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Pseudotetragonal Structure of $Li_{2+x}Ce_x^{3+}Ce_{12-x}^{4+}F_{50}$: The First Mixed Valence **Cerium Fluoride**

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The crystal structure of the new Li_{5.5}Ce₁₂F₅₀ compound has been fully characterized by single-crystal and synchrotron powder X-ray diffraction. An accurate pseudotetragonal structure was described in the monoclinic P2₁ space group with 68 independent crystallographic sites. The $\text{Li}_{5.5}\text{Ce}_{12}\text{F}_{50}$ composition belongs to the $\text{Li}_{2+x}\text{Ce}_x^{\ 3+}\text{Ce}_{12-x}^{\ 4+}\text{F}_{50}$ solid solution. Its structure consists of an opened fluorine framework where a channel network allows the intercalation of relatively mobile lithium cations, inducing the formation of the mixed-valence cerium (the intercalation of Li⁺ leads to the reduction of a part of Ce^{4+} to Ce^{3+}). One part of the lithium ions, necessary for the electroneutrality of the tetravalent equivalent cerium fluoride ($Li_2Ce_{12}F_{50}$ composition), is in a locked fluorine polyhedron. Only the supplementary x amount of lithium is able to be exchanged in $Li_{2+x}Ce_x^{3+}Ce_{12-x}^{4+}F_{50}$. The structure of $Li_{2+x}Ce_x^{3+}Ce_{12-x}^{4+}F_{50}$ is a rearrangement, due to lithium intercalation, of the base CeF_4 structure. Bond valence calculation on Ce sites, Ce coordination polyhedra volumes, and a calculated Ce cationic radius give the indication of a partial long-range ordering of trivalent and tetravalent cerium cations in specific slabs of the structure. 7Li NMR spectroscopy and XPS analyses have confirmed all of the structure details.

1. Introduction

In spite of their usual "rare earths" name, the lanthanide elements are now largely used in advanced technologies. The applications relate to optics with the development of solid lasers, optical fibers for telecommunications, and devices with electrochromic or electroluminescent diodes and so on. The remarkable magnetic properties of some inorganic fluorides result also in interesting potentialities. The diversity and the high added value of these applications justify the numerous research projects undertaken on these lanthanides and actinides fluorides. In particular, the crystal chemistry of terbium fluorides has been extensively studied because of the half-filled shell of the 4f' electronic configuration of terbium, which stabilizes the tetravalent cation. Compounds exhibiting mixed valences are subject to many studies because of their potential applications related to the electronic exchange and magnetism.

Contrary to the cerium ions for which the +4 oxidation state can be stabilized in solution, the Tb⁴⁺ cation exhibits a high instability, justifying the low number of studies on tetravalent terbium compounds. $^{1-4}$ Nevertheless, Tb^{4+} is more easily stabilized by fluorine anions than by oxide anions. The tetravalent Ce⁴⁺ cation can be stabilized in either an oxygenated or fluorinated medium.⁵⁻⁷ Up to now, very few mixed-valence terbium fluorides were reported in the literature. Among them, the Tb₄F₁₅ compound was synthesized by Brekhovskikh et al.⁸ and characterized by Popov et al. Other mixed-valence terbium fluorides have

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been synthesized and characterized in our laboratory: KTb₃F₁₂^{10,11} and K₂Tb₄F₁₇¹² present a long-range ordering between Tb³⁺ and Tb⁴⁺. For others, such as the aluminumcontaining terbium fluoride $Rb_2AlTb_3F_{16}^{13}$ and $RbAl_2Tb_4-F_{22}^{14}$ Tb^{3+} and Tb^{4+} are statistically distributed into the same crystallographic sites. Mixed-valence ternary terbium fluoride appears at a molar ratio of $TbF_4/AF > 1$ (A = Li, K, Rb, Cs). Chilingarov et al. 15 showed that the decomposition of TbF₄ into TbF₃ and atomic fluorine occurs at temperatures higher than 550 °C, even in a fluorine atmosphere. If the synthesis conditions allow it, TbF4 in excess transforms to trivalent fluoride by decomposition, which is suitable to immediately combine with the other components of the reactive mixture. This forms Tb³⁺/Tb⁴⁺ mixed-valence terbium fluoride. In other words, a thermally induced transformation occurs from the AF-TbF₄ binary system into the AF-TbF₃-TbF₄ ternary system. The obtained mixed-valence compounds are thermodynamically stable. Nevertheless, the reaction temperature and the cooling rate must be optimized. Thus, the terbium fluoride initially formulated, KTb_2F_9 ($TbF_4/KF = 2$), is actually a mixed-valence compound with the K₂Tb₄F₁₇

formula, that is, $K_2Tb^{3+}Tb^{4+}{}_3F_{17}$. In the LiF- MF_4 (M = Ce, Th, and U) binary systems, compounds formulated as LiM_4F_{17} were reported, $^{16-19}$ but no structural study confirmed their real stoichiometries. By analogy with terbium tetrafluorides, their location above the MF₄/LiF molar ratio equal to unity increases the ambiguity about their real stoichiometry. Indeed, the initial "LiTh₄F₁₇" composition has been corrected as Th₆F₂₄·H₂O, a hydrated thorium tetrafluoride, thanks to a structural study which has been carried out on a single crystal by Cousson et al. 19 Nevertheless, these authors mentioned that lithium ions are necessary for the synthesis of the hydrated tetrafluoride, although they are absent from the structure.

