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$K_3Mo_2O_{5.6}F_{3.4}$ and $K_3V_2O_{3.3}F_{5.7}$ – exploring transition metal cation valence and anion distribution in oxyfluorides†

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Oxyfluorides come in many different structures and are highly adaptable in composition, not least because of their mixed-anionic nature. Slight changes, unless specifically looked for, can easily go unnoticed. In this paper, we present two oxyfluorides, $K_3Mo_2O_{5.6}F_{3.4}$ and $K_3V_2O_{3.3}F_{5.7}$, synthesized under high-pressure/high-temperature conditions, and demonstrate the importance of careful analysis of composition, oxidation state and O/F anion distribution for an accurate description of oxyfluorides. Their crystal structures were determined by single-crystal X-ray diffraction and the transition metal cation valences analyzed by X-ray photoelectron spectroscopy (XPS). The O/F anion ratio was calculated using the principle of charge neutrality and the local distribution within the crystallographic framework was studied using bond valence (BV) and charge distribution (CHARDI) calculations. Madelung Part of Lattice Energy (MAPLE) calculations and magnetic measurements provide insight into phase stability and corroborate the mixed-valent nature of the compounds.

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Introduction

In contrast to their parent compounds, oxides and fluorides, oxyfluorides are a comparatively young and scarcely investigated substance class. Very rare in nature, often unstable and difficult to synthesize, they have been flying under the radar of popular research for a long time. It was only recently that oxyfluorides have started to attract more scientific interest due to their potential use in nonlinear optical materials,¹ batteries,^{2,3} transparent glass ceramics,⁴ solar cells^{5,6} and deep-red emitting Mn⁴⁺-activated phosphor materials,⁷ to name but a few.

What ultimately determines the properties of a new compound and its usefulness in the above applications is its crystal structure. Oxyfluorides are known for their structural diversity, due in no small part to their mixed-anionic nature.^{8,9} Oxygen and fluorine as anions are similar enough to share sites within the same crystallographic framework, yet different

enough to have a substantial impact on the local atomic environments and, as a consequence, the optical and electronic properties of the compound in question. An excellent way of probing the boundaries of what is structurally possible are high-pressure/high-temperature experiments, as they allow us to subject samples to extreme conditions and force them to adopt structures that they would not otherwise adopt.

A handful of oxyfluorides of the general formula $A_3M_2X_9$ (A = alkali metal, M = transition metal and X = O, F) have already been reported in the literature. Most of them crystallize in the structure type $Cs_3Cr_2Cl_9$,¹⁰ likely limited in terms of structural variety by the size of the caesium cation: $Cs_3V_2O_2F_7$,¹¹ $Cs_3V_2O_4F_5$,¹² $Cs_3Mo_2O_6F_3$,¹³ as well as the exchange variant $Cs_3V_{1.5}Mo_{0.5}O_3F_6$.¹² As for potassium variants, three representatives containing only pentavalent transition metal cations have been reported before: $K_3V_2O_4F_5$,^{14–16} $K_3Mo_2O_4F_5$ ¹⁷ and $K_3Nb_2O_4F_5$.¹⁸ The structure of $K_3V_2O_4F_5$ has never been elucidated in detail, and two different datasets are available for $K_3Mo_2O_4F_5$: a powder diffraction file (PDF) entry from 1983 and a complete structure determination from 2012.¹⁹ $K_3Nb_2O_4F_5$ is the only other known $A_3M_2X_9$ representative containing potassium and was first reported by our group early last year.¹⁸ While quite similar to $K_3V_2O_4F_5$ and $K_3Mo_2O_4F_5$ at first glance, the oxyfluorides reported in the present investigation were synthesized under high-pressure/high-temperature conditions and crystallize in a different structure type, namely $Ba_2RbFe_2F_9$.²⁰ Another common point of contention in oxy-

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fluorides is the oxidation state of the transition metal cations. This paper is the first report on $A_3M_2X_9$ type oxyfluorides that includes modern XPS techniques to quantitatively and reliably determine the oxidation states of the transition metal cations and, in turn, O/F ratios. Hereinafter, we explore the complex interplay between these two aspects and investigate their influence on local atomic environments through a detailed study of two new oxyfluorides $K_3Mo_2O_{5.6}F_{3.4}$ and $K_3V_2O_{3.3}F_{5.7}$.

Results and discussion

$K_3Mo_2O_{5.6}F_{3.4}$ and $K_3V_2O_{3.3}F_{5.7}$ were synthesized under high-pressure/high-temperature conditions at 8.0 GPa and 900 °C using a mixture of either KF and MoO_3 or KHF_2 and V_2O_5 . The V variant can also be synthesized utilizing certain blends of V_2O_5 and VO_2 . For a detailed description of the synthetic procedure, refer to the Experimental section. Transition metal cation oxidation states and their respective concentrations were determined by X-ray photoelectron spectroscopy (see section XPS measurements). The O/F ratio was subsequently calculated on the basis of charge neutrality, resulting in the final compositions $K_3Mo_2O_{5.6}F_{3.4}$ and $K_3V_2O_{3.3}F_{5.7}$.

The crystal structures of $K_3V_2O_{3.3}F_{5.7}$ and $K_3Mo_2O_{5.6}F_{3.4}$

$K_3V_2O_{3.3}F_{5.7}$ crystallizes in the rhombohedral space group $R\bar{3}m$ (no. 166) with the cell parameters $a = 5.7442(1)$, $c = 21.0484(6)$ Å, $V = 601.46(2)$ Å³ and $Z = 3$ at $T = 300(2)$ K. Except for the position of the transition metal cations within their octahedral coordination sphere, $K_3Mo_2O_{5.6}F_{3.4}$ crystallizes isotypically to $K_3V_2O_{3.3}F_{5.7}$. In accordance with the size difference between Mo^{6+} and $V^{4+/5+}$, its cell parameters are slightly larger with $a = 5.7983(1)$, $c = 21.1801(5)$ Å and $V = 616.68(2)$ Å³. Details on the single-crystal data and structure refinement process are presented in Table 1. Atomic coordinates, Wyckoff positions as well as equivalent isotropic displacement parameters are included in Table 2 (V) and Table 4 (Mo). Anisotropic displacement parameters can be found in Table 3 (V) and Table 5 (Mo), a comparison of the basic crystal structure parameters and interatomic distances is shown in Table 6.

The crystal structure of the title compounds can be described with cubic close packings of atoms. The anionic network atoms form distorted cubic close packed arrangements together with potassium atoms, while the transition metal cations occupy a total of $1/6^{\text{th}}$ of the octahedral voids. As there is an additional shift after every three layers, the cubic close packing is not continuous. A detailed description of the packing of atoms can be found in our previous publication for $K_3Nb_2O_4F_5$, which also crystallizes in the $Ba_2RbFe_2F_9$ structure type.¹⁸

Fundamentally, the crystal structures consist of corner sharing $[VA_6]^{n-}$ or $[MoA_6]^{n-}$ octahedra (M1) forming wavelike layers in the ab plane (Fig. 1). A second set of slightly elongated octahedra (M2, signified by lighter colors in Fig. 1) acts as bridges between those layers along c . The occupancy of the larger octahedral voids is quite minor at 2.5% for V and 8% for Mo, as they are generally too large to accommodate

Table 1 Single-crystal data and structure refinement of $K_3V_2O_{3.3}F_{5.7}$ and $K_3Mo_2O_{5.6}F_{3.4}$ with standard deviations in parentheses

