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Crystal Structures of α- And β-Nitrogen Trifluoride

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The Crystal Structures of α- and β-F₂ revisited


Dedicated to Professor Wolfgang Bensch on the occasion of his 65th birthday

Abstract: The crystal structures of α-F₂ and β-F₂ have been reinvestigated using neutron powder diffraction. For the low-temperature phase α-F₂, which is stable below circa 45.6 K, the monoclinic space group C2/c with lattice parameters a = 5.4780(12), b = 3.2701(7), c = 7.2651(17) Å, β = 102.088(18)°, V = 127.26(5) Å³, mS8, Z = 4 at 10 K can now be confirmed. The structure model was significantly improved, allowed for the anisotropic refinement of the F atom, and an F–F bond length of 1.404(12) Å was obtained which is in excellent agreement with spectroscopic data and high-level quantum chemical predictions. The high-temperature phase β-F₂, stable between circa 45.6 K and the melting point of 53.53 K, crystallizes in the cubic primitive space group Pm3n with the lattice parameter a = 6.5314(15) Å, V = 278.62(11) Å³, Z = 8, cP16, at 48 K. β-F₂ is isotypic to γ-O₂ and δ-N₂. The centers of gravity of the F₂ molecules are arranged like the atoms in the Cr₃Si structure type.

Introduction

The crystal structures of the chemical elements belong to the fundamental knowledge of chemistry. Atom distances, bond lengths and angles can be determined precisely and the data serve as benchmarks for quantum chemistry. To our striking surprise the crystal structure of α-fluorine, the polymorph stable below circa 45.55 K, has so far only once been investigated, and the β-allotrope, stable between the melting point (53.53 K) and the phase change temperature, also only once.[1–3] Both crystal structures were investigated using X-ray diffraction, for β-F₂ on single crystals grown from a mixture with argon and for α-F₂ only on powders as due to the phase change from β to α no single crystals could be obtained. Maybe it is not so surprising that the crystal structures of α- and β-F₂ have only been determined once as solid F₂ is still extremely reactive. In the case of the β-F₂ structure determination the authors reported several explosions when the solid F₂ reacted – due to the phase change at circa 45.6 K – with the copper ampoule that was used as the sample holder.[1] Additionally, the authors struggled with the strong X-ray reflection intensities of the Cu ampoule and the very weak reflections of α-F₂. The authors concluded that α-F₂ crystallizes in the monoclinic crystal system, probably in space group C2/m, but space group C2/c could not be ruled out. Years later, the original diffraction data were reinterpreted by Pauling and coworkers and space group C2/c was found to be more likely correct. All these assumptions were based on 36 collected reflections.[4] Thus, the space group of α-F₂ is still uncertain[5] and the problems due to the Cu sample holder naturally had an influence on the determined atom positions, bond lengths and displacement parameters.

Up to 1964 fluorine was the only stable element of which no crystal structure and not even a powder pattern had been reported but Lipscomb and coworkers succeeded after significant efforts to obtain a proper single crystal of β-F₂ in a sealed glass capillary.[2,3] In those days, diffraction intensities were measured visually, which was often ambiguous, and the reaction of F₂ with the residual moisture on the glass walls always led to the formation of HF. The authors faced the problem to distinguish between two likely cubic space groups, P4₃m (No. 218) and Pm₃n (No. 223), based on a total of 42 reflections which dropped significantly in intensity for higher diffraction angles. They chose the higher symmetric space group as they expected a disordered structure and explored in great efforts many structure models in this space group.[2,3] Neither the bond length of the F₂ molecules, and naturally not the anisotropic displacement parameters of the fluorine atoms could be refined.

We therefore reinvestigated the crystal structures of α- and β-fluorine using powder neutron diffraction and Rietveld refinement. We also selected Cu as a sample holder, as Cu can be thoroughly passivated with F₂. Aluminum would be almost transparent for neutrons and vanadium is almost exclusively an incoherent scatterer, but both cannot be passivated as thoroughly as Cu. Using neutron diffraction, the absorption by the Cu ampoule plays only a minor role and therefore the reflections of crystalline fluorine are much more easily detectable. Also in contrast to X-ray diffraction, the atom form factors for neutron scattering do not depend on the scattering angle and so the intensities of the reflections show a much less decrease (which may still happen due to absorption, texture, ...) with increasing scattering angle. We therefore obtained models providing much more precise lattice parameters, atomic coordinates as well as bond lengths for the two polymorphs of fluorine.