With the dual aim of unambiguously determining the real stoichiometry of the "Li M_4F_{17} " (M = Ce, Th, and U) compounds and investigating whether the mechanism of formation of mixed-valence terbium fluorides is transposable to LiF/MF_4 systems, a compound of nominal composition 1LiF, 1CeF₃, and 3CeF₄ was synthesized and characterized. The synthesized Li_{5.5}Ce₁₂F₅₀ compound has a pseudotetragonal symmetry. A first paper describes the weak monoclinic distortion of the lattice. The present paper gives its complete crystallographic description together with a ⁷Li magic-angle spinning (MAS) NMR investigation. The mixed

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valence has been investigated by XPS experiments in order to extract the Ce⁴⁺/Ce³⁺ ratio.

2. Experimental Section

- 2.1. Synthesis. Samples, used for single-crystal and synchrotron powder measurements, were synthesized by solid-state reactions involving a mixture of LiF (0.058 g), CeF₃ (0.444 g), and CeF₄ (1.459 g) starting materials in a 1:1:3 ratio. Pure CeF₄ tetrafluoride has been prepared by the following route: (i) dissolution of CeO₂ in a HF solution (40%), (ii) after evaporation of the solution, the dried resulting powder was fluorinated under a pure F₂ gas stream at 500 °C. CeF₃ was a commercial product (99.9% Aldrich). LiF (99.9%, Aldrich) was dehydrated by heating at 110 °C overnight in a primary vacuum. The 1:1:3 mixture was loaded into a platinum capsule (40 mm of length, 5 mm of diameter). The airtight platinum capsule was heated at 300 °C for 24 h, and then a temperature of 550 °C was applied over 72 h.
- 2.2. X-Ray Diffraction. Single-crystal X-ray diffraction intensity data were collected from two orange single crystals at room temperature by the use of a Stoe IPDS II image plate diffractometer. Both data were similar. The data set from the crystal given the best $R_{\rm int}$ was used to solve and refine the structure. The details of data collection and structure refinement are given in Table 1.

Synchrotron powder diffraction data of a freshly synthesized compound was obtained at the Swiss-Norwegian Beamline (SNBL) at the ESRF Grenoble (six analyzer crystal detector). More details are given in Table 1. Synchrotron powder diffraction was used to check the symmetry of the compound, that is, the weak lattice distortion unresolved with the image plate diffractometer. More details are given in ref 20.

- 2.3. ⁷Li MAS NMR Spectroscopy. The ⁷Li and ¹H spectra were recorded with magic angle spinning at room temperature using a Bruker MSL300 spectrometer operating at frequencies of 116.6 and 300.1 MHz for ⁷ Li and ¹ H nuclei, respectively. The spinning rate was equal to 14 kHz using a 4 mm Bruker probe. A single $\pi/2$ pulse sequence was used (τ -acquisition with $\tau = 6$ and $5 \mu s$ for ¹Li and ¹H nuclei, respectively). A total of 1000 and 128 scans were recorded with a recycling time of 1 and 5 s for ⁷Li and ¹H, respectively. The ¹H chemical shift refers to tetramethylsilane by using adamantane as an external reference. The 'Li chemical shifts are given with respect to solid LiCl.
- 2.4. XPS. XPS measurements were carried out with a Kratos Axis Ultra spectrometer using focused monochromatized Al Ka radiation (h $\nu = 1486.6 \text{ eV}$). For the Ag $3d_{5/2}$ line, the full width at half-maximum (fwhm) was 0.58 eV under the recording conditions. The analyzed area of the samples was $300 \times$ 700 μ m². Peaks were recorded with a constant pass energy of 20 eV. The pressure in the analysis chamber was ca. 5×10^{-5} To prevent the samples from moisture/air exposure on the analysis site, the XPS spectrometer was directly connected through a transfer chamber to a nitrogen drybox. Short acquisition time control spectra were recorded at the beginning and at the end of each experiment to check the nondegradation of the samples. The binding energy scale was calibrated from the carbon contamination using the C 1s peak at 285.0 eV. Core peaks were analyzed using a nonlinear Shirley-type background.²¹ The peak positions and areas were optimized with a weighted least-squares fitting method using 70% Gaussian/30% Lorentzian lineshapes. Quantification was performed on the basis of Scofield's relative sensitivity factors.

3. Structure Solution and Refinement

3.1. Structure Solution from Single-Crystal Data. A reciprocal lattice observed from both single crystals

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Table 1. Parameters of Data Collection and Treatment for Single-Crystal and Powder Sample of Li_{5.5}Ce₁₂F₅₀

		powde	er used for lattice determination	
formula molar mass (g/mol) space group		Li _{5.5} Ce ₁₂ F ₅₀ 2669.61 P2 ₁		
a [Å]	8.855(2)	-		
b [Å]	22.968(5)	8.82	2391(7)	
	· /	22.9	188(2)	
c [Å]	8.855(2)	8 85	384(7)	
β [deg]	90.0(-)			
$V[\mathring{A}^3]$	1800.94(2)	90.6	093(2)	
V [A]	1800.94(2)	1790	0.44(3)	
Z		2		
wavelength [Å]	0.71073 (Mo Kα)			0.37504
$\mu [\text{mm}^{-1}]$	8.63	0.4	2	1.40
cryst size [mm]	$0.103 \times 0.095 \times 0.060$	0.2		glass capillary diameter [mm]
absorption correction density (calcd) [g cm ⁻³]	numerical from crystal shape and size 4.92		an	alytical for cylindrical shape 4.95
data collection	Stoe IPDS II		SI	NBL powder diffractometer
detector distance [mm]	100	0.0		2θ step [deg]
exposure time [min]	3	0.:		time/step [s]
ω range; increment [deg]	0-180; 1.0			
2θ interval [deg]	2.29-59.53			1.0-17.5
range in hkl	$h = \pm 12; -28 < k < +31; l = \pm 12$			
reflns measured	15972			
reflns unique	8776			
R _{int}	0.042			
reflus with $I > 4\sigma(I)$	$6312 (R_{\sigma} = 0.039)$			
data/params structure solution	19.3 SIR92 ⁸ SHELXS-97 (Sheldrick, 1997a)			
structure refinement	SHELXL-97 (Sheldrick, 1997a) SHELXL-97 (Sheldrick, 1997b)			FullProf.2k 10
S	1.51	13.	Q	χ^2
R_F (all reflns)	0.075	0.04		$R_{\rm B}$ (all reflections)
R_{wF}^{2} (all reflns)	0.142	0.0		$R_{\rm wp}$ (background corrected)
extinction coefficient	0.00049(3)	0.0	-	
residual in difference	-3.43, 2.63			