Empirical formula	$K_3V_2O_{3.3}F_{5.7}$	$K_3Mo_2O_{5.6}F_{3.4}$
Molar mass/g mol ⁻¹	380.27	463.35
Crystal system	Rhombohedral	Rhombohedral
Space group	$R\bar{3}m$ (no. 166)	$R\bar{3}m$ (no. 166)
Powder data		
Powder diffractometer	STOE Stadi P	STOE Stadi P
Radiation; wavelength	MoK α_1 ; 70.93	MoK α_1 ; 70.93
λ /pm		
Temperature/K	295(2)	295(2)
a /Å	5.7390(2)	5.7830(1)
c /Å	21.0414(6)	21.1892(6)
V /Å ³	600.17(3)	613.69(3)
Single-crystal data		
Single-crystal diffractometer	Bruker D8 Quest	Bruker D8 Quest
Radiation; wavelength	MoK α_1 ; 71.073	MoK α_1 ; 71.073
λ /pm		
a /Å	5.7442(1)	5.7983(1)
c /Å	21.0484(6)	21.1801(5)
V /Å ³	601.46(2)	616.68(2)
Formula units per cell Z	3	3
Calculated density/g cm ⁻³	3.15	3.74
Crystal size/mm ³	$0.05 \times 0.04 \times 0.02$	$0.02 \times 0.02 \times 0.02$
Temperature/K	300(2)	300(2)
Absorption coefficient/mm ⁻¹	4.0	4.6
$F(000)/e$	546.4	641.4
2θ range/deg	5.8–75.0	5.8–65.0
Range in hkl	$-10 \leq h \leq 10$ $-10 \leq k \leq 10$ $-39 \leq l \leq 39$	$-10 \leq h \leq 10$ $-10 \leq k \leq 10$ $-39 \leq l \leq 39$
Total no. of reflections	10 117	11 993
Independent reflections	439	314
R_{int}	0.0338	0.0350
Reflections with $I \geq 2\sigma(I)$	385	291
Data; parameters	439; 23	314; 28
Absorption correction	Multiscan	Multiscan
Final R_1 ; wR_2 [$I > 2\sigma(I)$]	0.0271; 0.0622	0.0416; 0.1535
Final R_1 ; wR_2 (all data)	0.0310; 0.0655	0.0435; 0.1589
Goodness-of-fit on F^2	1.014	1.019
Largest diff. peak; hole/e Å ⁻³	0.96; -0.97	1.25; -0.79

Table 2 Wyckoff positions, atomic coordinates, equivalent isotropic displacement parameters U_{eq} (Å²) and site occupation factors for $K_3V_2O_{3.3}F_{5.7}$ (standard deviations in parentheses)

Atom	Wyckoff position	x	y	z	U_{eq}	s.o.f.
V1	6c	0	0	0.39708(2)	0.0235(2)	0.975
V2	6c	0	0	0.4814(8)	0.021(3)	0.025
K1	3a	0	0	0	0.0218(2)	1
K2	6c	0	0	0.20694(3)	0.0274(2)	1
O1	18h	0.4883(2)	-x	0.23110(5)	0.0221(2)	0.55
F1	18h	0.4883(2)	-x	0.23110(5)	0.0221(2)	0.45
F2	9e	$\frac{1}{2}$	0	0	0.0569(7)	1

$V^{4+/5+}$ ($r_{V^{4+}} = 0.72$ Å, $r_{V^{5+}} = 0.68$ Å) or Mo^{6+} ($r_{Mo^{6+}} = 0.73$ Å) cations. The greater degree of occupancy in the Mo variant can be attributed to the incorporation of Mo^{4+} , which possesses the perfect size to occupy these octahedral voids ($r_{Mo^{4+}} = 0.79$ Å). The amount of Mo^{4+} present within $K_3Mo_2O_{5.6}F_{3.4}$ was



Table 4 Wyckoff positions, atomic coordinates, equivalent isotropic displacement parameters U_{eq} (\AA^2) and site occupation factors for $\text{K}_3\text{Mo}_2\text{O}_{5.6}\text{F}_{3.4}$ (standard deviations in parentheses)

Atom	Wyckoff position	x	y	z	U_{eq}	s.o.f.
Mo1	$36i$	0.004(2)	0.046(2)	0.39116(4)	0.021(2)	0.16
Mo2	$3b$	0	0	$\frac{1}{2}$	0.036(2)	0.08
K1	$3a$	0	0	0	0.0336(6)	1
K2	$6c$	0	0	0.20564(9)	0.0327(5)	1
O1	$18h$	0.4889(2)	$-x$	0.2306(2)	0.0224(6)	0.933
F1	$18h$	0.4889(2)	$-x$	0.2306(2)	0.0224(6)	0.066
F2	$9e$	$\frac{1}{2}$	0	0	0.047(2)	1

Table 3 Anisotropic displacement parameters U_{ij} (\AA^2) for $\text{K}_3\text{V}_2\text{O}_{3.3}\text{F}_{5.7}$ (standard deviations in parentheses)

Atom	U_{11}	U_{22}	U_{33}	U_{23}	U_{13}	U_{12}
V1	0.0260(2)	U_{11}	0.0185(2)	0	0	0.01300(9)
V2	0.024(5)	U_{11}	0.015(5)	0	0	0.012(2)
K1	0.0212(2)	U_{11}	0.0229(3)	0	0	0.0106(2)
K2	0.0253(2)	U_{11}	0.0316(3)	0	0	0.0126(2)
O1 F1	0.0245(4)	U_{11}	0.0211(4)	$-0.0016(2)$	0.0016(2)	0.0152(4)
F2	0.0514(8)	0.104(2)	0.0331(8)	0.035(2)	0.0175(5)	0.052(2)

Table 5 Anisotropic displacement parameters U_{ij} (\AA^2) for $\text{K}_3\text{Mo}_2\text{O}_{5.6}\text{F}_{3.4}$ (standard deviations in parentheses)

Atom	U_{11}	U_{22}	U_{33}	U_{23}	U_{13}	U_{12}
Mo1	0.015(3)	0.021(2)	0.0262(6)	0.0007(4)	0.006(2)	0.010(2)
Mo2	0.022(2)	U_{11}	0.063(7)	0	0	0.011(2)
K1	0.0351(7)	U_{11}	0.031(2)	0	0	0.0176(4)
K2	0.0312(6)	U_{11}	0.0359(9)	0	0	0.0156(3)
O1 F1	0.0244(9)	U_{11}	0.025(2)	$-0.0021(4)$	0.0021(4)	0.0171(8)
F2	0.047(2)	0.070(3)	0.033(2)	0.021(2)	0.0104(9)	0.035(2)

determined by XPS and correlates directly with the occupancy of the M2 octahedral voids in between the main layers (see section XPS measurements). The main M1 octahedral voids are under-occupied in both compounds (to a large extent in $\text{K}_3\text{Mo}_2\text{O}_{5.6}\text{F}_{3.4}$, but only slightly in $\text{K}_3\text{V}_2\text{O}_{3.3}\text{F}_{5.7}$), so the sum of the occupancies over both crystallographic M positions was constrained to 1. The anion distribution within the title compounds was determined using BV and CHARDI calculations in conjunction with crystallographic data (see section BV, CHARDI and MAPLE calculations). Both compounds likely contain one mixed position occupied by both O and F and one discrete position occupied solely by F. Interestingly, the situation is reversed in $\text{K}_3\text{Nb}_2\text{O}_4\text{F}_5$ with only oxygen present at the discrete site. While this disparity might seem unintuitive at first glance, it is a direct result of the transition metal cation's position within the octahedron. The degree and direction of deflection from the "ideal" central position plays a crucial role in determining which crystallographic sites are suitable for an anion of a given charge and size.

The potassium atoms are coordinated either in a near-perfect cuboctahedral or a distorted anticuboctahedral

manner (Fig. 2). The bond lengths are quite similar when comparing the two variants, scaling ever so slightly with the size of the transition metal cation. In $\text{K}_3\text{Nb}_2\text{O}_4\text{F}_5$, the position of the K2 atoms is somewhat shifted, moving closer to the mixed positions and further away from the oxygen positions. In general, one would expect the opposite to be the case due to the more covalent nature of M–O bonds compared to M–F bonds. This specific discrepancy is most likely a result of the, on average, shorter M1–A2 (M = Nb, V, Mo and A2 = O, F) bond lengths in $\text{K}_3\text{Nb}_2\text{O}_4\text{F}_5$ compared to its V- and Mo-based counterparts (Table 6).