Results and Discussion

α-Fluorine

In α-fluorine there are four F₂ molecules per unit cell and the previous structure model is as follows: At 23 K, the lattice...
parameters were reported as $a = 5.50(1)$, $b = 3.28(1)$, $c = 7.284(1)$ Å, $\beta = 102.17(2)^\circ$, $V = 128.4(5) \text{ Å}^3$, mS8, the calculated density was 1.97 g/cm$^3$, and the calculated molar volume 19.3 cm$^3$/mol.$^{[1]}

The structure was reported to be very similar to α-O$_2$. Each F$_2$ molecule is surrounded by six others in the shape of a distorted hexagon, so a hexagonally packed layer results. The main difference to α-O$_2$ was that the F$_2$ molecules are tilted, probably only by about 11°, away from the normal onto the planes of the hexagons. As the tilts are to alternate sides in consecutive hexagonal layers, the unit cell is approximately doubled in the $c$ direction in comparison to α-O$_2$ ($a = 5.403$, $b = 3.443$, $c = 4.247$ Å, $\beta = 117.841$, $V = 69.7$ Å$^3$, $mC4\bar{1}$, $T = 22$ K).$^{[6]}$ It is said that this difference results from the antiferromagnetic ordering of the O$_2$ molecules, which is absent in case of F$_2$ molecules. α-O$_2$ crystallizes with an arrangement of the O$_2$ molecules similar to cubic closed packing.

We recorded powder neutron diffraction patterns of α-F$_2$ at 10 and 46 K. In the following, only the structural details of the 10 K measurement will be discussed. A Rietveld refinement was carried out (Error! Reference source not found.) and it unambiguously shows space group $C2/c$ (No. 15) to be correct as the refinement in space group $C2/m$ gave very poor results (see Supporting Information). The refined lattice parameters are $a = 5.4780(12)$, $b = 3.2701(7)$, $c = 7.2651(17)$ Å, $\beta = 102.088(18)^\circ$, $V = 127.26(5)$ Å$^3$, mS8, $Z = 4$ at 10 K. Therefore, α-F$_2$ is not isotypic to α-O$_2$.$^{[6]}$ The crystal structure of α-F$_2$ is shown in Error! Reference source not found.. Error! Reference source not found. holds the atomic coordinates, Wyckoff positions, site symmetries and occupancies, as well as the isotropic displacement parameters, Error! Reference source not found. the anisotropic displacement parameters of the F atom in α-F$_2$. Error! Reference source not found. holds selected crystallographic details.

![Figure 1](image1.png)

**Figure 1.** Powder neutron diffraction pattern of α-F$_2$ at 10 K. Black dots indicate measured intensity, fitted curve in red (Rietveld refinement for F$_2$, LeBail fit for Cu), difference curve in black. Tick marks indicate reflection positions, upper line for Cu, lower line for α-F$_2$. Excluded regions in grey. $R_0 = 0.054$, $wR_0 = 0.076$.

<table>
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<tr>
<th>Atom</th>
<th>Wyckoff position</th>
<th>Site symmetry</th>
<th>Site occupancy</th>
<th>x</th>
<th>y</th>
<th>z</th>
<th>$U_{\text{iso}}$</th>
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<tr>
<td>F</td>
<td>8f</td>
<td>1</td>
<td>1</td>
<td>0.27</td>
<td>0.31</td>
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<td>42(1)</td>
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</table>

| F    | 8f               | 1            | 1              | 0.28| 0.31| 0.10| 0 | | |
|      |                  |              |                |   |   |   | 5 | 9 | |

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<tr>
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<th>$U_{22}$</th>
<th>$U_{33}$</th>
<th>$U_{12}$</th>
<th>$U_{13}$</th>
<th>$U_{23}$</th>
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<tbody>
<tr>
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<td>0.018(4)</td>
<td>0.032(3)</td>
<td>-0.002(4)</td>
<td>0.007(2)</td>
<td>0.003(5)</td>
</tr>
</tbody>
</table>

**Figure 2.** The crystal structure of α-F$_2$ (left, projection along a-axis). The F–F bond length is 1.404(12) Å. The different colors in the right picture indicate the hexagonally packed F$_2$ layers to show the relation to cubic closed packed structures (the centers of gravity of the F$_2$ molecules). Red corresponds e.g. to the A layer, green to the B layer, yellow to the C layer. Anisotropic displacement ellipsoids are shown at 70% probability level at 10 K.