indicates a tetragonal lattice with a = 8.855 (2) Å and c =22.968(5) A. Analysis of the diffraction data set underlines a 91% probability of a centrosymmetric structure with the 4/m Laue symmetry ($R_{int} = 0.061$) and a primitive lattice. Automatic analysis of the extinction conditions using the Autostart Module of WinGX²³ did not lead to the determination of the space group. The first attempt to solve the structure was performed in the $P4_2/n$ space group. The starting atomic parameters of heavy cerium atoms were deduced from an automatic interpretation of direct methods with SIR92.24 Fluorine atoms were subsequently located from difference Fourier maps, and the structure was refined using SHELXL97.25 The solution obtained in the $P4_2/n$ space group seemed to be correct (with R = 0.197), but refinement did not decrease below $R_{\rm F} = 0.159$. Lithium atoms were not located in the structure by Fourier difference maps, and the refined composition corresponded to CeF₄ (all cerium atoms in an eight-fluorine coordination with a square-based antiprism neighboring). Refinement of the solution obtained with the $P\overline{4}$ space group brought a weak improvement with an $\hat{R}_{\rm F} = 0.154$ and a refined composition of CeF_{4.167}. Nevertheless, lithium atoms were not located,

electron-density map

and all cerium atoms presented a disordered nine-fluorine coordination. Finally, the best solution was found in the $P4_1$ space group, and refinement led to $R_F = 0.083$, $R_{\rm wF} = 0.155$, and S = 1.49. Lithium atoms were located by Fourier difference maps. The structure refinement was acceptable with a refined composition of Li₂Ce₆F₂₅, with two-thirds of cerium atoms coordinated in 8-fold coordination and one-third in 9-fold coordination. Because of the presence of some observed but normally systematically absent reflections, and the presence of numerous inconsistent equivalents, a supplementary resolution work was performed in monoclinic 2/m Laue symmetry ($R_{int} =$ 0.053). After transformation of the diffraction data set in a monoclinic lattice with unique axis b (instead of the measured \vec{c}), the Autostart Module of WinGX²³ assigned the centrosymmetric $P2_1/c$ space group, but solution in this $P2_1/c$ symmetry was not acceptable. Finally, the structure of this new fluoride compound was solved and refined in the noncentrosymmetric $P2_1$ space group with a final $R_{\rm F} = 0.050$ and a final refined composition of Li_{5.5}Ce₁₂F₅₀, indicating the presence of the Ce⁴⁺/Ce³⁺ mixed valence.

3.2. Synchrotron Powder Diffraction. Synchrotron powder diffraction data have been recorded on a freshly prepared powdered sample in order to check the monoclinic symmetry of the lattice. Numerous diffraction line splittings were observed, clearly indicating a lowering of the previously supposed tetragonal lattice. Figure 1 shows the synchrotron powder diffraction data and Rietveld

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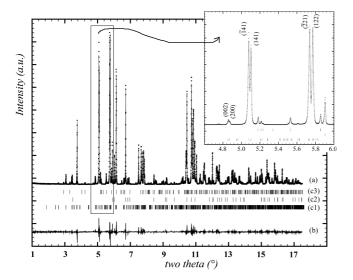


Figure 1. Rietveld plot of the fluoride sample ($R_{\rm wp} = 0.117$ and $\chi^2 =$ 13.7 for global contributions, $R_{\text{Bragg}} = 0.040$ and $R_{\text{F}} = 0.046$ for the new fluoride phase). Observed (a, dots) and calculated (a, solid line) synchrotron (SNBL) powder diffraction pattern ($\lambda = 0.37504 \,\text{Å}$) are shown with a difference curve (b) and Bragg peak positions for Li_{2+x}Ce₁₂F₅₀ (c1; 83(1) refined weight %) and impurities CeF₃ (c2) and CeF₄ (c3). The inset shows the splitting from the monoclinic distortion of the lattice.

plot performed with FullProf²⁶ by using the atomic positions from the single-crystal data analysis (P2₁ symmetry). Refined monoclinic lattice parameters are a = 8.82391(7) A, b = 22.9188(2) A, c = 8.85384(7) A,and $\beta = 90.6093(2)^{\circ}$. Cerium trifuoride CeF₃ and cerium tetrafluoride CeF₄ impurities are present in the sample with, respectively, 13(1) and 4(1) refined wt %.

