MoF5 revisited. A comprehensive study of MoF5

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MoFs revisited. A comprehensive study of MoFs.

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Dedicated to Professor Erhard Kemnitz to further acknowledge his ACS Award for Creative Work in Fluorine Chemistry

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Abstract: While the properties of molybdenum pentafluoride, MoF$_5$, have been investigated in the past, there exists no comprehensive study of the compound. Additionally, many of these studies appear incoherent and offer contradictory explanations of some of the observed properties of MoF$_5$. Consequently, a comprehensive examination of MoF$_5$ is presented here, including a redetermination of the crystal structure of MoF$_5$ using single crystal and powder X-ray diffraction, the reevaluation of its IR, Raman and UV-vis spectrum, and a study of its density (3.50(2) g/cm$^3$ @ 25 °C) and magnetic properties. Additionally, density functional theory (DFT) calculations were performed on the gas phase molecule MoF$_{20}$ to provide a discussion of properties realized during investigation. Single crystal X-ray diffraction showed MoF$_5$ to crystalize in the monoclinic, C$_{2}$/m space group, as isolated tetramers having the formula Mo$_4$F$_{20}$. Magnetic measurements showed that when “MoF$_5$” is cooled from the melt fast enough, paramagnetic species with $S = \frac{1}{2}$ are present together with $S = 0$ species. These species may be described using the formula (MoF$_5$)$_n$ ($n = \text{odd}$) and (MoF$_5$)$_n$ ($n = \text{even}$, presumably $n = 4$). From the measurements, the content of the $S = \frac{1}{2}$ species is estimated to be 6 %. The preferred species of MoF$_5$ under ambient conditions is MoF$_{20}$.

Keywords: crystal structure; IR spectroscopy; UV-vis spectroscopy; Raman spectroscopy; density; magnetic properties; molybdenum; fluoride; MoF$_5$ type; WOF$_4$ type

1. Introduction

The synthesis of molybdenum pentafluoride was first described by R. D. Peacock in 1957 as a result of passing dilute fluorine gas over molybdenum hexacarbonyl at −75 °C to produce Mo$_2$F$_9$, which decomposes at 170 °C under vacuum to give MoF$_5$ and MoF$_4$[1]. However, due to the use of fluorine gas, this reaction was described by preceding authors as quite exothermic. Since, simpler techniques have been discovered to produce MoF$_5$, such as the method used in this work which was briefly described by Geichman and coworkers [2] and which will be elaborated upon here. In short, MoF$_5$ is reduced using carbon monoxide and ultraviolet radiation. After the volatile products are discarded, pure MoF$_5$ can be collected from the reaction vessel.

MoF$_5$ is a sunflower yellow powder which, at high purities, melts at 45.7 °C (318.9 K) [3] and reacts with moisture to give a blue hydrolysis product, likely belonging to the molybdenum blues class of compounds. However, other sources report the melting point of MoF$_5$ to be between 63 and 67 °C (336 – 340 K) [1,4,5] but our observations are in agreement with the lower reported melting point.

MoF$_5$ is reported to be soluble in anhydrous HF without decomposition [5], however, through this work it has been found that, while MoF$_5$ is stable in anhydrous HF, it has limited solubility in
the solvent at room temperature. It sublimes at 50 °C [5] and disproportionates at 165 °C, giving MoF\textsubscript{5} and MoF\textsubscript{6}[4], however, no atmospheric or pressure conditions were reported for these measurements. Additional properties of MoF\textsubscript{5} have been reported, albeit not a lot of work has been done on MoF\textsubscript{5} since the 1970’s. UV-Vis [6], IR [7,8], and Raman data [7-9] are available for MoF\textsubscript{5}, however, little to no explanation of the bands arising from these spectra is given in these works. The magnetic properties and electronic properties [10-12], vapor pressure curve [4], and other thermodynamic properties [3] of MoF\textsubscript{5} have also been studied.

Edwards and coworkers investigated the crystal structure of MoF\textsubscript{5} in 1962 [13]. However, they state “the intensities of the spots [collected from single crystal oscillation and Weissenberg photographs] were difficult to estimate” and that the observed differences of Mo1—\(F_{\text{terminal}}\) and Mo2—\(F_{\text{terminal}}\) bond lengths are ”probably not significant”. In our efforts to explore the chemistry of MoF\textsubscript{5} and MoF\textsubscript{6} in greater detail, we obtained single crystals of the latter and re-determined its crystal structure to a much higher accuracy. We also set out to describe the properties of the absorption spectra of MoF\textsubscript{5}, and elaborate upon the magnetic measurements already described in the literature. Additionally, the density of MoF\textsubscript{5} was investigated and is reported here.

2. Results and Discussion

2.1 Single-crystal and powder X-ray structure analysis

MoF\textsubscript{5} crystallizes as fluorine-bridged tetramers having the formula Mo\textsubscript{5}F\textsubscript{20} (M = 763.76 g/mol). Single crystal X-ray diffraction showed the crystal structure of MoF\textsubscript{5} as belonging to the monoclinic space group C2/m (no. 12). The lattice parameters of MoF\textsubscript{5} were calculated by LeBail refinement from a powder diffraction measurement taken at room temperature: \(a = 9.6502(2)\), \(b = 14.2451(2)\), \(c = 5.3100(1)\) Å, \(\beta = 93.088(1)°\), \(V = 728.89(2)\) Å\(^3\). These lattice parameters are in good agreement with those obtained by Edwards and coworkers (\(a = 9.61(1)\), \(b = 14.22(2)\), \(c = 5.16(1)\) Å, \(\beta = 94.21(20)°\), \(V = 703\) Å\(^3\)) [13], however, no measurement temperature was reported for their data collection. The powder diffraction pattern of MoF\textsubscript{5} in Figure 1 shows the synthesized compound to be phase pure; the crystal parameters for MoF\textsubscript{5} are listed in Table 1.

