# Synthesis and Characterization of Two New Fluoroplumbate(II) Complexes: Tetrabutylammonium Fluorodihaloplumbate, (But)<sub>4</sub>N[PbX<sub>2</sub>F] (X = Cl, I)

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Tetrabutylammonium Fluorodichloroplumbate(II),  $N(C_4H_9)_4[PbCl_2F]$ , TBAFDiCP and Tetrabutylammonium Fluorodiiodoplumbate(II),  $[(C_4H_9)_4N][PbI_2F]$ , TBAFDiIP are the first examples of fluoroplumbate salts that have been prepared from the reaction of  $(C_4H_9)_4NF$  with  $PbCl_2$  and  $PbI_2$  respectively using either  $CH_3CN$  solvent. These new compound characterized by elemental analysis, IR, UV/Visible,  $^1H$  NMR, and  $^{19}F$  NMR techniques.

Key Words: Tetrabutylammonium fluoride. Lead complexes. Fluoride addition, Fluorocomplexes, Synthesis

### Introduction

The unique properties of fluorine impart an unusual reactivity to the metal-fluorine bond which can be exploited in preparative inorganic chemistry or in catalysis.<sup>1,2</sup> In addition, the development of main metal mediated M-F bond formation processes is still a virtually unexplored field. In recent years there has been a great deal of interest in the metal fluorocompounds. Particularly the fluorides have been subject of an intense scientific discussion since the first synthesis of such a compound. The important prerequisites for a fluorinating agent are mildness, versatility, selectivity and operational simplicity.3-6 One of the most famous fluorinating agent is tetramethylammonium fluoride. The main reaction for its synthesis is the reaction of Tetramethylammonium hydroxide with hydrofluoric acid. By using this reagent many fluorocompounds of main group elements produced such as: (CH<sub>3</sub>)<sub>4</sub>N[PF<sub>4</sub>].<sup>7</sup> (CH<sub>3</sub>)<sub>4</sub>N[SeF<sub>5</sub>]. (CH<sub>3</sub>)<sub>4</sub>N-[TeF<sub>5</sub>].8 (CH<sub>3</sub>)<sub>4</sub>N[IF<sub>8</sub>].9 and in fewer amount some of transition metal fluorocomplexes synthesized like (CH<sub>3</sub>)<sub>4</sub>N[MoO<sub>3</sub>F]. <sup>10</sup> [ReOF<sub>6</sub>]<sup>12</sup> and (CH<sub>3</sub>)<sub>4</sub>N[WO<sub>3</sub>F]. <sup>13</sup> There were two primary incentives for selection of (R)<sub>4</sub>N<sup>+</sup> as the counter ion. Firstly, quaternary ions such as Tetrabutylammonium which are often used as phase transfer catalysts. Secondly, quaternary ions such as Tetrabutylammonium are used as crystal growing agents. The numbers of lead(II) fluorocompounds are still scarce, and very few studies on their reactivity have been reported. In this paper, a direct, simple and one-step method has been used to synthesize two new lead fluorocompounds.

## **Results and Discussion**

Fluorides have been subject of many scientific discussions

firstly because of the synthetic possibility<sup>3,4</sup> and secondly, because of rare amounts of the spectroscopic data especially <sup>19</sup>F-NMR chemical shifts of these compounds.<sup>14-17</sup>

We now report the synthesis of the tetrabutylammonium fluorodihaloplumbate TBAFDiXP that are analog of the other halocompounds such as fluorochromate compounds. The method used for the synthesis does not involve direct use of HF or reaction of MHF<sub>2</sub> (M = NH<sub>4</sub>, K, Rb or Cs) with lead compounds and is based on the concept of strong action of tetrabutylammonium fluoride ( $C_4H_9$ )<sub>4</sub>NF and its power to fluoride addition to many compounds.