Within the layers, the F$_2$ molecules are hexagonally close packed. These layers are parallel to the $ab$-plane. When viewed along the $b$-axis (Error! Reference source not found., right), the ordering of the layers as ABC, and therefore the relationship to the cubic closed packing becomes obvious. Viewed along the $a$-axis, we observe a tilting of the F$_2$ molecules from the normal onto the $ab$-plane, best seen in Error! Reference source not found., left of
17.7(6)°. In α-O₂, this tilting angle is quite small (< 1°).[6] It is reported that these differences can be understood in terms of the larger quadrupole moment of the F₂ molecules and the lack of a magnetic interaction in comparison to O₂ molecules.[6,7] A F₂ molecule is surrounded by twelve F₂ molecules in the shape of a distorted cubeoctahedron, as expected for a structure that derives from the cubic closed packing. Within a layer, the centers of gravity of the F₂ molecules (on Wyckoff position 4d) are 3.1899(5) Å (4 times) and 3.2701(7) Å (2 times) apart, so the hexagon is slightly distorted. To the layer above and below, these distances lie in between 3.9834(8) and 4.0659(10) Å. The shortest intermolecular F···F distances are observed in between the layers with (two times) 2.849(11) and (one time) 2.985(10) Å. In the previous structure model, these F···F distances were reported as 2.82 Å (two times) and 2.87 Å (one time). Within the closed packed layer, the intermolecular F···F distances are (four times) 3.190(11) and (two times) 3.270(9) Å, which are in agreement with previously reported values of 3.20 and 3.28 Å.[4]

The F atoms occupy the 8f/Wyckoff position. We observe the F–F distance as 1.404(12) Å. Previously, it had been fixed at 1.44 Å[1] or refined to 1.49 Å.[4] The F–F distance of the F₂ molecule in the gas phase at 0 K has been calculated using CCSD(T) level of theory with basis sets up to aug-cc-pVTZ to 1.411 Å.[8] From rotational and vibrational Raman spectra 1.4177(15) Å were obtained at room temperature and a pressure of 1 atm,[8] whereas others report a well agreeing value of 1.4168(5) Å.[10] Therefore it seems that the F–F bond length is only little influenced by the state of matter. This has been assumed previously based on IR and Raman data, where the stretch vibration of 898 cm⁻¹ for solid F₂, and of 894 cm⁻¹ for liquid F₂ have been reported.[11] In summary, the F–F bond length observed by us is in excellent agreement with theory as well as spectroscopic experiments.

As can be seen from Error! Reference source not found., the anisotropic displacement ellipsoid of the F atom is elongated in the direction of the F–F bond, which may be due to vibration within the F₂ molecule and also due to librations between the F₂ molecules. The flattening of the anisotropic displacement ellipsoid may also be due to librations of the F₂ molecules.

**Quantum chemical investigation of α-F₂**

The harmonic and anharmonic stretching vibration of gas-phase F₂ have been reported to be 916.929(10) and 893.9416(18) cm⁻¹, respectively.[12] The corresponding frequencies from our *ab initio* CCSD(T)/aug-cc-pVTZ calculation are 916 and 892 cm⁻¹. The vibrational frequencies and the predicted F–F distance of 1.415 Å are therefore in very good agreement with the experiment. From systematic CCSD(T) studies on the F₂ molecule it is known that CCSD(T)/aug-cc-pVTZ benefits from some cancellations of errors in the case of F₂.[4]