MoF\textsubscript{5} crystallizes in the MoF\textsubscript{5} structure type (mS48, C2/m) which is sometimes referred to as the WOF\textsubscript{5} structure type [14] and is characterized by MoF\textsubscript{5}-units forming tetramers having the Niggli formula \(\text{[MoF}_{2/2}F_{4/1}]\). The fluorine atoms in the crystal structure of MoF\textsubscript{5} pack in a distorted cubic closed packed array. An identical structural motif is observed in the compounds M\textsubscript{2}F\textsubscript{20} (M = Nb, Ta), whereas the MoF\textsubscript{5} structure found in the crystals of RuF\textsubscript{5}, OsF\textsubscript{5}, RhF\textsubscript{5} and PtF\textsubscript{5} (RuF\textsubscript{5} type, mP48, P2\(_1\)/c) is characterized by corrugated tetramers and fluorine atoms which pack in distorted hexagonal closed packed arrays. Moreover, the atomic distances and angles of MoF\textsubscript{5} are comparable to those of NbF\textsubscript{5} and TaF\textsubscript{5}.

The crystal structure of MoF\textsubscript{5} can be seen in Figure 2 and the tetramer MoF\textsubscript{20} motif formed by MoF\textsubscript{5} is presented in Figure 3. In the specific case of MoF\textsubscript{20}, there are two distinct types of Mo atoms, one residing on the Wyckoff position 4\(i\) (site symmetry 2; Mo1) and the other residing on 4\(j\) (site symmetry \(m\); Mo2). In both cases, the Mo atoms experience octahedron-like coordination by six fluorine atoms, in which two of these fluorine atoms are bridging to other Mo atoms. The atomic distance between Mo1 to the \(\mu\)-bridging fluorine atoms F\textsubscript{4}, which reside on the 8\(j\) (1) positions, are observed to be 2.0423(11) Å, whereas the Mo2—\(\mu\)-F distance is 2.0463(11) Å. Both agree well within the 3σ criterion. The \(\mu\)-F—Mo—\(F_{\text{trans}}\) angles are essentially equal with a value of 178.04(6)° for Mo1 and 178.15(6)° for Mo2. Edwards and coworkers reported a Mo1—F\textsubscript{4} distance of 2.04(4) Å and a slightly longer Mo2—F\textsubscript{4} distance of 2.09(4) Å [13], both of which agree with our findings. However, their reported \(\mu\)-F—Mo1—\(F_{\text{trans}}\) and \(\mu\)-F—Mo2—\(F_{\text{trans}}\) angles are both 180.0(2)°.
The atomic distances of Mo1—F5, Mo2—F1 and Mo2—F2, which are the distances to the so-called axial F-atoms, are 1.8012(13), 1.8013(18) and 1.8011(18) Å, respectively, and are thus essentially identical. The distances of Mo1—F6 and Mo2—F3, where the respective F atoms are in trans position to the µ-F atoms and thus will be called equatorial F atoms, are slightly elongated with distances of 1.8236(14) and 1.8183(13) Å, respectively. Thus, the equatorial F atoms exhibit larger distances to the Mo atoms by about 0.02 Å in comparison to the axial F atoms (F1, F2, and F5). It is interesting to note that this finding may be a representation of the structural trans effect [15,16]. However, to the best of our knowledge this has not been reported for (pseudo)octahedral, homoleptic complexes. In NbF5 and TaF5 the distances of the metal atoms to the terminal F atoms are very similar, or the same, if the 3σ criterion is applied and no clear difference in length of the equatorial or axial bonds is observed. Thus, this structural trans effect may arise from the d1 electron configuration of the Mo atoms and will be elaborated upon in the Computational Results section.

Interestingly, the Mo1—F6 bond length reported by Edwards and coworkers (1.89(4) Å) [13], is rather unprecise than that reported here (1.8236(14) Å). It is unclear why Edwards and coworkers observed such an elongation for the equatorial F atoms on the Mo1 atom but this may be due to the problems they described for the estimation of spot intensities. In contrast to this elongation, their reported Mo2—F3 bond length is significantly shorter with a distance of 1.74(4) Å. For the axial F atoms (F5 on Mo1 and F1 and F2 on Mo2), atomic distances of 1.82(9), 1.66(1), and 1.69(1) Å were reported, respectively [13]. In our case, 1.8012(13), 1.8013(18) and 1.8011(18) Å are observed for these respective atomic distances.