3.3. Structure Refinement. Structure parameters of Li_{5.5}Ce₁₂F₅₀ have been refined in the monoclinic P2₁ space group using single-crystal data. All relevant details concerning data collection and treatment are listed in Table 1. Structure solution has indicated 12 independent cerium positions. A total of 50 independent fluorine positions were subsequently easily discovered with Fourier difference maps. All of these atomic coordinates were refined (using full-matrix least-squares refinement on F^2) with anisotropic atomic displacement parameters for cerium atoms. Careful analysis of new Fourier difference maps, in accordance with geometric considerations, allowed the localization of lithium atoms. This led to six refined independent lithium atomic positions. In the final runs of refinement, two fluorine atomic displacement parameters were constrained to $U_{\rm iso} = 0.005 \, \text{Å}^2$ because of an unrealistic quasi-zero refined value (F09 and F16). Due to similar geometric environments, Li1 and Li2 sites have been restrained to have the same atomic displacement parameter, as have the four similar Li3, Li4, Li5, and Li6 sites. Occupancy parameters have been checked for all of the atomic positions of the structure. The occupancy of the lithium position Li4 only deviated significantly from the unity at a value close to half occupancy. So, in the final run, its occupancy was fixed at 1/2, while all of the other sites were refined with the ideal occupancy corresponding to the composition Li_{5.5}Ce₁₂F₅₀ (Z = 2). The positional parameters of

the 68 independent atomic positions of the structure are listed in Table SI1 (in the Supporting Information), and the anisotropic displacement parameters of cerium atoms are given in Table SI2 (in the Supporting Information). In the final run, 328 independent parameters were allowed to vary. Selected interatomic distances are listed in Table 2.

4. Discussion

4.1. Structure Description. The structure of Li_{5,5}-Ce₁₂F₅₀ is built from stacking of the cerium polyhedra with lithium cations inserted in cavities and channels. Figure 2 shows a general view of the structure along the [101] direction, and the projections of four details along the b monoclinic axis. Cerium atoms present two coordination numbers: 8-fold coordinated by fluorine for eight Ce sites (CN8 for Ce81 to Ce88 with a square antiprism polyhedron) and 9-fold coordinated by fluorine for four Ce sites (CN9 for Ce91 to Ce94 with a monocapped square antiprism polyhedron). Interatomic Ce-F distances, indicated in Table 2, are quite homogeneous around each Ce site: 2.20 Å $\leq d_{\text{Ce81-F}} \leq 2.35$ Å, 2.27 Å $\leq d_{\text{Ce82-F}} \leq 2.39 \text{ Å}, 2.22 \text{ Å} \leq d_{\text{Ce83-F}} \leq 2.36 \text{ Å}, 2.29 \text{ Å}$ $\leq d_{\text{Ce84-F}} \leq 2.45 \,\text{Å}, 2.09 \,\text{Å} \leq d_{\text{Ce85-F}} \leq 2.31 \,\text{Å}, 2.17 \,\text{Å} \leq$ $d_{\text{Ce86-F}} \le 2.36 \text{ Å}, 2.10 \text{ Å} \le d_{\text{Ce87-F}} \le 2.32 \text{ Å}, 2.15 \text{ Å} \le$ $d_{\text{Ce88-F}} \le 2.31 \text{ Å}, 2.12 \text{ Å} \le d_{\text{Ce91-F}} \le 2.41 \text{ Å}, 2.22 \text{ Å}$ $\leq d_{\text{Ce92-F}} \leq 2.45 \text{ Å}, 2.14 \text{ Å} \leq d_{\text{Ce93-F}} \leq 2.37 \text{ Å}, \text{ and } 2.22$ $\rm \mathring{A} \leq d_{Ce94-F} \leq 2.48 \ \mathring{A}$. Nevertheless, average Ce-F distances around each Ce site, or the volume of each Ce polyhedron, show three cases: Ce91 to Ce92 present a similar behavior corresponding to 9-fold Ce⁴⁺, Ce85 to Ce88 present a similar behavior corresponding to 8-fold Ce⁴⁺, and Ce81 to Ce84 present a gradual evolution (sequence Ce81, Ce83, Ce82, Ce84) corresponding to an 8-fold mixed valence (see Table 2 and Figure 3b). The calculated cerium cationic radius (considering the average Ce-F distances and an anionic radius of 1.285 Å for F^{-27}) agrees fairly well with these observations: around 0.96 Å for CN8 Ce⁴⁺, around 1.01 Å for CN9 Ce⁴⁺, and between 0.97 Å and 1.14 Å for CN8 mixed Ce³⁺/Ce⁴⁺ (Figure 3a). These crystal chemical indications of the mixed valence correlate with the refined composition $(\text{Li}_{11}\text{Ce}_7^{\ 3+}\text{Ce}_{17}^{\ 4+}\text{F}_{100} \text{ per unit cell})$ and the orange color of our crystals. Bond valence sums (BVS²⁸) have been calculated for cerium sites by considering the values of 2.036, respectively 1.995, for the bond valence parameter R_{ij} assuming Ce³⁺, respectively Ce⁴⁺ (Table 2). Similar BVS calculations have been performed on CeF₃²⁹ and CeF₄³⁰ structures to validate the method. The calculated valences correlate well with our previous observations: tetravalent Ce⁴⁺ cations are located in the four 9-fold coordinated Ce91 to Ce94 sites and in the four 8-fold coordinated Ce85 to Ce88 sites; mixed Ce³⁺/Ce⁴⁺ valences are located in the four 8-fold coordinated Ce81 to Ce84 sites. Polyhedra volumes, calculated cationic radii, and BVS calculations for the two Ce82 and Ce84 sites

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 Table 2. Selected Interatomic Distances around Cerium Atoms, Polyhedron Volumes, and Bond Valence Sum Assuming Trivalent and Tetravalent Cerium Cations