The advantages of the new method are:

a) there is no side product. b) the reaction is quite fast. c) mild conditions and d) the accompanied color change that providing visual means for ascertaining the progress of the reaction

 $(C_4H_9)_4N]$ [PbCl<sub>2</sub>F] was prepared by the reaction of  $(C_4H_9)_4NF$  and PbCl<sub>2</sub> in a 1:1 ratio in MeCN solvent as follows:

$$(C_4H_9)_4NF + PbCl_2 \rightarrow (C_4H_9)_4N[PbCl_2F]$$

In the vibrational spectrum of this compound the known bands of cation were seen that confirmed with literature data (Table 1). Electronic spectrum of compound shows two absorptions at 225 nm ( $\varepsilon$  = 741 mol. <sup>-1</sup>lit.cm<sup>-1</sup>) and 275 nm ( $\varepsilon$  = 241 mol. <sup>-1</sup>lit.cm<sup>-1</sup>) that belongs to A' $\rightarrow$ A' and A' $\rightarrow$ A' transition

 $(C_4H_9)_4N[PbI_2F]$  was prepared by the reaction of  $(C_4H_9)_4NF$  and  $PbI_2$  in a 1:1.1 ratio in MeCN solvent as follows:

$$(C_4H_9)_4NF + PbI_2 \rightarrow (C_4H_9)_4N[PbI_2F]$$

In the vibrational spectrum of this compound the known bands of tetrabutylammonium cation and fluoroanion were seen that confirmed with literature data (Table 2).

**Table 1**. The frequencies (cm<sup>-1</sup>) and assignment of cation and anion of (C₁H₀)₁N[PbCl₂F]

Intensity	$v(\mathrm{cm}^{-1})$	Assignment	$v(\mathrm{cm}^{-1})$	Assignment	Intensity
	$(C_4H_9)_4N^-$		1435	и <sub>15</sub> , CH <sub>2</sub> , asym.def	(w)
3445	$\nu_{\rm CH2} + \nu_{19}$	(w, br)	1395	и <sub>6</sub> , CH <sub>2</sub> , sym.str	(w)
3305	$\nu_{\mathrm{CH2}} + \nu_{8}$	(w, br)	1170	ν <sub>rock</sub> , CH2, rocking ν14	(w)
3220	ν <sub>CH2</sub> , asym.str	(sh)	876	и <sub>8</sub> , NC <sub>4</sub> , asym.str	(w, br)
3015	$\nu_{13}, \nu_{\rm CH2}, {\rm asym.str}$	(w, br)	468	ν <sub>19</sub> , NC <sub>4</sub> , def.	(ms)
2950	ν <sub>14</sub> , CH <sub>2</sub> , asym.str	(s)	446	ν <sub>19</sub> , NC <sub>4</sub> , def.	(ms)
2850	$\nu_{14}$ , $CH_2$ , asym.str	(s)	[PbCl₂F]⁻		
2770	$\nu_7 + \nu_{16}$	(w)	.122	Dt. T (A)	(m. 1-)
2390	13 + 18 + 1/16	(w)	433	Pb-F (A)	(w, br)
1995	1/8 + 1/15	(w, br)	303	Pb-C1(E)	(w)

**Table 2**. The frequencies (cm<sup>-1</sup>) and assignment of cation and anion of (C₁H₀)₁N[Pbl₂F]

$v(\mathrm{cm}^{-1})$	Assignment	Intensity	ν(cm <sup>-1</sup> )	Assignment	Intensity
	(C₄H <sub>9</sub> )₄N⁻		1463	ν <sub>15</sub> , CH <sub>2</sub> , asym.def	(s)
3435	V <sub>C H2</sub> + V <sub>19</sub>	(w, br)	1376	ν <sub>16</sub> , CH <sub>2</sub> , sym.str	(m)
3350	И <sub>С Н2</sub> + И <sub>8</sub>	(W)	1146	иоск, CH2, rocking и4	(m)
3110	ν <sub>CH2</sub> , asym.str	(sh)	873	148, NC4, asym.str	(W)
3080	ν <sub>13</sub> , ν <sub>CH2</sub> , asym.str	(W)	525	119, NC4, def.	(W)
2950	14, 16H2, asym.str	(\$)	403	119, NC4, def.	(ms)
2840	$\nu_{14}$ , $\nu_{CH2}$ , asym.str	(s)			
2590	$\nu_7 + \nu_{16}$	(w)	483	Pb-F (A)	(w, br)
2420	13 + 18 + 176	(w)			
1860	$u_8 + u_{15}$	(w, br)			