As may be expected, the Mo—F atomic distances for the terminal F-atoms are smaller in comparison to those of the bridging fluorine atoms (2.0423(11) and 2.0463(11) Å) by about 0.22 Å for those in equatorial position and 0.24 Å for the axial ones.
<table>
<thead>
<tr>
<th></th>
<th>MoF₅ (SCXRD)</th>
<th>MoF₅ (PXRD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>empirical formula</td>
<td>MoF₅</td>
<td>MoF₅</td>
</tr>
<tr>
<td>color and appearance</td>
<td>transparent yellow cuboid</td>
<td>yellow powder</td>
</tr>
<tr>
<td>molecular mass [g/mol]</td>
<td>190.94</td>
<td>190.94</td>
</tr>
<tr>
<td>crystal system</td>
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<td>monoclinic</td>
</tr>
<tr>
<td>space group</td>
<td>C2/m (12)</td>
<td>C2/m (12)</td>
</tr>
<tr>
<td>a [Å]</td>
<td>9.4719(7)</td>
<td>9.6502(2)</td>
</tr>
<tr>
<td>b [Å]</td>
<td>14.1200(7)</td>
<td>14.2451(2)</td>
</tr>
<tr>
<td>c [Å]</td>
<td>5.0856(4)</td>
<td>5.3100(1)</td>
</tr>
<tr>
<td>β [°]</td>
<td>96.191(6)</td>
<td>93.088(1)</td>
</tr>
<tr>
<td>V [Å³]</td>
<td>676.20(8)</td>
<td>728.89(2)</td>
</tr>
<tr>
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<td>8</td>
<td>8</td>
</tr>
<tr>
<td>ρ&lt;sub&gt;calc&lt;/sub&gt; [g cm⁻³]</td>
<td>3.751 @ 100 K</td>
<td>3.4789 @ 293 K</td>
</tr>
<tr>
<td>ρ&lt;sub&gt;exp&lt;/sub&gt; [g cm⁻³] @ 25 °C</td>
<td>—</td>
<td>3.50(2)</td>
</tr>
<tr>
<td>λ [Å]</td>
<td>0.71073 (MoKα)</td>
<td>1.54051 (CuKα)</td>
</tr>
<tr>
<td>T [K]</td>
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<td>293</td>
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<tr>
<td>μ [mm⁻¹]</td>
<td>3.854 (MoKα)</td>
<td>30.368 (CuKα)</td>
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<tr>
<td>θ&lt;sub&gt;max&lt;/sub&gt;</td>
<td>34.856</td>
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<td>2θ range measured (min, max, increment)</td>
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<td>5.007, 75.972, 0.015</td>
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<tr>
<td>2θ range refined (min, max)</td>
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<td>5.007, 75.972</td>
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<tr>
<td>hkl max</td>
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<td>—</td>
</tr>
<tr>
<td>Size [mm³]</td>
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</tr>
<tr>
<td>R&lt;sub&gt;ave&lt;/sub&gt;, R&lt;sub&gt;r&lt;/sub&gt;</td>
<td>0.047</td>
<td>—</td>
</tr>
<tr>
<td>R (F) (I ≥ 2σ(I), all data)</td>
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<td>—</td>
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<tr>
<td>wR&lt;sub&gt;ave&lt;/sub&gt; (F²) (I ≥ 2σ(I), all data)</td>
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<td>—</td>
</tr>
<tr>
<td>Rp, wRp</td>
<td>—</td>
<td>0.0437, 0.0610</td>
</tr>
<tr>
<td>S (all data)</td>
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<td>4732, 23, 0</td>
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<tr>
<td>Δρ&lt;sub&gt;max&lt;/sub&gt;, Δρ&lt;sub&gt;min&lt;/sub&gt; [e Å⁻³]</td>
<td>1.08, −1.54</td>
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</tr>
</tbody>
</table>
Figure 2. A section of the crystal structure of MoF₅ viewed along the c-axis. Displacement ellipsoids shown at the 70% probability level at 100 K.

Figure 3. Structure of the MoF₅ tetramer unit displaying the MoF₄ units bridged through two fluorine atoms to two neighboring Mo atoms. Atom labeling in accordance to the manuscript by Edwards and coworkers [13]. Displacement ellipsoids are shown at the 70% probability level at 100 K. [Symmetry codes: (i) x, −y, z; (ii) −x, y, 1 − z; (iii) −x, −y, 1 − z.]

The point symmetry of the MoF₂₀ molecule is 2/m, C₂ᵥ, with the twofold rotation axis running through the Mo1 atoms and the perpendicular mirror plane bisecting the Mo2, F1 and F2 atoms. The Mo−Mo distance is 4.0881(3) Å, and the four membered ring formed by the Mo atoms is a flat square.

2.2. Electronic and Vibrational Spectroscopy

The electronic spectrum of MoF₅ dissolved in perfluoroether (C₈F₁₆O, FC-75, 3M) was studied in the range of 190 to 1100 nm (see Figure 4). The spectrum obtained was rather simple with absorption occurring in the near UV and blue regions (beginning at about 500 nm), and the
beginning of another absorption band starting at about 770 nm and extending into the near IR region of the spectrum. Peakcock and Sleight studied the electronic spectrum of molten MoFs in the region of 4000 to 26000 cm⁻¹ (2500 to 385 nm) [6]. They observed a single band in the spectrum centered at 7500 cm⁻¹ (1333 nm). The single absorption band found in the work of Peacock and Sleight may arise from the same electronic transition giving rise to the absorption in the red and near IR regions in this work, however, a direct comparison cannot be made since both works obtained spectra using different sample conditions.

![Graph showing absorption vs. wavelength](image)

**Figure 4.** Visible and near IR spectrum of MoFs dissolved in perfluoroether.

IR spectroscopy was studied on a polycrystalline sample of MoFs at 25 °C in the region of 4000 to 400 cm⁻¹. The IR spectrum obtained from this work shows five bands at approximately 762, 739, 720, 675 and 496 cm⁻¹. It can be said that the bands at 762, 739, 720 and 675 cm⁻¹ are all overlapping to some degree. Upon comparison of the measured IR spectrum to that of a IR spectrum calculated for the solid state of MoFs using the DFT-PBE0 method (a more detailed description of these calculations can be found in the Computational Results section), it can be seen that the peak positions correlate very well, however, the peak shapes and intensities vary. A comparison of the experimental and theoretical spectra is shown in Figure 5. Furthermore, peak assignments for the bands in the experimental IR spectrum were made utilizing the results of the theoretically calculated IR spectrum in which the theoretical vibrations were visualized using the Jmol software [17]. Band assignments are listed in Table 2.
Figure 5. Top: Experimentally obtained IR spectrum of crystalline MoF$_5$; bottom: Theoretically calculated IR spectrum of the solid state of MoF$_5$ calculated using the DFT-PBE0 method.