						VS	
central atom	fluoride ligands	distance (Å)	CN^a	polyhedron volume (Å ³)	Ce ³⁺	Ce ⁴⁺	valence
Ce81 ^b	F37	2.20(2)			0.642	0.575	
	F47	2.20(1)			0.642	0.575	
	F46	2.24(2)			0.576	0.516	
	F48	2.25(2)	8	20.47	0.561	0.502	III/IV
	F27	2.31(2)			0.477	0.427	
	F04	2.33(2)			0.452	0.404	
	F25 F42	2.34(2) 2.35(2)			0.440 0.428	0.394 0.383	
	<ce81-f></ce81-f>	2.33(2)			4.22	3.77	
Ce82	F43	2.27(1)			0.531	0.476	
0002	F29	2.28(1)			0.517	0.463	
	F18	2.29(1)			0.503	0.451	
	F26	2.32(2)	8	22.13	0.464	0.415	III
	F38	2.36(1)			0.417	0.373	
	F07	2.37(1)			0.405	0.363	
	F10	2.38(1)			0.395	0.353	
	F12	2.39(1)			0.384	0.344	
$a \circ a^h$	<ce82-f></ce82-f>	2.33			3.62	3.24	
Ce83 ^b	F45	2.22(2)			0.608	0.544	
	F35 F20	2.23(1)			0.592	0.530	
	F44	2.25(1) 2.29(1)	8	20.77	0.561 0.503	0.502 0.451	III/IV
	F19	2.33(2)	o	20.77	0.303	0.404	111/1 V
	F17	2.35(2)			0.428	0.383	
	F21	2.36(2)			0.417	0.373	
	F15	2.36(2)			0.417	0.373	
	<Ce83 $-$ F $>$	2.30			3.98	3.56	
Ce84	F41	2.29(2)			0.503	0.451	
	F33	2.32(2)			0.464	0.415	
	F24	2.32(2)			0.464	0.415	
	F40	2.32(2)	8	22.91	0.464	0.415	III
	F39	2.38(1)			0.395	0.353	
	F13	2.40(1)			0.374	0.335	
	F11	2.45(1)			0.327	0.292	
	F08 < Ce84-F>	2.45(1) 2.37			0.327 3.32	0.292 2.97	
Ce85	F11	2.09(1)	8	19.49	0.864	0.774	IV
CC63	F28	2.17(1)	0	19.49	0.696	0.623	1 V
	F16	2.24(1)			0.576	0.516	
	F29	2.25(2)			0.561	0.502	
	F46	2.25(1)			0.561	0.502	
	F03	2.29(2)			0.503	0.451	
	F22	2.31(1)			0.477	0.427	
	F19	2.31(2)			0.477	0.427	
	<Ce85 $-$ F $>$	2.24			4.72	4.22	
Ce86	F22	2.17(1)			0.696	0.623	
	F13	2.20(1)			0.642	0.575	
	F43	2.24(1)	0	10.02	0.576	0.516	13.7
	F48 F09	2.25(2) 2.25(1)	8	19.93	0.561 0.561	0.502 0.502	IV
	F30	2.28(1)			0.517	0.302	
	F15	2.28(2)			0.517	0.463	
	F05	2.36(2)			0.417	0.373	
	<ce86-f></ce86-f>	2.25			4.49	4.02	
Ce87	F10	2.10(2)			0.841	0.753	
	F32	2.21(1)			0.625	0.559	
	F41	2.21(2)			0.625	0.559	
	F20	2.23(1)	8	19.30	0.592	0.530	IV
	F04	2.24(2)			0.576	0.516	
	F14	2.28(1)			0.517	0.463	
	F23	2.30(1)			0.490	0.439	
	F01	2.32(2)			0.464	0.415	
Ce88	<ce87-f></ce87-f>	2.24			4.73	4.23	
	F23	2.15(1)			0.735	0.658	
	F07 F33	2.20(2)			0.642 0.608	0.575	
	F33 F44	2.22(2) 2.25(1)	8	19.39	0.561	0.544 0.502	IV
	F06	2.25(1)	o	17.37	0.561	0.502	1 4
	F25	2.26(2)			0.546	0.302	
	F31	2.27(1)			0.531	0.476	
	F02	2.31(2)			0.477	0.427	
					J /		

Table 2. Continued

$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	/S	valence
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	Ce ⁴⁺	
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	0.713	
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	0.658	
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	0.476	
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	0.427	
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	0.404	IV
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	0.383	
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	0.353	
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	0.335	
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	0.326	
Ce92 F18 2.22(1) 9 24.33 0.608 F34 2.23(1) 0.592 F39 2.24(1) 0.576 F17 2.24(2) 0.576 F28 2.28(1) 0.517 F05 2.30(2) 0.490 F47 2.33(1) 0.464	4.07	
F34 2.23 (1) 0.592 F39 2.24 (1) 0.576 F17 2.24 (2) 0.576 F28 2.28 (1) 0.517 F05 2.30 (2) 0.490 F47 2.33 (1) 0.464	0.544	IV
F39 2.24(1) 0.576 F17 2.24(2) 0.576 F28 2.28(1) 0.517 F05 2.30(2) 0.490 F47 2.33(1) 0.464	0.530	1 1
F17 2.24(2) 0.576 F28 2.28(1) 0.517 F05 2.30(2) 0.490 F47 2.33(1) 0.464	0.516	
F28 2.28(1) 0.517 F05 2.30(2) 0.490 F47 2.33(1) 0.464	0.516	
F05 2.30(2) 0.490 F47 2.33(1) 0.464	0.463	
F47 2.33 (i) 0.464	0.439	
	0.415	
F09 2.43 (1) 0.345	0.309	
F49 2.45 (1) 0.327	0.292	
<ce92-f> 2.30 4.50</ce92-f>	4.02	
Ce93 F31 2.14(1) 0.755	0.676	
F12 2.14(1) 0.755	0.676	
F42 2.23(2) 0.592	0.530	
F01 2.31 (2) 0.477	0.427	
F40 2.33(2) 9 23.75 0.452	0.404	IV
F35 2.33 (1) 25.75 0.452	0.404	1 4
F50 2.36(2) 0.417	0.373	
F36 2.37(1) 0.405	0.363	
F14 2.37(1) 0.405	0.363	
<ce93-f> 2.29 4.71</ce93-f>	4.22	
Ce94 F27 2.22(2) 0.608	0.544	
F36 2.22(1) 0.608	0.544	
F24 2.24(2) 0.576	0.516	
F38 2.24(1) 0.576	0.516	
F32 2.26(1) 9 23.82 0.546	0.489	IV
F45 2.29 (2) 25.82 0.340 F45	0.451	1 4
F02 2.29 (2) 0.503	0.451	
F06 2.40(1) 0.303	0.431	
F50 2.48(1) 0.301 <ce94-f> 2.29 4.60</ce94-f>	0.270 4.11	