Structure optimization of the α-F₂ (C2/c) solid state structure at the DFT-PBE0/TZVP level of theory with D3(B+ABC) dispersion corrections results in a true local minimum and a reasonable agreement with the experimental crystal structure determined by neutron diffraction. The lattice parameters a and b are underestimated by 2.2 and 3.3 %, respectively, while c and β are overestimated by 1.1 and 1.3 %, respectively (see Supporting information for full comparison of structural parameters). We also evaluated the cohesive energy of α-F₂, that is, the energy difference between solid α-F₂ and gas-phase F₂. Using a larger def2-TZVP basis set, the cohesive energy is calculated to be –3.9 kJ/mol (see Supporting Information for details). The absolute value 3.9 kJ/mol can be compared with the experimental sublimation enthalpy of F₂ adjusted to 0 K, which is 8.2 ± 0.3 kJ/mol.[13,14] Our quantum chemically calculated cohesive energy is in the same range as the experimental 0 K sublimation enthalpy, but somewhat underestimated. Müller *et al.* have shown that very good agreement with experiments can be reached using high-level quantum chemical methods.[15]

The harmonic vibrational frequencies of solid α-F₂ could only be calculated with the DFT-PBE0 method, which is not as accurate as CCSD(T) used for the F₂ molecule. In fact, previous benchmarks of several DFT methods have shown that hybrid DFT methods such as DFT-PBE0 overestimate the F–F bond strength and the F–F stretching frequency.[16] This can already be seen in the optimized F–F distance within the α-F₂ structure, which is underestimated by 1.8 % in comparison to the experiment (1.38 Å vs. 1.404 Å). The Raman frequency of the F–F stretching vibration in solid α-F₂ has been reported at 895 cm⁻¹ and 894 cm⁻¹ in previous studies.[11,17] Here the F–F stretching frequency predicted with DFT-PBE0 is 1096 cm⁻¹. Because there are two F₂ molecules in the unit cell, there are in fact two F–F stretching modes: An A₂-symmetric mode where the F₂ molecules are vibrating in the same phase and a B₂-symmetric mode where the F₂ molecules are vibrating in anti-phase. The frequency difference between these two modes is less than 0.5 cm⁻¹ and the intensity of the B₂-mode is only 8% of the A₂-mode. In the experimental Raman spectrum measured with spectral slit width of ca. 0.5 cm⁻¹ the low-intensity B₂-mode could not be distinguished from the A₂-mode.[17]

Even though the F–F stretching frequency calculated on the DFT-PBE0 level of theory is clearly overestimated, the calculated Raman and IR spectra enable comparisons of the low-energy librational modes with the experimental spectra.[17] Four librational modes have been observed in the lattice vibration region of the α-F₂ Raman spectrum: 93, 77, 55, and 44 cm⁻¹. In the calculated spectrum, the corresponding values are 101, 71, 58, and 16 cm⁻¹. The experimental IR spectrum showed librational modes at 80, 42.5, and 28 cm⁻¹. The corresponding DFT-PBE0 modes are 90, 41, and 32 cm⁻¹, with low absorbances of 0.02, 0.03, and 0.01 km mol⁻¹.

Structural optimization of α-F₂ in the alternative space group C2/m discussed above shows it to be essentially identical in energy and C2/m is also a true local minimum. However, the optimization yields much larger deviations of the predicted cell parameters in comparison to the experimentally determined ones. The deviation of the b axis increases from ~3.3 to ~4.6 %, the one of the c axis increases from 1.1 to 1.9 %, and for the angle β from 1.3 to 2.7 %. Only in the case of the lattice parameter a, the deviation...
decreases from $-2.2$ to $-1.0 \%$. The predicted F–F distance however remains exactly the same as for the C2/c structure ($1.38 \text{Å}$).