As mentioned earlier, IR spectroscopy of MoF$_5$ has been previously studied. Acquista and Abramowitz studied differences in the IR spectra of MoF$_5$ vapors in argon matrices at liquid helium temperatures [7]. These differences occurred by allowing the MoF$_5$ vapors to reach equilibrium at different temperatures in the range of 25 – 150 °C before being trapped at liquid helium temperatures for measurement. They observed bands at 768, 716, 704 and 231 cm$^{-1}$ during the room-temperature measurements. At higher temperatures, though, these bands began to decrease in intensity while bands at 713, 683, 261 and 112 cm$^{-1}$ began to appear. They attributed this observation to the disappearance of polymeric species at higher temperatures and the increase of the monomeric MoF$_5$ unit. However, a comparison of these IR spectra in argon matrices to the polycrystalline spectrum reported here cannot be made due to the extreme differences in sample handling and measurement techniques. Acquista and Abramowitz also studied a solid-state spectrum of MoF$_5$ at liquid helium temperatures, which gave rise to broad bands at 725, 700, 660, and 525 cm$^{-1}$. Due to its absence in the spectra of heated samples described above, they attributed the band at 525 cm$^{-1}$ as arising from the fluorine-bridge bond of the tetrameric unit. This band at 525 cm$^{-1}$ may correspond to the band observed in this work at 496 cm$^{-1}$, which arises from the symmetric and asymmetric stretching of the Mo$\text{--}\mu$-F bonds, in which case the observations of Acquista and Abramowitz may be confirmed. However, the solid-state spectrum of MoF$_5$ received
from Acquista and Abramowitz [7] is completely different from that reported in this work. This difference, however, may arise from different sample handling techniques and temperature of analysis.

Ouellette and coworkers studied the IR spectrum of a super-cooled liquid of MoF$_5$ at 25 °C [8]. This spectrum showed bands at approximately 725, 690, 490, 250, 180 and 130 cm$^{-1}$. Analysis of their liquid MoF$_5$ data was made on the assumption that monomeric MoF$_5$ species (of $D_3h$ symmetry) were present in their melted sample. The liquid IR spectrum obtained from the work of Ouellette and coworkers has similar features when compared to the polycrystalline spectrum obtained in this work, although the peak positions and intensities are different. They also studied the IR spectrum of solid MoF$_5$ which showed weak bands at 970, 890, 845, 520 and 480 cm$^{-1}$, medium bands at 200 and 160 cm$^{-1}$, and very strong bands at 745, 698 and 647 cm$^{-1}$. The authors did not attempt to make band assignments for the spectrum of solid MoF$_5$ due to the complexity of the molecule’s solid state structure, nor do they state the temperature at which the measurement was performed. Nevertheless, similar features can also be seen in their solid MoF$_5$ spectrum when compared to the polycrystalline spectrum reported here. The IR bands obtained from this work and previous works can be compared in Table 2.

Furthermore, an interesting statement was found in the work of Ouellette and coworkers [8] in which they report the unlikelihood that a reversible change between the tetrameric unit in the solid and a polymeric unit (n > 4) in the liquid could be made [8]. This statement was tested in this work through the use of IR spectroscopy to determine if a crystalline sample of MoF$_5$ decomposes after several melting and cooling cycles to a glassy state characterized by an increased presence of monomers, dimers, or trimers. In this experiment, a sample of MoF$_5$ was melted at 90 °C and then immediately cooled in liquid nitrogen to produce glassy MoF$_5$. It was observed that this glassy product quickly reverted back to the crystalline form at room temperature within minutes. Sequential IR spectra taken after nine melting, cooling and annealing (at room temperature) cycles looked identical to the one shown in Figure 5; no new peaks appeared in the range measured. This suggests that any polymeric species present in the melt either recombine or decompose to form the tetramer unit at room temperature. However, our magnetic measurements (see below) showed that cooling rates are of great importance in the conservation of the (MoF$_5$)$_n$ species.

Lastly, a Raman spectrum was obtained for a crystalline sample of MoF$_5$ at room temperature (see Figure 5). This spectrum showed eleven discernible bands at 757, 736, 703, 694, 401, 288, 248, 239, 197, 181 and 134 cm$^{-1}$. It was compared to a theoretically calculated spectrum obtained for the solid state of MoF$_5$ using the DFT-PBE0 level of theory. These spectra can be compared in Figure 6. The band locations correlate very well between the experimental and theoretical spectra, however, the band intensities are different. A peak at 401 cm$^{-1}$ is present in the experimentally obtained Raman spectrum but absent in the theoretically calculated spectrum. This peak may be an overtone of two vibrational modes, or it may arise from a small amount of polymers and/or monomers in the sample, or from an impurity. Band assignments were made using the theoretical Raman spectrum and can been seen in Table 2. Raman spectroscopy of crystalline MoF$_5$ has previously been studied by Acquista and Abramowitz [7], Ouellette and coworkers [8], and Bates [9]. The spectrum obtained from this work agrees very well with the spectra obtained from these previous studies; they can be compared in Table 2.
Table 2. Comparison of IR and Raman data obtained in this work and previously reported works [7–9]. Band assignments are given for the IR and Raman spectrum obtained from this work. Where two or more band assignments are given, coupling of these optical modes were obtained from the quantumchemical calculations. Some vibrational assignments are reported without a “symmetrical” or “asymmetrical” classification due to overlap.