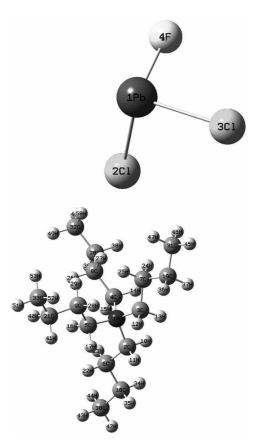


Figure 1. The calculated structure for (C<sub>4</sub>H<sub>9</sub>)<sub>4</sub>N[PbCl<sub>2</sub>F].

Table 3. A number of the theoretical frequencies (cm  $^{-1}$  ) of cation and anion of  $(C_4H_9)_4N[PbCl_2F]$ 

ν(cm <sup>-1</sup> )	Assignment	$\nu  (cm^{-1})$	Assignment
	$(C_4H_9)_4N^-$	1449	ν <sub>15</sub> , CH <sub>2</sub> , asym.def
3014	ν <sub>1</sub> , ν <sub>CH2</sub> , asym.str	1392	ν <sub>16</sub> , CH <sub>2</sub> , sym.str
2953	ν <sub>1+</sub> , CH <sub>2</sub> , asym.str	1176	ν <sub>rock</sub> , CH <sub>2</sub> , rocking ν <sub>14</sub>
2770	V7+ V16	870	v18, NC4, asym.str
2390	13 + 18 + 1/16		[PbCl <sub>2</sub> F] <sup>-</sup>
1995	18 + 1/15	433	Pb-F

For more clear characterization the theoretical calculation of vibration modes have been done by the method and basis set: B<sub>3</sub>LYP/LANL2DZ. The calculations shown that the structures of these two fluorodihaloplumbate salts, are not formed dimer, trimer, or more multi nuclear structures in solid states. The calculated data and structure of one of these fluorodihaloplumbate salts are shown in Figure 1 and Table 3.

Electronic spectrum of compound shows four absorptions at 245 nm ( $\varepsilon$ = 1330 mol. <sup>-1</sup>lit.cm<sup>-1</sup>). 320 nm ( $\varepsilon$ = 124 mol. <sup>-1</sup>lit.cm<sup>-1</sup>). 336 nm ( $\varepsilon$ = 230 mol. <sup>-1</sup>lit.cm<sup>-1</sup>) and 369 nm ( $\varepsilon$ = 226 mol. <sup>-1</sup>lit.cm<sup>-1</sup>), the first two absorptions belong to A' $\rightarrow$ A', A' $\rightarrow$ A' and two another seconds belong to A" $\rightarrow$ A' transitions. Tables 4 and 5 show the electronic transitions data of these two fluorodihaloplumbate compounds respectively. In the <sup>19</sup>F NMR of these compounds a signal is seen in the –60 ppm that confirmed the bonding between fluoride and lead.

Table 4. Transitions specifications of TBAFDiCP

$(\varepsilon, \mathbf{M}^{-1}\mathbf{cm}^{-1})$
225(741) A'→A'

Table 5. Transitions specifications of TBAFDiIP

$\hat{\lambda}_4$ $(\varepsilon, M^{-1}cm^{-1})$	$\lambda_3$ $(\varepsilon, \mathrm{M}^{-1}\mathrm{cm}^{-1})$	$\lambda_2$ $(\varepsilon, \mathrm{M}^{-1}\mathrm{cm}^{-1})$	$\lambda_{\rm l}$ ( $\varepsilon$ , ${ m M}^{-1}{ m cm}^{-1}$ )
245(1330)	320(124)	336(230)	369(226)
A'→A'	A'→A'	A"→A'	A"→A'

