

MoSi₂ (*C*11_b) Structure:

AB2_tI6_139_a_e-001

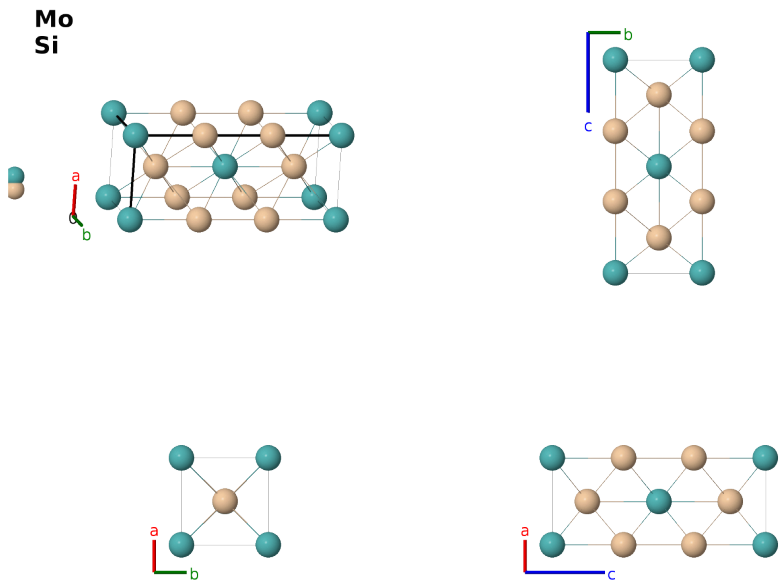
This structure previously used the label(s) **AB2_tI6_139_a_e**. Calls to these addresses will be redirected here.

Cite this page as:

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<https://aflow.org/prototype-encyclopedia/NG7F>

https://aflow.org/prototype-encyclopedia/AB2_tI6_139_a_e-001



Prototype	MoSi ₂
AFLOW prototype label	AB2_tI6_139_a_e-001
<i>Strukturbericht</i> designation	<i>C</i> 11 _b
ICSD	85756
CCDC	1645669
Pearson symbol	tI6
Space group number	139
Space group symbol	<i>I</i> 4/ <i>mmm</i>
AFLOW prototype command	<code>aflow --proto=AB2_tI6_139_a_e-001</code> <code>--params=<i>a</i>, <i>c</i>/<i>a</i>, <i>z</i>₂</code>

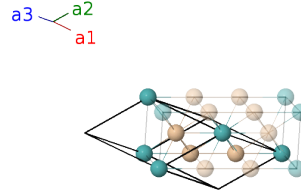
Other compounds with this structure

AgZr₂, AuHf₂, AuMn₂, AuZr₂, BeAu₂, CdTi₂, CuHf₂, CuTl₂, CuZr₂, DyAg₂, DyAu₂, DyTl₂, ErAu₂, GdAg₂, GdAu₂, HfAu₂, LuAu₂, MnAu₂, MoGe₂, MoU₂, β -PdBi₂, PdHf₂, PdTi₂, PtHf₂, ReSi₂, TaNi₂, TiAu₂, WSi₂, ZrAu₂, ZrPd₂

- When $c = 3a$ and $z_2 = 1/3$ the atoms are on the sites of a body-centered cubic (*A2*) lattice. For MoSi₂ itself, (Harada, 1998) gives $c/a = 2.45$ and $z_2 = 0.3353$. Other compounds in this structure have very different values of c/a and even z_2 .
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Body-centered Tetragonal primitive vectors

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



Basis vectors

	Lattice coordinates		Cartesian coordinates	Wyckoff position	Atom type
$\mathbf{B}_1$	=	0	=	0	(2a) Mo I
$\mathbf{B}_2$	=	$z_2 \mathbf{a}_1 + z_2 \mathbf{a}_2$	=	$cz_2 \hat{\mathbf{z}}$	(4e) Si I
$\mathbf{B}_3$	=	$-z_2 \mathbf{a}_1 - z_2 \mathbf{a}_2$	=	$-cz_2 \hat{\mathbf{z}}$	(4e) Si I

References

- [1] Y. Harada, M. Morinaga, D. Saso, M. Takata, and M. Sakata, *Refinement of crystal structure in MoSi₂*, Intermetallics **6**, 523–527 (1998), doi:10.1016/S0966-9795(97)00102-7.

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