Powder X-ray diffraction

Rietveld refinements on powder diffraction measurements of product mixtures 1 (vanadium) and 2 (molybdenum) reveal that both $\text{K}_3\text{V}_2\text{O}_{3.3}\text{F}_{5.7}$ and $\text{K}_3\text{Mo}_2\text{O}_{5.6}\text{F}_{3.4}$ can be synthesized in high purity. In addition to the bulk target phase, product mixture 1 contains small quantities of an unknown by-product (Fig. 3). Product mixture 2 is composed of approximately 98.5(1) wt% $\text{K}_3\text{Mo}_2\text{O}_{5.6}\text{F}_{3.4}$ and 1.5(1) wt% $\text{K}_3\text{MoO}_3\text{F}_3$ as well as trace amounts of a third unidentified phase (Fig. 4).

BV, CHARDI and MAPLE calculations

Theoretical oxidation states of atoms within inorganic compounds can be calculated using mathematical concepts such as BV (bond valence)^{21–24} and CHARDI (charge distribution).^{25–27} These calculations allow for an evaluation of specific crystallographic positions based on their local environment and may offer valuable information in cases where certain distinctions are difficult or impossible to make using only crystallographic data. For the title compounds, these calculations first hinted at the presence of V^{4+} in $\text{K}_3\text{V}_2\text{O}_{3.3}\text{F}_{5.7}$ and allowed for a clear assignment of fluorine and oxygen to its two anion positions. Both methods are based on Pauling's well-known concept of bond length/bond strength combined with Hoppe's theory of effective coordination numbers.^{28–30} While BV relies on the use of empirically determined parameters for its calculations,^{31–33} CHARDI adopts a Madelung-type (point-charge) description of ionic crystal structures using only real observed distances, eliminating the need for pre-determined parameters altogether.³⁴

Bond valence sums (BVSs) for both compounds were calculated for all possible bonding scenarios and subsequently weighted according to their probability of occurrence. Mixed as well as incomplete occupancies were treated by multiplying the bond valence contribution with the occupancy for all affected bonds. CHARDI values were calculated using the program CHARDI2015 developed by Nespolo and Guillot.³⁵ At only 2.5%, occupancy of the V2 site was too low to allow for an accurate prediction of its oxidation state, so it had to be removed for the calculation of CHARDI values.

The calculation results deliver a clear picture regarding the anion distribution in both $\text{K}_3\text{V}_2\text{O}_{3.3}\text{F}_{5.7}$ and $\text{K}_3\text{Mo}_2\text{O}_{5.6}\text{F}_{3.4}$. Large differences in metal–anion bond lengths typically make identification of the nature of the anion straightforward. In $\text{K}_3\text{V}_2\text{O}_{3.3}\text{F}_{5.7}$, V1 is deflected along a special axis towards the



Table 6 Comparison of the basic crystallographic data of $K_3V_2O_{3.3}F_{5.7}$, $K_3Mo_2O_{5.6}F_{3.4}$ and $K_3Nb_2O_4F_5$ ¹⁸

Parameters	$K_3V_2O_{3.3}F_{5.7}$	$K_3Mo_2O_{5.6}F_{3.4}$	$K_3Nb_2O_4F_5$ ¹⁸
Eff. ionic radius (M^{n+})/Å for CN = 6	0.68 (V^{5+}), 0.72 (V^{4+})	0.75 (Mo^{6+}), 0.79 (Mo^{4+})	0.78 (Nb^{5+})
$a/\text{Å}$	5.7442(1)	5.7983(1)	5.799(2)
$c/\text{Å}$	21.0484(6)	21.1801(5)	21.371(4)
$V/\text{Å}^3$	601.46(2)	616.68(2)	622.4(4)
Atoms	A1 = O F, A2 = F	A1 = O F, A2 = F	A1 = O F, A2 = O
M1 occupancy	97.5%	92.0%	100%
M1–A1/Å	1.741(2) 3×	1.66(2) 1.83(2) 2.034(5)	1.816(2) 2.007(2) 2×
M1–A1 average/Å	1.741	1.84	1.943
M1–A2/Å	2.1331(3) 3×	1.895(4) 2.106(5) 2.253(9)	1.9338(8) 2×
M1–A2 average/Å	2.1331	2.085	1.998
M2 occupancy	2.5%	8.0%	Not occupied
M2–A1/Å	1.819(9) 3×	2.068(2) 6×	
M2–A1 average/Å	2.33(2) 3×	2.07	
K1–A1/Å	2.789(2) 6×	2.814(2) 6×	2.823(2) 6×
K1–A2/Å	2.8721(2) 6×	2.8991(2) 6×	2.900(2) 6×
K2–A1/Å	2.830(2) 3×	2.818(3) 3×	2.743(2) 3×
K2–A1 average/Å	2.9191(2) 6×	2.9492(5) 6×	2.966(2) 6×
K2–A2/Å	2.889	2.905	2.892
K2–A2 average/Å	3.1348(5) 3×	3.181(2) 3×	3.2813(8) 3×

mixed positions, simplifying the process of distinction even more so (Table 7). In $K_3Mo_2O_{5.6}F_{3.4}$, the deflection of the Mo atom is more nuanced, resulting in a total of six possible positions within a single octahedron. Although this makes the assignment of anions to specific positions somewhat more difficult and leaves more room for interpretation, the differences in bond lengths to A1 and A2 are still large enough on average to allow a reasonable distinction (Table 8). Anisotropic displacement parameters also support the assignment (see Fig. 1, Tables 3 and 5).

Regarding cation charges, an average calculated charge of +4.50 for V1 was the first indication that +5 might not be the sole oxidation state of V as we had originally assumed (due to the use of V_2O_5 as a starting material). This finding prompted further investigations by magnetic measurements, which indicated a distribution of approximately 1:1 for $V^{4+}:V^{5+}$ (see section magnetic measurements on $K_3V_2O_{3.3}F_{5.7}$) as well as a quantitative determination of the distribution by XPS (see section XPS measurements), suggesting an average charge of +4.65 for the V atoms. While CHARDI makes good predictions across the board, the BV method seems to have some difficulty accurately predicting the charges for positions Mo1 and K1 (both variants) in particular, underestimating the former and overestimating the latter.

In accordance with the increase in octahedron size, the charges for V2 and Mo2 are calculated to be significantly lower at +3.19 and +3.89, respectively, outlining the inadequacy of the larger M2 octahedral void as a host for $V^{4+/5+}$ and Mo^{6+} . While the residual electron density at that position is quite minor in $K_3V_2O_{3.3}F_{5.7}$ at $\approx 2.3 \text{ e } \text{Å}^{-3}$ (equating to $\approx 2.5\% \text{ V}$), it is substantial in $K_3Mo_2O_{5.6}F_{3.4}$ at $\approx 9.8 \text{ e } \text{Å}^{-3}$ (equating to $\approx 8\%$

Mo). In contrast, $K_3Nb_2O_4F_5$, which shares all the main structural features with the two oxyfluorides presented here, but exhibits a different anion distribution and deflection of the transition metal cation, shows essentially no residual electron density at that position.¹⁸ The main M1 octahedral void in $K_3V_2O_{3.3}F_{5.7}$ can host both V^{5+} ($r_{V^{5+}} = 0.68 \text{ Å}$) and V^{4+} ($r_{V^{4+}} = 0.72 \text{ Å}$), making it so only very few atoms find their way into the larger, less adequate M2 octahedral void in between the main layers. For $K_3Mo_2O_{5.6}F_{3.4}$, the size difference between the two available transition metal cations is more pronounced ($r_{Mo^{6+}} = 0.73 \text{ Å}$, $r_{Mo^{4+}} = 0.79 \text{ Å}$), rendering one octahedral void much more suitable for a cation of a given charge than the other.