^a Cerium coordination number. ^b Mixed valence cerium sites.

correspond to a trivalent Ce^{3+} cation (by comparison of the BVS values for Ce^{4+} and Ce^{3+} in Table 2).

The connection of the 8- and 9-fold coordinated polyedra forms an opened tridimentional framework. The structure is composed of a stacking of two kinds of slabs along the monoclinic axis. One kind of slab (labeled B and D in Figure 2 composed of Ce81 and Ce82 and of Ce83 and Ce84 polyhedra) is opened due to the presence of perpendicular channels along the [101] and [101] directions. Two-thirds of the lithium sites (Li3 and Li4 in slab B and Li5 and Li6 in slab D) are located at the crossing of this channel network. These lithium cations seem to be relatively mobile along the channels. The second kind of slab (labeled A and C in Figure 2, respectively composed by Ce85, Ce86, Ce91, and Ce92 and Ce87, Ce88, Ce93, and Ce94 polyhedra) is compact and contains locked lithium cations in the Li1 and Li2 sites, respectively in slab A and C. Off-center Li1 and Li2 sites are 8-foldcoordinated by fluorides. Compact A and C slabs are composed of tetravalent Ce⁴⁺ cations only. Each of these Ce polyhedra (Ce85 to Ce88 and Ce91 to Ce94) are linked to seven neighboring polyhedra (mainly cornerbut also edge-sharing). The mixed valence cerium sites (Ce81 to Ce84) fully compose the opened B and D slabs. These cerium sites are in the vicinity of the mobile Li sites (Li3 to Li6). A long-range ordering of Ce³⁺ and Ce⁴⁺ sites is then present in Li_{1+x}Ce₆F₂₅ (i.e., with a preferential location of Ce³⁺ in the four Ce81 to Ce84 sites when Li⁺ cations are intercalated), as already observed in the Tb-based definite mixed-valence compounds $\mathrm{KTb_3F_{12}}^{10,11}$ and $\mathrm{K_2Tb_4F_{17}}^{12}$ An explicit formula of the studied compound is $\mathrm{Li_{2+x}Ce_x}^{3+}\mathrm{Ce_{12-x}}^{4+}\mathrm{F_{50}}$, indicating a minimum amount of lithium (two per unit formulas) and a progressive reduction of cerium (from Ce⁴⁺ to Ce³⁺) when intercalating the x supplementary lithium cations.

The structure of $\text{Li}_{5.5}\text{Ce}_{12}\text{F}_{50}$ is related to CeF_4 (monoclinic C2/c symmetry²⁰). The CeF_4 structure presents a succession of opened and locked slabs also. "Opened" slabs in CeF₄ show similarity with the opened slabs B and D in Li_{5.5}Ce₁₂F₅₀. Locked slabs are different. They are composed of 8-fold coordinated Ce atoms only. Adjacent locked slabs are corner-sharing across the opened slab region, blocking the "opened" slabs. The Li₅ 5Ce₁₂F₅₀ structure is a rearrangement of the CeF₄ structure, allowing the insertion of lithium cations into channels of the opened slabs. Large similitude with the

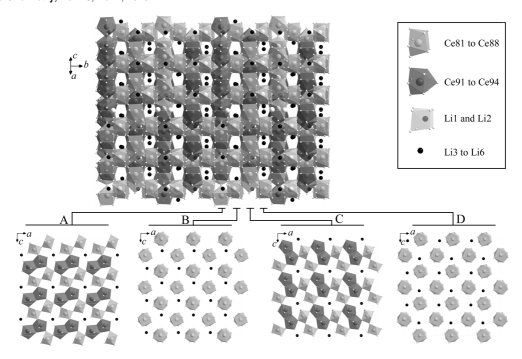


Figure 2. General representation (along [101]) and details (along [010]) of the Li_{5.5}Ce₁₂F₅₀ structure. Dark polyhedra and light polyhedra correspond respectively to cerium atoms with a CN9 and CN8 coordination number. Small white, large light gray, dark gray, and black balls correspond respectively to fluoride, cerium with CN8, cerium with CN9, and lithium atoms. Drawings A, B, C, and D are four details corresponding to the four independent structural slabs constituting the whole structure. The inset in the right top of the figure shows the fluoride neighboring of cations (coordination of lithium atoms, distorted cubic environment for Li1 and Li2, and bonds for Li3 to Li6 are not represented in the general and detailed views).

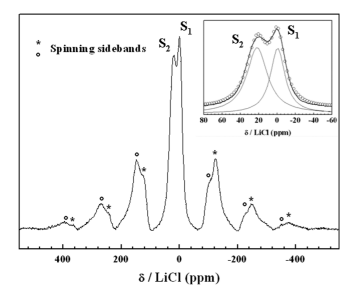


Figure 3. ⁷Li MAS NMR spectrum of the Li_{5.5}Ce₁₂F₅₀ compound (spinning speed of 14 kHz, O and * mark the spinning sidebands).

thorium fluoride $Th_6F_{24} \cdot H_2O^{16}$ is evidenced. The hydrate compound $Th_6F_{24} \cdot H_2O$, I4/m symmetry, shows a similar network with all thorium cations 8-fold-coordinated (square antiprism) and water molecules located into the channel network of the opened slabs, at the location of the mobile lithium cations in the Li_{5.5}Ce₁₂F₅₀ structure.