<table>
<thead>
<tr>
<th>This work</th>
<th>Acquista et al.</th>
<th>Acquista et al.</th>
<th>Ouellette et al.</th>
<th>Ouellette et al.</th>
<th>This work</th>
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<tr>
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<td>IR (solid) [cm⁻¹]</td>
<td>IR (argon matrix) [cm⁻¹]</td>
<td>IR (solid) [cm⁻¹]</td>
<td>IR (liquid) [cm⁻¹]</td>
<td>Raman (crystalline) [cm⁻¹]</td>
<td>Raman (crystalline) [cm⁻¹]</td>
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<tr>
<td>762 ν(Mo—F(eq))</td>
<td>725</td>
<td>768</td>
<td>745</td>
<td>725</td>
<td>757 ν(Mo—F(eq)); νas(Mo—F(ax))</td>
<td>758</td>
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<tr>
<td>739 ν(Mo—F(eq)); νas(Mo—F(ax))</td>
<td>700</td>
<td>735</td>
<td>733</td>
<td>736 ν(Mo—F(eq)); νas(Mo—F(ax))</td>
<td>735</td>
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<tr>
<td>720 ν(Mo—F(eq)); ν(Mo—F(eq))</td>
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<td>713</td>
<td>704</td>
<td>703 νas(Mo—F(eq)); νas(Mo—F(ax))</td>
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<tr>
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<td>660</td>
<td>683</td>
<td>698</td>
<td>690</td>
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<td>496 ν(Mo—µ-F); νas(Mo—µ-F)</td>
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</table>

*Indicates two vibrational modes which occur so closely to each other that they could not be resolved into two peaks.
Figure 6. Top: Measured Raman spectrum of crystalline MoF$_5$; bottom: Theoretical Raman spectrum of the solid state of MoF$_5$ calculated using CRYSTAL17 calculations.

2.3. Magnetic Measurements

Magnetic data of MoF$_5$ was obtained with the application of the VMS option of a Quantum Design physical property measurement system (ppms). The material had to be measured in a homemade FEP sample holder due to the enhanced reactivity of MoF$_5$ with moisture residues present in traditional sample holders. The data was corrected with respect to the contribution of the sample holder as well as the diamagnetic contribution of the sample through utilization of both experimental data and Pascal constants. The molar diamagnetic susceptibility of MoF$_5$ was calculated to be $-6.8 \times 10^{-5}$ cm$^3$ mol$^{-1}$. Field and temperature dependent magnetic data was also recorded.

Temperature-dependent scans with applied fields of 1 and 5 Tesla were performed in the temperature range from 1.8 K to 300 K. The curves show a broad maxima at about 260 K. A decrease in temperature is met with decreasing magnetization, however, at temperatures below 50 K, the magnetization increases once again and reaches values close to the room temperature measurements (see Figure 7).
Figure 7. Magnetization curve of “MoFs” at 1 (lower curve) and 5 T (upper curve).

Field-dependent measurements of MoFs were also carried out at 1.8, 5, 10, 15, 50, 100, 150, 200 and 300 K. Up to 10 K the magnetization data show significantly sigmoidal curves. Above 10 K the behavior is linear (Figure 8 and Figure S1).

Figure 8. Field dependent measurements on a crystalline sample of MoFs.

This data describes the magnetic behavior of the crystalline state of MoFs, which is in good agreement with the previously reported [12] antiferromagnetism of MoFs that is responsible for the observed maximum at 260 K. The increase of the magnetization below 50 K is an indicator towards the presence of an additional paramagnetic phase in the sample. This observation indicates the breakdown of the MoF$_{20}$ unit into smaller subunits (i.e. monomers, dimers, and trimers). If the paramagnetic behavior is assumed to stem from species with $S = \frac{1}{2}$, that is, (MoFs)$_n$ $(n = \text{odd})$ subunits, their amounts can be estimated to circa 0.3% for $n = 1$, and circa 1% for $n = 3$. These amounts were determined based on the Brillouin-fit shown in Figure S1. Such amounts are not observable, or would be very difficult to observe, in a powder X-ray diffraction pattern, so it is unclear whether the paramagnetic phase is crystalline or amorphous. However, due to the observed
ability of MoF$_5$ to be readily super-cooled [8] and easily transferred to an amorphous state [12], we
assume it to also be amorphous in our case.

This cooled MoF$_5$ sample was then warmed above its melting point, leading to an increase in the
observed magnetization. The rise in the magnetization curve observed at approximately 321 K is,
however, dependent upon the heating rate, as shown in Figure 9. In all cases, though, the content of
the paramagnetic species with $S = \frac{1}{2}$ seems to have increased abruptly to approximately 6%
(Brillouin-fit in Figure S1), indicating cleavage of the tetramer unit. Upon rapidly cooling the heated
sample (20 K/min, indicated by blue curve in Figure 9) a retention of these paramagnetic species
seems to occur, as indicated by an observed Curie-Weiss-like magnetic behavior. A Curie-Weiss-fit
(Figure S2) yielded a $\theta$ value of $-141.7(2)$ K, indicating the presence of somewhat strong
antiferromagnetic coupling within this phase. If a Curie-Weiss-fit with two phases, one being a
putatively strong antiferromagnetic phase and the other being a paramagnetic phase, is carried out,
then the ratio of these two phases can be determined, leading to the calculation of approximately
6% paramagnetic, $S = \frac{1}{2}$ species present in the sample.