The Madelung Part of Lattice Energy (MAPLE) of a new compound can be used to validate the proposed structure model by comparing the calculated value to the sum of MAPLE values of the simple (most often binary) components.^{36–38} For fluorido- and oxidometalates, a difference between those two values of less than 1% indicates a plausible structure model.³⁹ The values of the target compounds as well as all theoretical components were calculated using the program MAPLE V4 developed by Hübenthal.⁴⁰ Due to a limitation of the program, the heavily under-occupied M2 octahedral sites could not be considered in the calculations. Instead, occupancy of the main M1 sites was increased to 1 to compensate for the missing transition metal cations and achieve charge neutrality. Because both V^{4+} as well as V^{5+} are located at the same crystallographic position in $K_3V_2O_{3.3}F_{5.7}$, the exact composition could be used for the calculation. The sum of all theoretical components could, however, only be calculated to $K_3V_2O_3F_6$, as the complexity of the real composition



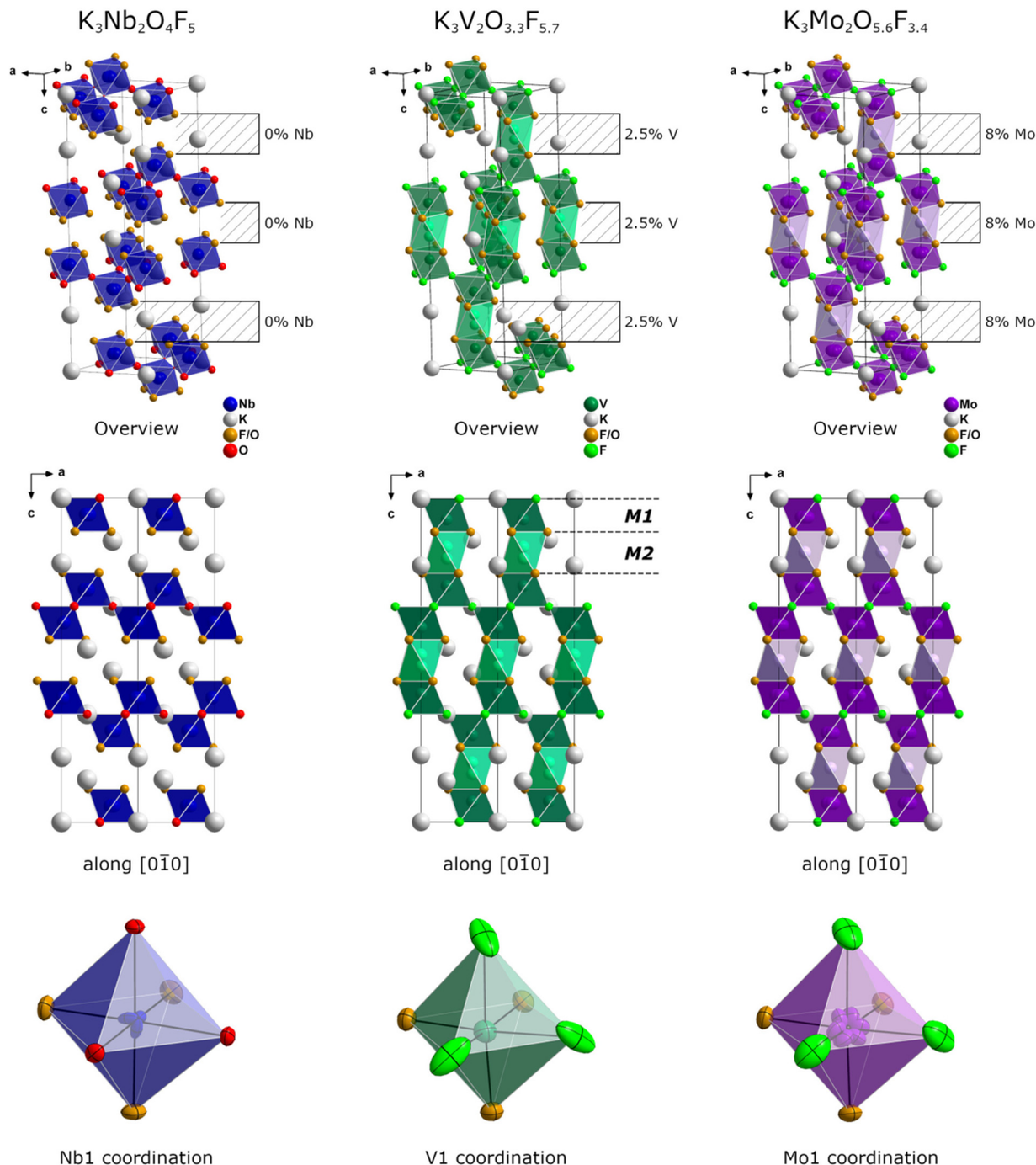


Fig. 1 Comparison of the crystal structures of the previously reported oxyfluoride $K_3Nb_2O_4F_5$,¹⁸ and its two siblings $K_3V_2O_{3.3}F_{5.7}$ and $K_3Mo_2O_{5.6}F_{3.4}$. An octahedral void, which remains empty in $K_3Nb_2O_4F_5$ and $Ba_2RbFe_2F_9$,²⁰ the compound that gives the structure type its name, is occupied to varying degrees in the title compounds. The exact positions of the transition metal cations also vary from compound to compound depending on the occupancy of the anion site. All possible positions are depicted, but of course only one can be occupied per octahedron.

could not be matched using simple binary and ternary components. Oxygen and fluorine are similar enough to still yield a difference in the MAPLE values of only 0.81%, indicating a plausible structure model (Table 9).

In the case of $K_3Mo_2O_{5.6}F_{3.4}$, consideration of the “real” sum formula would have necessitated a reduction of the main

Mo atoms’ charge from +6 to +5.8. Since we suspect the vast majority of Mo^{4+} to be situated at the M2 octahedral site, this reduced charge model does not represent reality, as a difference in the MAPLE values of 6.22% further emphasizes. To compensate for that, a second calculation was carried out using the sum formula $K_3Mo_2O_6F_3$, essentially ignoring the



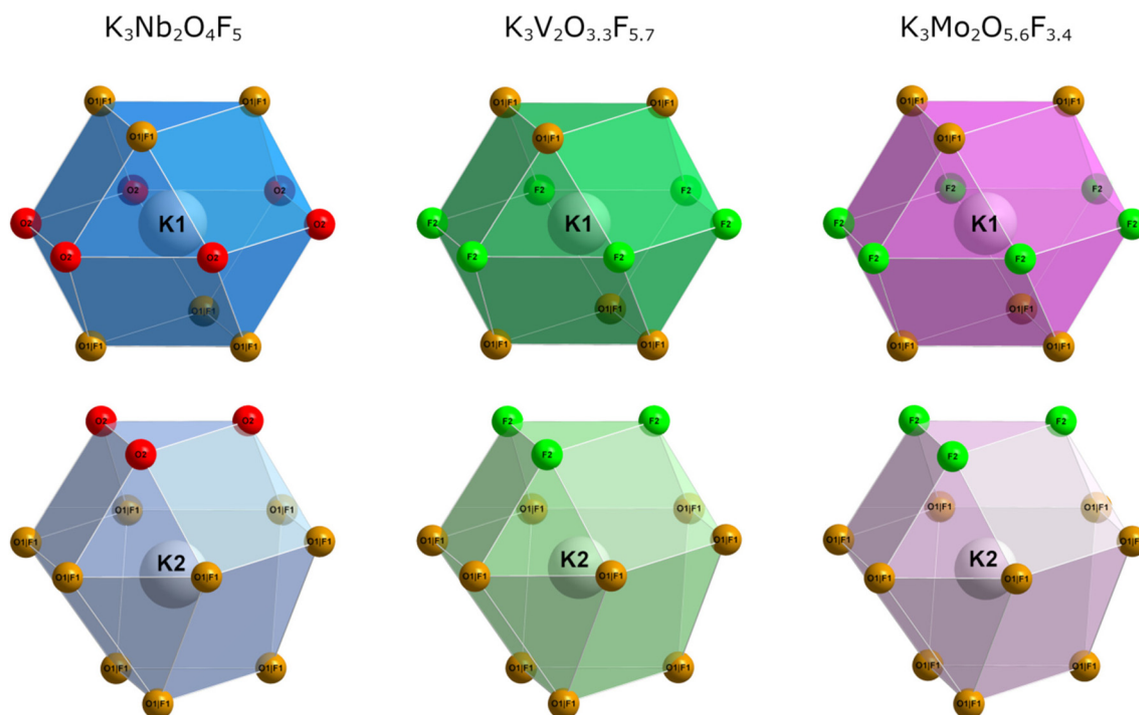


Fig. 2 Cuboctahedral coordination sphere of K1 and anticuboctahedral coordination sphere of K2 in $K_3Nb_2O_4F_5$ (left), $K_3V_2O_{3.3}F_{5.7}$ (middle) and $K_3Mo_2O_{5.6}F_{3.4}$ (right).