The effective formation of a cerium mixed-valence fluoride is mainly supported by the refined Li⁺ amount (which presents a large uncertainty due to its extremely weak scattering contrast, especially in the presence of cerium) and by crystal chemistry consideration (which are

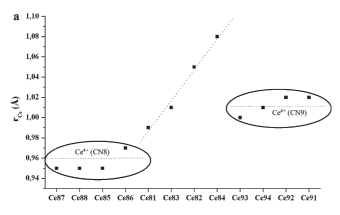
good indicators, but, in the present case, cannot be considered as definitive). The presence of two different environments for lithium cations has been checked with ⁷Li MAS NMR spectroscopy, and the simultaneous presence of Ce³⁺ and Ce⁴⁺ has been investigated using XPS analysis (which allows qualitative and quantitative

4.2. ⁷Li NMR MAS Spectroscopy. Figure 4 displays the ⁷Li NMR spectrum of the Li_{5.5}Ce₁₂F₅₀ compound. The NMR spectrum exhibits two peaks centered at 0 and 21.1 ppm noted S_1 and S_2 , respectively, indicating the presence of two types of lithium ions. Although two types of lithium ions with different environments are highlighted by the structural study, the significant separation of the NMR lines is probably ascribable to the presence of Ce³⁺ paramagnetic ions, which results in the shift of the S₂ line. The paramagnetism effect on the chemical shifts has been discussed for lithium in several works, in particular for various compositions of the Li(Ni,Co)O₂ solid solution³¹ and for nonstoichiometric LiCoO₂,³² where the presence of the Ni³⁺ or Co²⁺ cations in the low spin state is accompanied by the appearance of new resonance lines in addition to the line at 0 ppm. Paramagnetic "external" species such as adsorbed oxygen molecules result in a shift for zeolites with a high silica content.³³ Considering the location of the Li⁺ ions in the Li_{5.5}- $Ce_{12}F_{50}$ structure, two-thirds of them (Li3, Li4, Li5, and Li6) are localized in the B and D layers, which contain

⁽³¹⁾ Marichal, C.; Hirschinger, J.; Granger, P.; Menetrier, M.; Rougier, A.; Delmas, C. Inorg. Chem. 1995, 34, 1773–1778.

⁽³²⁾ Levasseur, S.; Menetrier, M.; Suard, E; Delmas, C. Solid State Ionics 2000, 128, 11-24.

⁽³³⁾ Accardi, R. J.; Lobo, R. F. Microporous Mesoporous Mater. 2000, 40, 25 - 34.



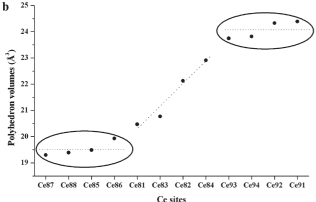


Figure 4. Environments of the 12 independent Ce sites: Calculated cerium ionic radius (a) and polyhedron volumes (b).

Ce³⁺ trivalent cerium ions. The other Li⁺-including Li1 and Li2 ions are in the A and C layers, which are exclusively built with tetravalent cerium ions. One may assume that only the nuclear resonance of the first type of lithium is affected by the presence of paramagnetic Ce³⁺ because these ions are located within channels delimited by the mixed Ce³⁺/Ce⁴⁺ ions in statistical distribution. In addition to the "paramagnetic" shift, the line width is also affected in the case of the S_2 line with a broadening; the fwhm's are equal to 2220 and 3170 Hz for S_1 and S_2 , respectively.

Taking into account the assignment of the S_1 and S_2 lines and the number of involved lithium ions, the theoretical ratio of their integrated surfaces should be equal to 2:3.5, that is, 0.57. Indeed, even if the lithium nucleus is quadrupolar, integrated surfaces of the lines are almost proportional to the relative concentration of the Li⁺ cations. The experimental ratio of 0.69 has been obtained by considering two Lorentzian lines for the fitting. The weak difference between the theoretical and experimental values could be explained by the particular structure of the compound. Indeed, the opened structure of Li_{5.5-} Ce₁₂F₅₀ fluoride can be compared to an intercalation compound with stoichiometry $\text{Li}_{2+x}\text{Ce}_x^{3+}\text{Ce}_{12-x}^{4+}\text{F}_{50}$, where x can vary in large proportions. It is worthy of note that ¹H MAS NMR confirms the absence of water molecules within the opened channels of the Li_{5.5}Ce₁₂F₅₀ structure. As a matter of fact, no significant signal was obtained for its spectrum (not shown here).

4.3. X-Ray Photoelectron Spectroscopy (XPS). The more relevant peak to identify the simultaneous presence of Ce³⁺ and Ce⁴⁺ is the Ce3d core peak, located in the

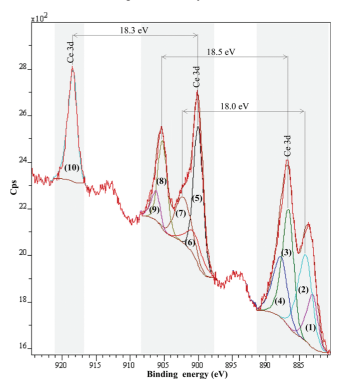


Figure 5. XPS Ce3d core level peaks of Li_{5.5}Ce₁₂F₅₀.