Upon warming of this increased-paramagnetic, metastable phase to 300 K, with a heating rate of
2 K/min, a recombination of the (MoF$_5$)$_n$ species (with $n = \text{odd}$) to (MoF$_5$)$_n$ (with $n = \text{equal}$; most
likely $n = 4$) is likely occurring as indicated by the decrease in susceptibility to the former value of
the crystalline phase (as shown by the violet curve in Figure 9). As we cannot observe the sample
visually during the magnetic measurements, it remains an open question as to whether the increase
in paramagnetic species corresponds to the melting of the compound, or if the tetramers begin to
decompose in the solid state. However, melting of the sample is suspected as the temperature
nicely corresponds to the reported melting point of high purity MoF$_5$ [3].

**Figure 9.** Magnetic susceptibility of two different MoF$_5$ phases (black). The magnetic behavior
during the phase transitions is measured with rising temperatures. The cleavage of the tetramers is
shown by the red and orange curves (two separate measurements), the recombination of (MoF$_5$)$_n$ ($n
= \text{odd}$) to (MoF$_5$)$_n$ ($n = \text{equal}$; very likely $n = 4$) is shown by the violet curve. The rapid cooling of the
heated sample with a cooling rate of 20 K/min, which presumably leads to an amorphous state, is shown by the blue curve.

2.4 Computational Results

Solid MoF₅ (MoF₂₀) was investigated at the DFT-PBE0/TZVP level of theory (see Materials and Methods for full computational details). First, the geometries of two spin configurations were fully optimized, corresponding to the spins of the four Mo atoms in either a ferromagnetic (FM) or antiferromagnetic (AFM) configuration. Both spin configurations could be studied in the original space group C2/m. The AFM configuration turned out to be the magnetic ground state, being 5 kJ/mol lower in energy in comparison to the ferromagnetic configuration. The absolute value of the spin-only magnetic moment of the Mo atoms is 0.94 μB. All solid-state results discussed below have been obtained for the antiferromagnetic ground state.

The lattice parameters of the optimized crystal structure are slightly overestimated in comparison to the SCXRD structure (differences in parentheses): a = 9.74 Å (2.8%), b = 14.39 Å (1.9%), c = 5.21 Å (2.5%) and β = 95.9° (−0.3%). The calculated Mo–F distances showed the same trend as observed in the experimental structure: Mo–F_{bridge} = 2.07 Å, F_{eq} = 1.84 Å, and F_{ax} = 1.82 Å. The IR and Raman spectra calculated for MoF₅ are shown in Figure 5 and Figure 6, respectively. The calculated spectra are in good agreement with the experimental spectrum and enabled the assignment of the vibrational modes.

In addition to solid-state MoF₅, the MoF₂₀ molecule was investigated in the gas-phase (DFT-PBE0/def2-TZVP level of theory). The D₄h-symmetric antiferromagnetic ground state is 7 kJ/mol more stable than the D₄h-symmetric ferromagnetic spin configuration. We investigated the chemical bonding with the help of Intrinsic Atomic Orbital (IAO) population and Intrinsic Bond Orbitals (IBOs) analysis. The partial atomic charges (e⁻) are as follows: Mo = +2.09, F_{bridge} = −0.55, F_{eq} = −0.40, and F_{ax} = −0.37. The larger the partial charge of an F atom, the longer the calculated Mo–F distance: Mo–F_{bridge} = 2.06 Å, F_{eq} = 1.82 Å, and F_{ax} = 1.80 Å. The longer Mo–F_{eq} distance in comparison to Mo–F_{ax} distance is in agreement with the experimental and computational solid-state results. The elongation of the Mo–F_{eq} can be understood by analyzing the Mo–F bonding in terms of IBOs, which are a physically well-defined form of localized molecular orbitals (MOs) and more suitable for chemical interpretation than the delocalized canonical molecular orbitals. The IBO analysis shown in Figure S3 shows that the highest-energy lone pair orbital of the F_{eq} atom is in complete anti-phase to the Mo d²₋ₓ²ᵧ₂ orbital (unpaired Mo d¹ electron). In the case of the F_{ax} atom, the highest energy lone pair orbital and the Mo dₓ²−ᵧ² orbital are not in complete anti-phase and some constructive overlap arises (the IBO has ~10% contribution from Mo). As a consequence, the repulsion between the Mo d¹ electron and the F lone pair electrons is smaller for the F_{ax} atoms than the F_{eq} atoms and the Mo–F_{ax} distance is slightly shorter than the Mo–F_{eq} distance.

The energetics of the following (MoF₃)ₙ species were also studied (molecular geometries are included in the Supporting Information): MoF₃ (C₁); MoF₁₀ (D₂₀); MoF₁₅ (D₃₀); MoF₂₀ (D₄₀); MoF₂₅ (D₅₀). These were gas-phase calculations at 0 K and a ferromagnetic spin configuration was used for all systems since the aim of the study was to understand the structural strain in (MoF₃)ₙ species and not to investigate the much smaller FM/AFM energy differences. This study gave the relative energies (kJ/mol) per MoF₅ unit, which for n = 1–5 are as follows: 67, 85, 41, 0, and 40. Thus, the concatenation of MoF₅ units is energetically favorable and the MoF₂₀ species is clearly the (MoF₃)ₙ species with the smallest structural strain.