β-Fluorine
In the previous single crystal X-ray structure determinations of β-F$_2$, fluorine gas was condensed into glass ampoules and flame sealed.\textsuperscript{[2,3]} The authors reported difficulties due to the formation of HF with the H$_2$O residues on the glass. We therefore can conclude that also SiF$_2$ must have been formed during those experiments, as glass decomposition with fluorine is an autocatalytic process. Besides these experimental problems, the reflection intensities had been evaluated visually in those times, which of course can be very difficult. The authors of the previous single crystal structure study tried their very best in using many elegant models to describe the disorder of the F$_2$ molecules, and a section of the crystal structure with their final model is shown in Error! Reference source not found.. The F–F distance was given as 1.418 Å and 1.417 Å. In the previous structure model only a single F molecule was used as the center of gravity of almost spherically disordered F$_2$ molecules on Wyckoff position 2a (m3.), which is at the corners and the center of the unit cell (Error! Reference source not found.). The F atoms on the 12h (mm2.) and 48l (1) positions, which are close to the faces of the unit cell, led only to an unsatisfactory disorder model as seen in Error! Reference source not found.. We investigated some different cubic space groups and employed various disorder models (see Supporting Information), but in the end, we agree with the literature and also describe the crystal structure of β-F$_2$ in the cubic primitive space group Pm3n (No. 223) with the lattice parameter $a = 6.5314(15) \text{Å}$, $V = 278.62(11) \text{Å}^3$, $Z = 8$, cP16, at 48 K. Therefore, β-F$_2$ is isotypic to γ-$\text{Q}_2$ and δ-$\text{N}_2$.\textsuperscript{[3,18,19]} The crystal structure of β-F$_2$ is shown in Error! Reference source not found..

![Figure 3. A section of the old structure model of β-F$_2$. Note the single F atom (a spherically disordered F$_2$ molecule) on the corners and in the center of the unit cell which is drawn with a diameter corresponding to the previously reported F–F atom distance of 1.418 Å. The radii of the other F atoms are chosen arbitrarily. The numbers shown correspond to the z coordinate and give the height of the center of gravities of the F$_2$ molecules in the cell.](image)

We avoided the problems of single crystal growth of F$_2$ and its handling in glass, however we are therefore "only" able to obtain a structure model based on powder and not on single crystal diffraction. As can be seen from the powder neutron diffraction pattern of β-F$_2$ (Error! Reference source not found.), its reflections are of small intensity in comparison to the ones of the Cu sample holder. Additionally, the intensity of the reflections decreases rapidly with increasing scattering angle 2θ, which already indicates that “disorder”, vibrations and librations are present. This assumption is supported by the presence of the diffuse scattering in the powder diffraction pattern. We were therefore unfortunately unable to obtain high-angle diffraction data of sufficient signal to noise ratio for a better refinement.

![Figure 4. Powder neutron diffraction pattern of β-F$_2$ at 48 K. Black dots indicate measured intensity, fitted curve in red (Rietveld refinement for F$_2$, reflections of Cu had to be omitted (grey) from the refinement), difference curve in black. Tick marks indicate reflection positions for β-F$_2$. Rwp = 0.014, wR = 0.020.](image)
Figure 5. A section of the current model of the crystal structure of β-F₂ at 48 K. F atoms are shown isotropic with arbitrary radii. The numbers shown correspond to the z coordinate and give the height of the centers of gravity of the F₂ molecules in the cell.

Table 3. Atom coordinates, Wyckoff positions, site symmetries and occupancies, and isotropic displacement parameters of β-F₂ in comparison to the literature.

<table>
<thead>
<tr>
<th>Atom</th>
<th>Wyckoff position</th>
<th>Site symmetry</th>
<th>Site occupancy</th>
<th>x</th>
<th>y</th>
<th>z</th>
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<tbody>
<tr>
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<td>1/8</td>
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<td>0.06</td>
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<tr>
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<tr>
<td>F2</td>
<td>12/f mm2</td>
<td>1/6</td>
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<td>0.08</td>
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<tr>
<td></td>
<td>2a</td>
<td>m3.</td>
<td>1</td>
<td>0</td>
<td>0.10</td>
<td>0.08</td>
<td>0.08</td>
</tr>
<tr>
<td>F3</td>
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<td></td>
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</table>

Overall, the F1 and F2 atoms arrange around the 2a position in a shape similar to a rhombic dodecahedron. A section of the crystal structure showing the disorder model is available from Error! Reference source not found.. The F1 atoms (yellow in Error! Reference source not found.) form a cube with the F–F bonds of the F₂⁻¹ molecules oriented along the space diagonals of the cube. A distance restraint had to be used for the refinement of the bond lengths preventing them from becoming unphysically short. This leads to (restrained) F1–F1 distances of 1.403(3) Å within the F₂⁻¹ molecule. The F2 atoms form an octahedron (green in Error! Reference source not found.) around the 2a position with an F2–F2 distance of 1.404(6) Å for the F₂² molecules. As in total only a single F₂ molecule can occupy the space around the 2a position, the site occupation factors given in Error! Reference source not found. result. Unfortunately, we could not freely refine the occupancy ratio of the F₂⁻¹ / F₂² molecules but had to fix it at 50 %. Overall, the disorder of the F₂⁻¹ and F₂² molecules is reminiscent of a spherical disorder around the 2a position.