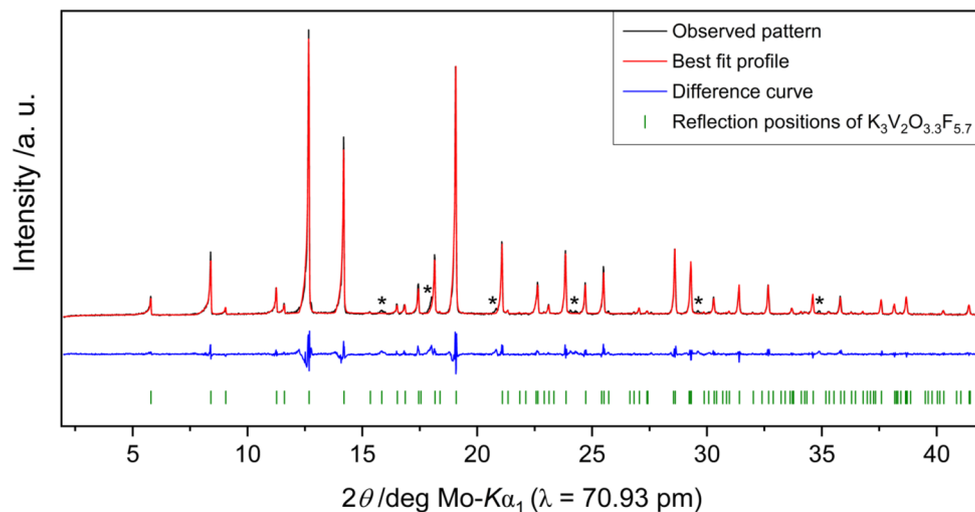


Fig. 3 Rietveld plot of product mixture 1. The observed diffraction pattern is plotted in black, the best fit profile in red and the difference curve in blue. Reflection positions of $K_3V_2O_{3.3}F_{5.7}$ are marked in green. Asterisks denote reflections belonging to an unknown side phase. ($R_{exp} = 6.73\%$, $R_{wp} = 12.55\%$, $R_p = 9.61\%$, $GoF = 1.87$, background: Chebyshev polynomial of 12th order).

existence of the approximately 8% Mo^{4+} present within the compound. This special treatment leads to a difference in the MAPLE values of 0.44%, indicating a sensible structure solution in the absence of Mo^{4+} (Table 10). The small amount of Mo^{6+} reduced to Mo^{4+} during synthesis can easily be hosted by the much larger M2 octahedral void, as evidenced by BV and CHARDI calculations.

The two sets of calculations for the title compounds further reinforce some of our experimental observations. The V variant is only stable when vanadium is present in both oxidation states +5 and +4 simultaneously. Any attempts at synthesis of the target phase using only V^{4+} lead to the formation of K_2VOF_4 instead. Blends of V_2O_5 and VO_2 can, however, be used to synthesize samples of equal



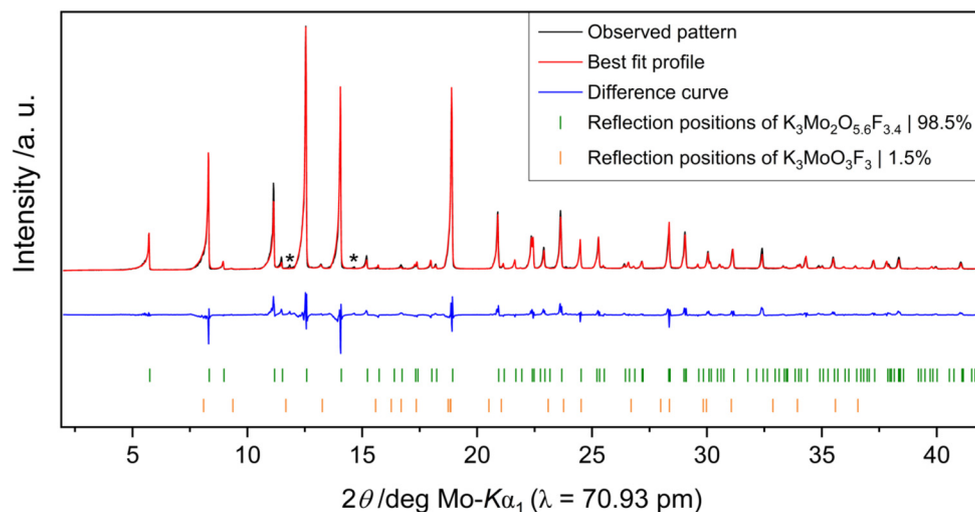


Fig. 4 Rietveld plot of product mixture 2. The observed diffraction pattern is plotted in black, the best fit profile in red and the difference curve in blue. Reflection positions of $K_3Mo_2O_{5.6}F_{3.4}$ and the minor side phase $K_3MoO_3F_3$ are marked in green and orange, respectively. Asterisks denote reflections belonging to an unknown side phase. ($R_{exp} = 1.34\%$, $R_{wp} = 13.46\%$, $R_p = 9.76\%$, $Goof = 10.08$, background: Chebyshev polynomial of 12th order).

Table 7 Charge distribution in $K_3V_2O_{3.3}F_{5.7}$ calculated using BV ($\sum V$) and CHARDI ($\sum Q$)

	V1	V2	K1	K2	O1 F1	F2
$\sum V$	+4.34	+3.19	+1.42	+1.13	-1.54	-0.91
$\sum Q$	+4.65	—	+1.00	+1.00	-1.55	-1.00
Average	+4.50	—	+1.21	+1.07	-1.55	-0.96
Theoretical value	+4.65	+4.00	+1.00	+1.00	-1.55	-1.00

Table 8 Charge distribution in $K_3Mo_2O_{5.6}F_{3.4}$ calculated using BV ($\sum V$) and CHARDI ($\sum Q$)

	Mo1	Mo2	K1	K2	O1 F1	F2
$\sum V$	+5.39	+3.82	+1.44	+1.22	-1.86	-1.28
$\sum Q$	+5.94	+3.96	+0.99	+0.99	-1.96	-1.02
Average	+5.67	+3.89	+1.22	+1.11	-1.91	-1.15
Theoretical value	+6.00	+4.00	+1.00	+1.00	-1.94	-1.00

Table 9 Comparison between the calculated MAPLE values for $K_3V_2O_{3.3}F_{5.7}$ and the corresponding theoretical starting materials KF ($Fm\bar{3}m$), VO_2F ($R\bar{3}c$) and $KVOF_3$ ($C2/m$) according to the equation: $2KF + VO_2F + KVOF_3 \rightarrow K_3V_2O_{3.3}F_{5.7}$. Due to the complex sum formula of the target phase, the O/F ratio does not match exactly

KF ($Fm\bar{3}m$), ⁴³ kJ mol ⁻¹	908 (×2)
VO_2F ($R\bar{3}c$), ⁴⁴ kJ mol ⁻¹	18 296
$KVOF_3$ ($C2/m$), ⁴⁵ kJ mol ⁻¹	12 384
Σ of starting materials, kJ mol ⁻¹	32 496
$K_3V_2O_{3.3}F_{5.7}$, kJ mol ⁻¹	32 762
Difference, kJ mol ⁻¹	266
Difference in %	0.81

purity, as long as the V^{4+} concentration does not exceed 35%. Beyond that critical point, K_2VOF_4 is formed as a side phase proportional to the amount of excess V^{4+} . This

Table 10 Comparison between the calculated MAPLE values for $K_3Mo_2O_6F_3$ and the corresponding theoretical starting materials KF ($Fm\bar{3}m$) and MoO_3 ($Pbnm$) according to the equation: $3KF + 2MoO_3 \rightarrow K_3Mo_2O_6F_3$

KF ($Fm\bar{3}m$), ⁴³ kJ mol ⁻¹	908 (×3)
MoO_3 ($Pbnm$), ⁴⁶ kJ mol ⁻¹	25 344 (×2)
Σ of starting materials, kJ mol ⁻¹	53 412
$K_3Mo_2O_6F_3$, kJ mol ⁻¹	53 177
Difference, kJ mol ⁻¹	235
Difference in %	0.44

suggests either that the V^{4+} content is variable within a certain window of approx. 35–58% due to percentage-based *in situ* reduction of V^{5+} , or more likely that the reduction is conditional and ceases at approx. 35% to facilitate the formation of the new phase. V_2O_5 has been reported to go through a sequence of transformations from its original state *via* VO_2 to V_2O_3 upon heating to 600 °C in a vacuum.⁴¹ While obviously influenced by the extreme synthesis conditions in our case, we suspect that this process is likely cut short by the formation of the title phase, limiting the thermal decomposition of V_2O_5 .

