<code>aflow --proto=AB2_oI6_71_a_e-002 --params=a,b/a,c/a,x₂</code>

- The original references describe MoPt₂ and ReSi₂ in different orientations, so they have nominally different AFLOW prototype labels, AB2_oI6_71_a_g and AB2_oI6_71_a_i, respectively. When we apply our AFLOW prototype label rules, however, the label for both structures becomes AB2_oI6_71_a_e. The two structures are generated by the same symmetry operations with different sets of parameters (**--params**) specified in their corresponding CIF files.

Body-centered Orthorhombic primitive vectors

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



Basis vectors

	Lattice coordinates		Cartesian coordinates	Wyckoff position	Atom type
$\mathbf{B}_1$	=	0	=	0	(2a) Re I
$\mathbf{B}_2$	=	$x_2 \mathbf{a}_2 + x_2 \mathbf{a}_3$	=	$ax_2 \hat{\mathbf{x}}$	(4e) Si I
$\mathbf{B}_3$	=	$-x_2 \mathbf{a}_2 - x_2 \mathbf{a}_3$	=	$-ax_2 \hat{\mathbf{x}}$	(4e) Si I

References

- [1] T. Siegrist, F. Hulliger, and G. Travaglini, *The crystal structure and some properties of ReSi_2* , J. Less-Common Met. **92**, 119–129 (1983), doi:10.1016/0022-5088(83)90233-3.

First cited in

M. J. Mehl, D. Hicks, C. Toher, O. Levy, R. M. Hanson, G. L. W. Hart, and S. Curtarolo, *The AFLOW Library of Crystallographic Prototypes: Part 1*, Comput. Mater. Sci. **136**, S1-828 (2017). doi: 10.1016/j.commatsci.2017.01.017