Table 3. Labeling and Mean Positions of the 10 Components of Li_{5.5}Ce₁₂F₅₀

		spin-orbit	characteristic	binding energy (eV)
	1	3d _{5/2}	shake down of 3	883.1
Ce ³⁺	6	$3d_{3/2}$	shake down of 8	900.5
	3	$3d_{5/2}$		886.7
	8	$3d_{3/2}$		905.2
	2	$3d_{5/2}$		884.2
	7	$3d_{3/2}$		902.2
Ce ⁴⁺	4	$3d_{5/2}$	satellite	887.7
	9	$3d_{3/2}$	satellite	906.2
	5	$3d_{5/2}$	satellite	900.0
	10	$3d_{3/2}$	satellite	918.4

energetic window (Binding Energy), 880-920 eV. Note that XPS spectra of cerium compounds are known to exhibit rather complex features due to shake-up or -down lines, multiplet splitting or hybridization with ligand orbitals, and fractional occupancy of the valence 4f orbitals.³⁴ The Ce3d XPS spectrum of Li_{5.5}Ce₁₂F₅₀ (Figure 5) exhibits 10 components (labeled from 1 to 10) with the assignment of each component defined in Table 3. The theoretical basis for all of the components has been reported in the literature.³⁵

Four peaks labeled 1 and 3 (for $3d_{5/2}$) and 6 and 8 (for $3d_{3/2}$) are related to the Ce³⁺ oxidation state. The satellite components 1 and 6 are related to the ligand-to-metal charge transfer phenomenon by the primary photoionization process. Six peaks labeled 2, 4, and 5 (for $3d_{5/2}$) and 7, 9, and 10 (for 3d_{3/2}) referring to the three pairs of

⁽³⁴⁾ Shyu, J. Z.; Otto, K.; Watkins, W. L. H.; Graham, G. W.; Belitz, R. K.; Gandhi, H. S. J. Catal. 1988, 114, 23-33.

^{(35) (}a) Fujimori, A. J. Magn. Magn. Mater. 1985, 47-48, 243-247. (b) Allen, J. W. J. Magn. Magn. Mater. 1985, 47-48, 168-174. (c) Fujimori, A. Phys. Rev. B 1983, 28, 2281-2283. (d) Fujimori, A. Phys. Rev. B 1983, 27, 3992-4001.

spin-orbit doublets are characteristic of Ce⁴⁺ final states. ^{36,37} The literature reports that the peak labeled 10 arises exclusively from Ce⁴⁺ due to a transition from the 4f⁰ initial state to the 4f⁰ final state. This fact is pretty interesting, as this component is isolated from the others and located around 918 eV. Then, this peak could be used as a quantitative probe of the amount of Ce⁴⁺. ^{36,38,39} Indeed, it has been demonstrated that the integral area of the peak labeled 10 with respect to the total Ce3d area could be translated into the percentage of Ce⁴⁺. The mean percentage of Ce⁴⁺ can be then calculated through the following relation:

$$Ce^{4+}\% = \frac{(10)\%}{14} \times 100$$

where (10)% is the percentage of the peak area corresponding to the component 10 with respect to the total Ce3d area. By using this formula, the experimental ratio Ce^{4+}/Ce^{3+} is equal to 1.75. By taking into account the refined weight amounts (13% for CeF₃ and 4% for CeF₄), we can reach the calculated formula $\text{Li}_{5,2}\text{Ce}_{3,2}^{3+}\text{Ce}_{8,8}^{4+}\text{F}_{50}$, very close to the refined one ($\text{Li}_{5,5}\text{Ce}_{3,5}^{3+}\text{Ce}_{8,5}^{4+}\text{F}_{50}$).

5. Conclusion

The structure of Li_{5.5}Ce₁₂F₅₀ has been successfully solved, refined, and described by combining single-crystal X-ray

diffraction and high-resolution synchrotron powder diffraction data. The synchrotron powder diffraction pattern has been essential in determining the effective monoclinic $P2_1$ symmetry of the lattice, due to a weak distortion of a pseudotetragonal lattice. The structure of $Li_{5.5}Ce_{12}F_{50}$ is related to the structure of CeF_4 , ¹⁴ with the intercalation of lithium cations allowed by opening a channel network. One part of Li⁺ is locked, and the other part is relatively free to move along the channel network. The insertion of Li⁺ into the channel leads to the reduction of one equivalent part of Ce⁴⁺ to Ce³⁺. This is the first mixed-valence cerium fluoride. A long-range ordering of trivalent and tetravalent cerium cations has been observed. Structural details deduced from the crystal chemistry have been proven by spectroscopic analyses. The two kinds of Li⁺ environments have been clearly identified by ⁷Li NMR spectroscopy. And the presence of both Ce³⁺ and Ce⁴⁺ valences has been clearly established by XPS. The observed Ce⁴⁺/Ce³⁺ ratio is in perfect agreement with the previously refined Li_{5.5}Ce₁₂F₅₀ composition. The structure description indicates that Li_{1+x} Ce₁₂F₅₀ is a good candidate to realize the electrochemical intercalation-disintercalation of Li⁺ in order to change and to control the Ce³⁺/Ce⁴⁺ ratio. Investigations are in progress.

Supporting Information Available: (Table SI1) Atomic positional and displacement parameters for Li_{5.5}Ce₁₂F₅₀. The equivalent isotropic displacement parameter $U_{\rm eq}$, defined as one-third of the trace of the orthogonalized U_{ii} tensor, is given for cerium positions. (Table SI2) Anisotropic displacement parameters (pm²) for cerium atoms in Li_{5.5}Ce₁₂F₅₀. This material is available free of charge via the Internet at http:// pubs.acs.org.

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