Figure 6. A section of the crystal structure of β-F₂ showing the used disorder model around the 2a position. The F1 atoms forming the cube are drawn yellow, the F2 atoms forming the octahedron are drawn in green. Isotropic displacement parameters are shown with arbitrary radii.

The disorder of the F₂³ molecule with its center of gravity on Wyckoff position 12h (mm2..) can be described using only one F atom (F3). The (restrained) F²⁻³ distances are 1.389(13) Å. In contrast to the previous structure model the refinement of the disordered F₂³ molecules leads to a quite flat arrangement with all F3 atoms almost coplanar (Error! Reference source not found.). We also attempted the refinement with other positions for the F3 atom, however that always led to an insufficient structure model (see the Supporting Information for details).
The structure model was significantly improved, allowed for the application of arbitrary radii.

Quantum chemical calculations on β-F₂
As quantum chemical calculations on the disordered β-modification are enormously demanding and highly difficult to carry out and other groups are successfully working since several years on this task, we decided not to investigate this topic.

Conclusions
Using neutron diffraction and quantum chemical solid-state calculations we have shown α-F₂ to unambiguously crystallize in space group C2/m, mS8, with four F₂ molecules in the unit cell. The lattice parameters obtained at 10 K are a = 5.4780(12), b = 3.2701(7), c = 7.2651(17) Å, β = 102.088(18)°, V = 127.26(5) Å³. The structure model was significantly improved, allowing for the anisotropic refinement of the F atom, and an F–F bond length of 1.40(4) Å was obtained. β-F₂ crystallizes in space group Pm3̅n with the lattice parameter a = 6.5314(15) Å, V = 278.62(11) Å³, Z = 8, cP16, at 48 K. The centers of gravity of the F₂ molecules are arranged like the atoms of the Cr₃Si structure type. β-F₂ is isotypic to γ-O₂ and δ-N₂. Due to the heavy disorder of the structure, only restrained F–F atom distances of 1.403(3), 1.404(6), and 1.389(13) Å could be refined, which still present a significant enhancement in comparison to the literature data.

Table 5. Crystallographic details of α- and β-fluorine and a comparison to the literature data.

<table>
<thead>
<tr>
<th>Crystal system</th>
<th>a/Å</th>
<th>c/Å</th>
<th>β/°</th>
<th>α-F₂ (this work)</th>
<th>α-F₂ (this work)</th>
<th>α-F₂ (this work)</th>
<th>β-F₂ (this work)</th>
<th>β-F₂ (cP16 single crystal)</th>
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<td>127.26</td>
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<td>C2/m (15)</td>
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<td>Pm3̅n (223)</td>
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<td>Pm3̅n (223)</td>
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</table>

Experimental Section
General: All operations with fluorine were carried out in a stainless steel, perfluoroelastomer (FEP) or Monel line. As an inert atmosphere either dry and purified argon (5.0, Westfalen AG, Germany) or helium (5.0, Westfalen AG, Germany) were used so that a possible contact of the inner surfaces of the apparatus and the fluorine with moisture or air was excluded. As vacuum pumps either “fluorine-resistant” two-stage rotary vane pumps (pₘₜₐₓ = 10⁻³ mbar) or turbomolecular pumps (pₘₜₐₓ = 10⁻⁸ mbar) were used. Fluorine was kindly donated by Solvay (> 99.9 %). 5.33 g of fluorine were condensed from a prepassivated nickel bottle into the prepassivated...
copper sample holder which was kept at 50 K so that all fluorine was condensed as a liquid.