For the Mo-based compound, we measured several single crystals and found varying amounts of Mo located at the larger M2 octahedral site ranging from 5–8%. With MAPLE calculations predicting a perfectly stable structure for $K_3Mo_2O_6F_3$ without the presence of any Mo^{4+} or occupancy of the M2 site, we do not believe the reduction of Mo^{6+} to play a major role in the formation of the new compound. Instead, we suspect that a small but varying amount of MoO_3 naturally decomposes to MoO_2 and O_2 at high temperatures,⁴² leading to the incorporation of Mo^{4+} into the M2 octahedral void, which happens to be perfectly sized to host a cation of this nature.



Magnetic measurements on $K_3V_2O_{3.3}F_{5.7}$

The $K_3V_2O_{3.3}F_{5.7}$ sample behaves like a Curie–Weiss paramagnet above 50 K (Fig. 5, top). Fitting of the 10 kOe measurement according to a modified Curie–Weiss law reveals an experimental magnetic moment of $0.95(1)\mu_B$ V atom⁻¹. The Weiss constant is $\theta_P = 3.3(1)$ K and the temperature independent contribution $\chi_0 = -0.003(1) \times 10^{-5}$ emu mol⁻¹. For the ionic formula splitting $3K^+ 2V^{4+} 2O^{2-} 7F^-$ with solely tetravalent vanadium (d^1 system) one would expect an effective magnetic moment of $1.73\mu_B$ per V^{4+} ion.⁴⁷ The simultaneous presence of both fluorine and oxygen, distributed over the same crystallographic sites, does, however, provide a lot of leeway in terms of possible oxidation states for vanadium. Considering a composition of $3K^+ V^{4+} V^{5+} 3O^{2-} 6F^-$ with half of the vanadium atoms in the pentavalent state, only half the moment is expected to occur. We can thus conclude, that the sample most likely contains a distribution of domains of compositions $K_3V^{4+}_{2-x}V^{5+}_xO_{2+x}F_{7-x}$ with x around 1. Mixed-valence vanadate compounds have previously been reported to possess magnetic moments corresponding to non-integer numbers of electrons.^{48–50}

The recorded magnetization isotherms are shown at the bottom of Fig. 5. For the 10 and 50 K isotherms, we observe

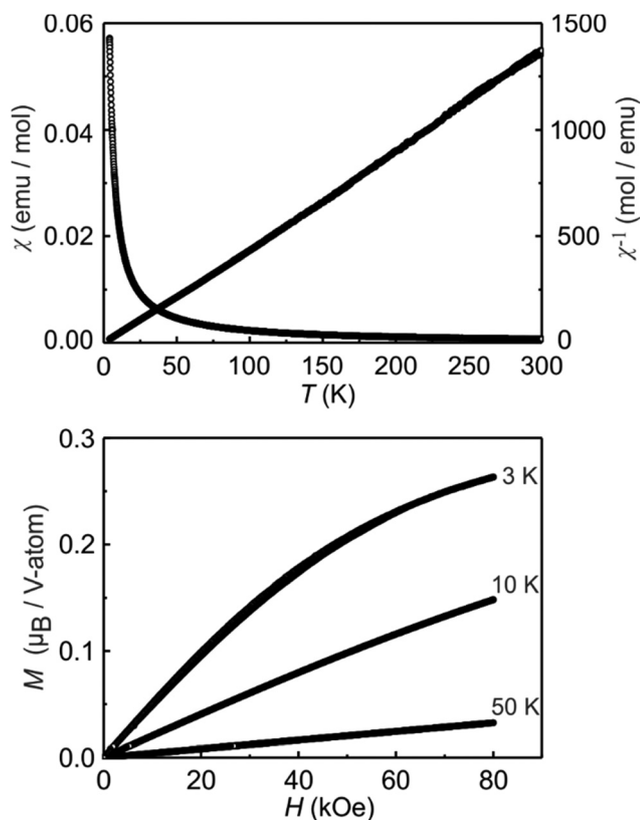


Fig. 5 Magnetic properties of a sample with the composition $K_3V_2O_{3.3}F_{5.7}$ (top) Temperature dependence of the magnetic susceptibility and its inverse (χ and χ^{-1} data) recorded with an external field of 10 kOe (bottom) magnetization isotherms recorded at 3, 10 and 50 K.

linear increases as expected for paramagnetic materials. The 3 K magnetization isotherm shows a continuous, field-induced increase of the magnetization with the highest magnetization of only $0.26(1)\mu_B$ V atom⁻¹ at 3 K and 80 kOe. Thus, much higher field strengths are necessary for full parallel spin alignment. The low value of the magnetization is also an indication of the low proportion of V^{4+} in the studied sample.

XPS measurements

As BV calculations as well as magnetic measurements indicated the simultaneous presence of multiple oxidation states in $K_3V_2O_{3.3}F_{5.7}$, XPS measurements were carried out on both title compounds to be able to experimentally determine the oxidation states as well as their relative concentration. Initial survey spectra of $K_3V_2O_{3.3}F_{5.7}$ and $K_3Mo_2O_{5.6}F_{3.4}$ confirmed the presence of all expected elements along with adventitious carbon (Fig. 6a).

In order to distinguish between individual oxidation states of Mo and V atoms, their respective 3d (Mo) and 2p (V) core level peaks were measured and the data fitted with Voigt profiles to calculate the atomic ratios from the integrated areas (Fig. 6b and c). For details on the fitting procedure, refer to the Experimental section. As indicated by our previous investigations, vanadium is present in two different oxidation states, namely V^{5+} and V^{4+} in a ratio of approximately 1.9 : 1. Molybdenum also exhibits two different oxidation states: Mo^{6+} and Mo^{4+} . The highly charged Mo^{6+} cation is heavily favored, with a ratio of 10 : 1. The amount of Mo^{4+} determined by XPS (approx. 9%) is directly proportional to the occupancy of the larger M2 octahedral void in $K_3Mo_2O_{5.6}F_{3.4}$ (approx. 8%).

Having determined concentrations and charges of the cations, it is possible to calculate the amount of oxygen and fluorine present in each compound using the principle of charge neutrality: $K_3Mo_{1.8}^{6+}Mo_{0.2}^{4+}O_{5.6}F_{3.4}$ and $K_3V_{1.3}^{5+}V_{0.7}^{4+}O_{3.3}F_{5.7}$.