3. Conclusions

MoF₅ was synthesized by the reduction of MoF₅ in the presence of CO under UV irradiation. It was confirmed that MoF₅ crystallizes in the monoclinic C2/m space group as MoF₂₀ tetramers.

Single crystal analysis showed a discernable difference in atomic distance between the Mo–F_{eq} and...
the Mo—F$_{eq}$ bonds, with the Mo—F$_{eq}$ atomic distances being longer by about 0.02 Å. Theoretical calculations of the MoF$_{20}$ molecule in the gas-phase (run at the DFT-PBE0/def2-TZVP level of theory) were analyzed using IAO and IBOs. The IBO analysis showed the highest-energy lone pair orbital of the F$_{eq}$ atoms to be in complete anti-phase with the Mo d$^2$F$^2$ orbital, whereas the highest energy lone pair orbital of the F$_{ax}$ atoms are not in complete anti-phase with the Mo d$^2$F$^2$ orbital and instead exhibit some constructive overlap. This constructive overlap between orbitals in the Mo—F$_{ax}$ bond is what leads to the slightly shorter bond length of this bond when compared to the Mo—F$_{eq}$ bond.

Furthermore, the vibrational spectroscopy study of MoF$_5$ showed that great differences in the IR spectrum of the molecule can be observed depending on the sample handling and measurement techniques. Polycrystalline IR and Raman spectra are reported here, in which theoretical calculations of solid-state MoF$_5$ were used to make band assignments. It was shown that of the 69 possible optical modes of MoF$_{20}$, 29 optical modes (18 Raman; 11 IR) could be seen in the computational spectra and 16 (11 Raman; 5 IR) could be seen in the experimental spectra. Band assignments for the experimental spectra can be seen in Table 2.

The study of the magnetic properties of a polycrystalline sample of MoF$_5$ showed the compound to exhibit antiferromagnetic properties, with a maximum observed at 260 K. An increase in magnetism was also observed at temperatures below 50 K. This increase in magnetism is thought to occur from paramagnetic (MoF$_5$)$_n$ ($n = odd$) subunits with $S = \frac{1}{2}$. Warming of the sample lead to an increase of these $S = \frac{1}{2}$ subunits, most likely belonging to the monomers and trimers of cleaved MoF$_5$ units. Upon rapid cooling of this sample with increased-paramagnetic behavior, it was shown that this increase in paramagnetism could be preserved. However, upon heating of this increase-paramagnetic, metastable phase back to room temperature, a decrease in magnetism was observed, leading to the conclusion that the paramagnetic, (MoF$_5$)$_n$ ($n = odd$), species recombined to give the MoF$_{20}$ tetramers. In support of this conclusion, it was shown through our computational study that the MoF$_{20}$ unit is the preferred species of “MoF$_5$”. It seems a fair assessment to conclude from our investigations that, while a small percentage of MoF$_5$ may be found in the solid state as (MoF$_5$)$_n$ ($n = 1-3$) units, the preferred species of MoF$_5$ under ambient conditions is MoF$_{20}$.

4. Materials and Methods

4.1. General Procedures and Materials

All operations were performed in either stainless steel (316L) or Monel metal Schlenk lines, which were passivated with 100% fluorine at various pressures before use. Preparations were carried out in an atmosphere of dry and purified Argon (5.0, Praxair). Carbon monoxide (5.0, Air Liquide Deutschland) was used as supplied. Molybdenum hexafluoride (99%, ABCR) was distilled once prior to usage. Several UV light bulbs (254 nm, HNS S 11 W G23, Osram Puritec) were used in a homemade chamber to promote MoF$_6$ reduction.

4.2. Synthesis of MoF$_5$

In order to synthesize molybdenum pentafluoride, MoF$_6$ (3.74 g, 17.80 mmol) was distilled into a 300 mL quartz vessel that was previously evacuated and flamed-dried three times. To this vessel 800 mbar (9.69 mmol) of carbon monoxide was added and the reaction vessel was placed into a chamber to allow irradiation by ultraviolet light (254 nm). After 12 hours of irradiation, the product was cooled using liquid nitrogen, the gaseous species in the vessel were evacuated, 800 mbar of fresh carbon monoxide was added, and the vessel was irradiated with UV light for an additional 12 hours. After the reaction was complete, all remaining volatile species were evacuated at room temperature and the product (a yellow powder) was transferred into a glovebox for further sample handling. The yield of MoF$_5$ was quantitative, except for mechanical losses. A powder X-ray
diffraction pattern was taken directly from the product with no further sample preparation and confirmed that phase-pure MoF₅ was obtained. The product was found to decompose in air to form a blue hydrolysis product, as expected.

In an inert atmosphere, about 200 mg of MoF₅ was transferred to an FEP (fluorinated ethylene propylene) tube and sealed. This tube was placed in an oil bath set to 90 °C in order to melt the sample. Once melted, the MoF₅ was allowed to slowly reach room temperature. This method was used to produce the single crystals used in this study. All other measurements of MoF₅ were performed directly on the powder obtained from the reaction.