Copper was selected as the sample holder as it is the metal of choice in the work with F₂ if passivated properly. Nickel or Monel (a Ni-Cu-alloy) would be superior (also for working the metals), however Ni scatters and absorbs neutrons to a significantly higher extent compared to Cu. We did not dare to use the usual sample holder out of vanadium for neutron diffraction as it is not or not easily passivated, and explosions at the nuclear reactor providing the neutrons should be avoided. In addition to the sample holder, the capillary inside the cryostat stick, allowing for the transfer of F₂, was built out of stainless steel and was thoroughly passivated with F₂ in order to exclude any reactions.

Powder Neutron Diffraction: The powder patterns of F₂ were recorded in a pre-passivated copper ampoule of 12 mm outer and 10 mm inner diameter and of approximately 60 mm height at temperatures of 10 K, 46 K (both for α-F₂) and 48 K (and β-F₂) using the SPODI neutron powder diffractometer (λ = 1.5482 Å) at the research reactor FRM II. After condensation of the fluorine it was cooled below the melting point. A first quick measurement was carried out at 50 K and the positions of the reflections confirmed the presence of the cubic β-phase of F₂. Due to the Cu sample holder, its reflections showed an inhomogeneous intensity distribution along their Debye-Scherrer rings which was due to texture of the Cu sample holder. The sample was then cooled quickly to 10 K. A quick measurement at this temperature showed that the cubic β-phase was present no longer.

Refinement of α-F₂:

The structure solution and refinement of the crystal structure of α-F₂ were carried out using the Jana2006 software [22,23] and the SUPERFLIP algorithm. For the Cu sample holder a Le Bail refinement was used whereas the reflections of α-F₂ were treated with a Rietveld refinement. A manual background was chosen. The pseudo-Voigt functions were used to treat the peak shapes. The slight asymmetry was described by the divergence algorithm implemented in the Jana2006. https://www.ccdc.cam.ac.uk/ CSD 1874484.

Refinement of β-F₂:

The refinement of the crystal structure of β-F₂ was carried out similar to the procedure described above with the exception that the reflections of the Cu sample holder had to be excluded from the refinement. Several disorder models of the F₂ molecules were tested (see Supporting Information details) https://www.ccdc.cam.ac.uk/ CSD 1874485.

Computational details: The quantum chemical calculations on solid alpha-F₂ were done with the CRYSTAL17 program package. We applied the PBE0 hybrid density functional method and the weak intermolecular interactions were treated with the D3 dispersion correction of Grimme (Becke-Johnson damping and three-body correction). A polarized triple-ζ-valence basis set based on the molecular Karlsruhe def2-TZVP basis set was used for the fluorine atoms. The full def2-TZVP basis set was used for single-point calculations on the cohesive energy (see Supporting Information). The reciprocal space of the primitive cell was sampled using an 8x8x4 Monkhorst-Pack-type k-point grid. For the evaluation of the Coulomb and exchange-integrals (TOLINTEG), tight tolerance factors of 8, 8, 8, 8, 16 were used. Both the atomic positions and lattice-constants were fully optimized within the constraints imposed by the space group symmetry. Very tight optimization criteria were applied in the structural optimization: root-mean-square (RMS) of gradient 0.00003 a.u. (TOLDEG), RMS of estimated displacements 0.00012 a.u. (TOLDEX), and the maximum energy change between optimization steps 10⁻¹⁰ a.u. (TOLDEE). Default extra-large integration grid was used for the DFT exchange-correlation functional (XGRID). The harmonic vibrational frequencies were obtained by using the computational scheme implemented in CRYSTAL [34,35]. The CFOUR program package was used to carry out ab initio CCSD(T) coupled-cluster calculations on the F₂ molecule with an augmented and polarized correlation-consistent triple-zeta-valence quality aug-cc-pVTZ basis set [36–41]. The structure of F₂ was fully optimized and both the harmonic and anharmonic vibrational frequencies were evaluated at the CCSD(T) aug-cc-pVTZ level of theory. [42]

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Keywords: fluorine • crystal structure • neutron diffraction • quantum chemical calculations

The crystal structures of α-F₂ and β-F₂ have been reinvestigated using neutron powder diffraction.

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