Conclusion

Oxyfluorides exhibit an incredible degree of compositional adaptability due to their mixed-anionic nature. Even seemingly straightforward synthetic routes can lead to unexpectedly complex products. When both oxygen and fluorine are present as anions, it may no longer be easy to derive cation oxidation states. Similarly, it can be difficult to predict exactly how certain starting materials will behave under a given set of reaction conditions, especially extreme ones. Partial oxidation or reduction of suitable cations during synthesis can alter local atomic environments and influence the distribution of O/F anions in the final product. In some cases, such as with $K_3V_2O_{3.3}F_{5.7}$, mixed valence is crucial for phase stability. In other cases, such as with $K_3Mo_2O_{5.6}F_{3.4}$, cations of different oxidation states can be accommodated in different ways and to varying degrees. The exact influence of cation oxidation state and O/F anion distribution on compound formation and stability is complex and not easily predicted. To accurately describe



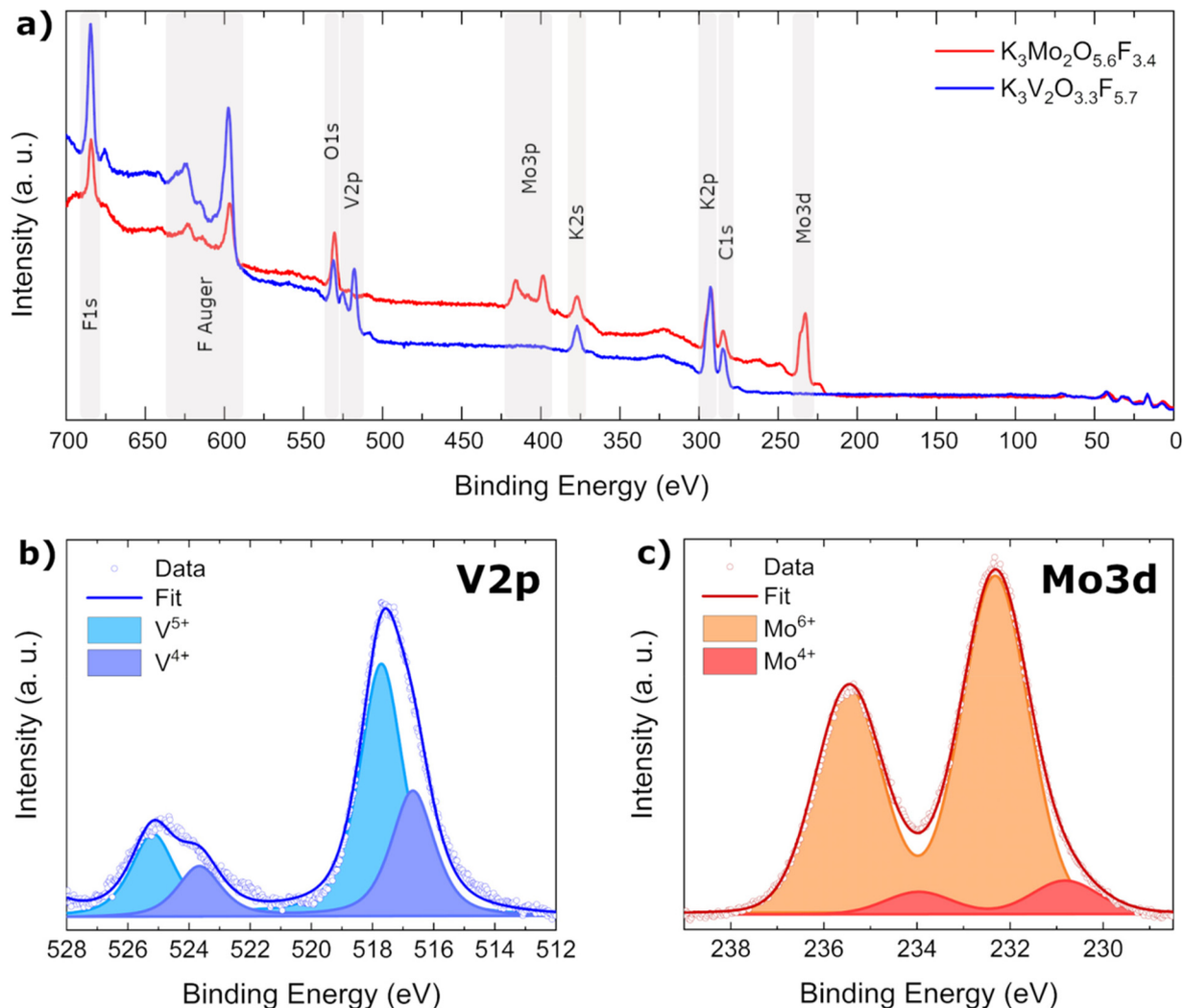


Fig. 6 (a) XPS survey spectra of $\text{K}_3\text{Mo}_2\text{O}_{5.6}\text{F}_{3.4}$ (red) and $\text{K}_3\text{V}_2\text{O}_{3.3}\text{F}_{5.7}$ (blue) with the most intense core level signals labeled. Detailed scans of the core level peaks are presented in (b) for V2p in $\text{K}_3\text{V}_2\text{O}_{3.3}\text{F}_{5.7}$ and in (c) for Mo3d in $\text{K}_3\text{Mo}_2\text{O}_{5.6}\text{F}_{3.4}$.

oxyfluorides and understand their nuances, it is important to consider how the two interact with and influence each other.

Experimental section

Synthetic procedure

Both $\text{K}_3\text{V}_2\text{O}_{3.3}\text{F}_{5.7}$ and $\text{K}_3\text{Mo}_2\text{O}_{5.6}\text{F}_{3.4}$ were synthesized under high-pressure/high-temperature conditions employing a special multianvil setup for a uniaxial 1000 t frame press (Max Voggenreiter GmbH, Mainleus, Germany). The starting mixtures were prepared in a glovebox (MBraun Inertgas-System GmbH, Garching, Germany) under argon atmosphere by simple weighing and thorough homogenization in an agate mortar. 50.00 mg (0.6402 mmol) KHF_2 (Alfa Aesar, 99+% metals basis) and 33.27 mg (0.1829 mmol) V_2O_5 (Merck, extra pure) in a stoichiometric ratio of 3.5 : 1 served as the starting materials for the synthesis of $\text{K}_3\text{V}_2\text{O}_{3.3}\text{F}_{5.7}$. Alternatively, a mixture of V_2O_5 and VO_2 (Alfa Aesar, 99% metals basis) may

be used, provided that the concentration of tetravalent vanadium does not exceed 35%. In the case of $\text{K}_3\text{Mo}_2\text{O}_{5.6}\text{F}_{3.4}$, a stoichiometric (3 : 2) mixture of 30.00 mg (0.5164 mmol) KF (Fluka, >99.0%) and 49.55 mg (0.3443 mmol) MoO_3 (Merck, >99.5%) proved to yield the purest product. The resulting powders were subsequently transferred into a platinum capsule (99.95%, Ögussa, Vienna, Austria), which was, in turn, placed inside a boron nitride crucible (Henze Boron Nitride Products AG, Lauben, Germany). In the next step, the crucible was inserted into a ZrO_2 sleeve, whose inside is coated with graphite. The graphite acts as a resistive heating element with the ZrO_2 being responsible for keeping the heat inside the reaction chamber. Molybdenum platelets on either side of the cylindrical construction enable conductivity from the graphite oven to the outside. The construction was then inserted into the pressure medium – an octahedron consisting of MgO doped with 5% Cr_2O_3 . The octahedral assembly was further surrounded by eight tungsten carbide cubes (Hawedia, Marklkofen, Germany) with truncated corners and held



together by six glass fiber reinforced plastic panels. The experimental setup was completed upon placement of the cube into a Walker-type module alongside six steel anvils to allow for a quasi-hydrostatic compression of the sample using a uniaxial (vertical) hydraulic press. More detailed descriptions of various multianvil assemblies can be retrieved from the literature.^{51–54}

The reaction conditions for the syntheses of the title compounds are identical. First, the starting materials were compressed to 8.0 GPa in 215 min followed by an increase in temperature to 900 °C. The temperature was held constant for an additional 30 min before it was reduced to 350 °C in 60 min. At that point the reaction was quenched, reaching room temperature in a matter of minutes. To complete the experiment, the pressure acting on the assembly was gradually reduced to atmospheric pressure over a period of 900 min. The octahedral assembly was removed from the experimental setup and transferred into a glovebox. The platinum capsule holding the sample was then recovered, separated from other assembly materials and opened to reveal a dark green ($\text{K}_3\text{V}_2\text{O}_{3.3}\text{F}_{5.7}$) or deep red ($\text{K}_3\text{Mo}_2\text{O}_{5.6}\text{F}_{3.4}$) polycrystalline sample (Fig. 7).

Single-crystal X-ray diffraction

A glovebox-stored sample of each title compound was transferred onto a microscope slide and covered with perfluoropolyether to prevent oxidation and/or hydrolysis during the selection of suitable single crystals under a polarization microscope outside of the glovebox. Each crystal was subsequently mounted onto a Bruker D8 Quest single-crystal diffractometer (Bruker, Billerica, USA) equipped with a Photon III C14 detector system. Measurements were carried out at an ambient temperature of 300(2) K using $\text{MoK}\alpha$ radiation ($\lambda = 71.073$ pm) generated by an Incoatec Microfocus X-ray tube (Incoatec, Geesthacht, Germany) in conjunction with a multilayer monochromator.