4.3. Single Crystal X-ray Diffraction

X-ray structure analysis of the single crystals of MoF₅ was carried out with a STOE IPDS 2T diffractometer with plane graphite-monochromated molybdenum radiation (Mo-Kα, λ = 0.71073 Å) generated by a sealed X-ray tube (12×0.4 mm long fine focus), and a detector resolution of 6.67 pixels mm⁻¹. Evaluation and integration of the diffraction data was carried out using the X-Area software, and an absorption correction was made through integration using the X-Red³ and X-Shape program within the parent software [18]. The structure was solved using the previous structure model of MoF₅ and refined against F² in the SHELXL software [19,20]. All atoms were located by Difference Fourier synthesis and refined anisotropically. Representations of the crystal structure were created using the Diamond software [21]. Further details of the crystal structure investigation may be obtained from the Fachinformationszentrum Karlsruhe, 76344 Eggenstein-Leopoldshafen, Germany (Fax: +49-7247-808-666; E-Mail: crysdata@fiz-karlsruhe.de, http://www.fiz-karlsruhe.de/request for deposited data.html) on quoting the depository numbers CSD-433190.

4.4. Powder X-ray Diffraction

Powder X-ray diffraction patterns were obtained with a Stadi-MP-Diffractometer (STOE) using Cu-Kα radiation (λ = 1.54051 Å), a germanium monochromator, and a Mythen1K detector. The data were handled using the WINXPow software [22]. The compound was filled into borosilicate capillaries, which were previously flamed dried under vacuum, and sealed using a hot tungsten wire. The powder diffraction pattern presented in this work was obtained using a ring collimator, a step size of 0.195° 2θ, and a measurement time of 125 seconds per step between 0 and 78° 2θ. Using the Jana2006 software [23], a Le Bail refinement was used to fit the powder diffraction pattern. A Chebyshev polynomial employing 15 terms was used for background correction, however, 3 of these 15 terms were omitted from refinement. In order to fit peak shape, a Pseudo-Voigt function was employed. Asymmetry was corrected through divergence.

4.5. UV/VIS Spectroscopy

The electronic spectrum was measured on a SPECORD 210 PLUS UV/Vis spectrophotometer. The sample was dissolved in perfluoroether and placed into a homemade sample holder with sapphire windows. The spectrum was processed with the WinASPECT Plus software [24].

4.6. IR Spectroscopy

The IR spectrum was measured on an alpha FTIR spectrometer (Bruker) using a diamond ATR unit under an Ar atmosphere. The spectrum was processed with the OPUS software package [25].

4.7. Raman Spectroscopy

MoF₅ was loaded into a 0.3 mm borosilicate capillary and the Raman spectrum was measured in backscattering geometry by means of a Raman microscope inVia (Renishaw), using a frequency-
doubled Nd:YAG laser (532 nm wavelength). The spectrum was recorded in confocal mode between 2 cm\(^{-1}\) and 1792 cm\(^{-1}\). The laser power was reduced to 5% to prevent degradation of the sample.

4.8. Density Determination

The density of MoF\(_5\) was measured using the automated gas displacement pycnometry system AccuPyc II 1340 (micromeritics) with a calibrated 0.1 cm\(^3\) sample holder and helium as the displacement gas. The number of preliminary purges was set to 30, while the subsequent density measurements were performed 50 times with measurement averaging.

4.9. Magnetic Measurements

DC-magnetic data were recorded with the VSM option in a Physical Property Measurement System (PPMS Dynacool, Quantum Design, SanDiego, USA). Temperature and field dependent scans were performed in the range from 1.8 to 400 K and from −9 to 9 Tesla. The maximum cooling rate of 20 K/min allows to convert the MoF\(_5\) from its liquid to a super-cooled state.

4.10. Computational details

Periodic quantum chemical calculations for MoF\(_5\) were carried out using the PBE0 hybrid density functional method (DFT) [26,27]. A polarized triple-zeta-valence (TZVP) level basis set for all atoms was applied. The basis set for molybdenum was derived for this study from the def2-TZVP basis set (details and full basis set listing in Supporting Information) [28]. The basis set for fluorine was taken from our previous study on sodium hydrogen fluorides [29]. All calculations were carried out using the CRYSTAL17 program package [30]. The reciprocal space was sampled using a 5x5x5 Monkhorst-Pack-type \(k\)-point grid [31]. All calculations were run as spin-unrestricted (the magnetic ground state is described in the main text). For the evaluation of the Coulomb and exchange integrals (TOLINTEG), tight tolerance factors of 8, 8, 8, 8, and 16 were used. Both the atomic positions and lattice constants were fully optimized within the constraints imposed by the space group symmetry. Default optimization convergence thresholds and an extra-large integration grid (XLGRID) were applied in all calculations.

The harmonic vibrational frequencies, IR intensities, and Raman intensities were obtained through the use of the computational scheme implemented in CRYSTAL [32–35]. The Raman intensities have been calculated for a polycrystalline powder sample (total isotropic intensity in arbitrary units). The Raman final spectrum was obtained by using pseudo-Voigt peak profile (50:50 Lorentzian:Gaussian) and FWHM of 8 cm\(^{-1}\). When simulating the Raman spectrum, the temperature and laser wavelength were set to values corresponding to the experimental setup (\(T = 298.15\) K, \(\lambda = 532\) nm). For the IR spectrum, Lorentzian lineshape and FWHM of 8 cm\(^{-1}\) was used. The peak assignment was carried out by visual inspection of the normal modes (Jmol program package [17]).

In addition to the periodic calculations, molecular gas-phase calculations were carried out at the DFT-PBE0/def2-TZVP level of theory using TURBOMOLE program package [28,36,37]. Resolution of Identity (RI) approximation was used to speed up the calculations [38,39]. Intrinsic Atomic Orbitals (IAOs) and Intrinsic Bond Orbitals (IBOs) were used in the bonding analysis of the cations [40].

Appendix A. Supplementary data

Supplementary material related to this article can be found, in the online version, at doi:

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