Diffraction data was collected with the program APEX V2021.4-0.⁵⁵ Data reduction, integration and unit cell refinement were carried out in SAINT-V 8.40B.⁵⁶ A multi-scan absorption correction was applied to the intensity data using the software SADABS-2016/2.⁵⁷ Initially, space groups $R3$ (no. 146) and $R\bar{3}$ (no. 148) were considered for structure solution. An ADDSYM^{58,59} check in PLATON⁶⁰ indicated the presence of higher symmetry, suggesting space group $R\bar{3}m$ (no. 166) instead. Using the structure refinement and analysis program OLEX2,⁶¹ the structure was solved in the proposed space group $R\bar{3}m$ (no. 166) using SHELXTL-2018/2⁶² and refined with the OLEX2.REFINE⁶³ refinement package employing Gauss-Newton minimization. The datasets used for refinement were restricted to angles below 75° for $\text{K}_3\text{V}_2\text{O}_{3.3}\text{F}_{5.7}$ and 65° for $\text{K}_3\text{Mo}_2\text{O}_{5.6}\text{F}_{3.4}$ to account for limited diffraction intensity at larger angles due to comparatively small crystal size (especially in the case of $\text{K}_3\text{Mo}_2\text{O}_{5.6}\text{F}_{3.4}$). Atomic coordinates were standardized using the PLATON subroutine STRUCTURE TIDY⁶⁴ to allow for effortless comparison between the two title compounds and other related structures. All visual representations of the crystal structures were created with DIAMOND 4.6.8.⁶⁵

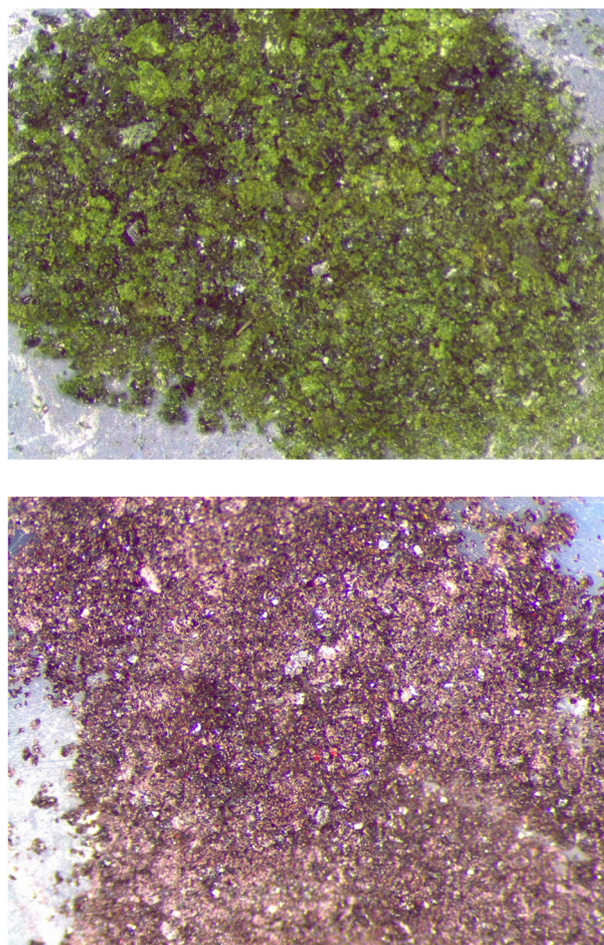


Fig. 7 Photographs of $\text{K}_3\text{V}_2\text{O}_{3.3}\text{F}_{5.7}$ (left) and $\text{K}_3\text{Mo}_2\text{O}_{5.6}\text{F}_{3.4}$ (right) under an optical microscope at 1.5 \times magnification. The shade of green of $\text{K}_3\text{V}_2\text{O}_{3.3}\text{F}_{5.7}$ can vary quite heavily from experiment to experiment, sometimes appearing almost entirely black.

Powder X-ray diffraction

Powder diffractograms of high-purity $\text{K}_3\text{V}_2\text{O}_{3.3}\text{F}_{5.7}$ and $\text{K}_3\text{Mo}_2\text{O}_{5.6}\text{F}_{3.4}$ were recorded on a STOE STADI P diffractometer (STOE & CIE GmbH, Darmstadt, Germany) operating in Debye-Scherrer transmission geometry. To prevent reactions with oxygen and/or air moisture, the samples were finely ground and sealed inside soda-lime glass capillaries prior to removal from the glovebox. Ge(111)-monochromatized $\text{MoK}\alpha_1$ radiation ($\lambda = 70.93$ pm) diffracted by the sample was recorded on a DECTRIS MYTHEN2 DCS4 detector system. The measurement was carried out at an ambient temperature of 295(2) K across a 2θ range of 2.0–42.0° with a step size of 0.015°. Rietveld refinements of the recorded diffractograms were performed using the software package DIFFRAC^{plus}-Topas 4.2 (Bruker AXS, Karlsruhe, Germany).⁶⁶

XPS measurements

For XPS measurements, finely ground powders of both title compounds were applied as a thin layer to a conductive adhesive tape using a fine brush. The tape was then stuck onto



the sample holders and transferred to an ultra-high vacuum system. Measurements were performed using a non-monochromatic MgK α source (VG) with an excitation energy of 1253.6 eV. The emitted photoelectrons were detected by a hemispherical analyzer (Specs, Phoibos 100) at pass energies of 90 eV for the survey scans and 20 eV for the detailed scans. Due to the inherently low conductivity of powdered samples, a charge induced shift of all binding energies by 5.5 eV ($K_3V_2O_{3.3}F_{5.7}$) and 5.07 eV ($K_3Mo_2O_{5.6}F_{3.4}$) was observed. This was corrected during data processing by shifting the 1s core level peak of adventitious carbon to 284.7 eV and adjusting the rest of the dataset accordingly. To determine the oxidation state and relative concentration of the individual V and Mo atoms, the core level peaks of Mo3d and V2p were fitted using the software XPSPEAK 4.1⁶⁷ using Voigt profiles and assuming a Shirley background. To distinguish between oxidation states of vanadium, O1s core level peaks were used as a reference point.⁶⁸ The obtained fit has a full width at half maximum (FWHM) of 1.88 eV with a Lorentzian/Gaussian ratio of 23%. The spin-orbit splitting of the peaks is 7.58 and 7.30 eV for V⁴⁺ and V⁵⁺, respectively. The Mo3d core level doublet was fitted with a FWHM of 1.7 eV, a Lorentzian/Gaussian ratio of 11% and a spin-orbit splitting of 3.15 eV.

Magnetic measurements

The sample with the initial composition $K_3V_2O_{3.3}F_{5.7}$ was pure on the level of X-ray powder diffraction and ground to a fine powder. It was filled into a polypropylene capsule for investigation, using the Vibrating Sample Magnetometer (VSM) option of a Physical Property Measurement System (PPMS) by Quantum Design. The capsule was inserted into a brass sample holder, which was attached to the sample holder rod of the VSM. The sample was then investigated in a temperature range from 2.5 to 300 K through measurements of $M(T,H)$ with applied fields up to 80 kOe. Fitting and plotting of the data was performed with OriginPro 2016G⁶⁹ and graphical editing with the program CorelDRAW2017.⁷⁰

Author contributions

Conceptualization, single-crystal structure solution, powder X-ray diffraction, BV, CHARDI, MAPLE calculations and original draft by FZ. Synthetic work by EW and FZ. BÖ and SO carried out the XPS analysis and contributed to the drafting of the corresponding section. MKR and RP carried out the magnetic measurements and provided a complete first draft of this section. HH supervised and guided this project, in addition to providing the necessary laboratory equipment and tools. The manuscript was reviewed and edited by all authors.

Data availability

Deposition Numbers 2320478 and 2320483† contain the supplementary crystallographic data for $K_3V_2O_{3.3}F_{5.7}$ and $K_3Mo_2O_{5.6}F_{3.4}$, respectively.

Conflicts of interest

The authors declare no competing interests related to this article.

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