# Experimental

Material and instruments. Acetonitrile (Fluka, P.A.) was distilled several times from phosphorus pentaoxide before using, thereby reducing its water content to <4 ppm. Tetrabutylammonium fluoride was bought from Merck. PbX<sub>2</sub> (Merck, P.A.) were used without further purification. Solvents were purified by standard methods. Infrared spectra were recorded as KBr disks on a Shimadzu model 420 spectrophotometer. The UV/Visible measurements were made on an Uvicon model 922 spectrometer. <sup>1</sup>H and <sup>19</sup>F-NMR were recorded on a Bruker AVANCE DRX 500 spectrometer. All the chemical shifts are quoted in ppm using the high-frequency positive convention; <sup>1</sup>H NMR and <sup>19</sup>F-NMR spectra were referenced to external SiMe<sub>4</sub> and CFCl<sub>3</sub> respectively. The percent compositions of elements were obtained from the Microanalytical Laboratories. Department of Chemistry, OIRC, Tehran.

Synthesis of Tetrabutylammonium Fluorodichloroplumbate (TBAFDiCP), (C<sub>4</sub>H<sub>9</sub>)<sub>4</sub>N[PbCl<sub>2</sub>F]. To a solution of tetrabutylammonium fluoride (0.50 g. 1.9 mmol) was added a solution of lead dichloride. PbCl<sub>2</sub> (0.51 g. 1.83 mmol) in MeCN under stirring. A yellowish white precipitate was formed. After 24 hours stirring, the mixture was filtered, washed with ether and dried at room temperature. The tetrabutylammonium salts are somewhat hygroscopic, and it was better to be stored under a layer of hexane. Mp: 245 °C. Cacld. C<sub>16</sub>H<sub>36</sub>Cl<sub>2</sub>FNPb: C. 35.60; H. 6.67; N. 2.59; % found C. 36.49; H. 6.78; N. 2.61; A 0.001 mole solution conductivity is 129 Ω<sup>-1</sup>cm<sup>2</sup> mol<sup>-1</sup>. UV/Visible, IR. <sup>1</sup>H-NMR and <sup>13</sup>C-NMR were all consistent with the TBAFDiCP structure. In <sup>19</sup>F-NMR of this compound a broad peak was seen near -61 ppm.

Tetrabutylammonium fluorodichloroplumbate is soluble in ethanol and dimethyl sulphoxide (DMSO) and not soluble in dichloromethane, ether and toluene.

Synthesis of Tetrabutylammonium Fluorodiiodoplumbate (TBAFDiIP), (C<sub>4</sub>H<sub>9</sub>)<sub>4</sub>N[PbI<sub>2</sub>F]. Tetrabutylammonium fluorodiiodoplumbate (II), [(C<sub>4</sub>H<sub>9</sub>)<sub>4</sub>N][PbI<sub>2</sub>F] was prepared by dissolving PbI<sub>2</sub> (0.51 g. 1.83 mmol) in MeCN and addition of this solution to a solution of tetrabutylammonium fluoride (0.50 g. 1.9 mmol) in MeCN under stirring at room temperature until yellow precipitate was formed. After

24 hours stirring, the mixture was filtered, washed with ether and dried at room temperature. Mp: 180 °C, for  $C_{16}H_{36}FNPbI_2$ ; C. 26.58; H. 4.98; N. 1.93; % found C. 27.38; H. 5.00; N. 1.95; A 0.001 mole solution conductivity is 118  $\Omega^{-1} \text{cm}^2 \text{mol}^{-1}$ . UV/Visible, IR, <sup>1</sup>H-NMR and <sup>13</sup>C-NMR were all consistent with the TBAFDiIPstructure. Tetrabutylammonium fluorodiiodoplumbate is soluble in diethylether, metanol (DMSO), acetone and not soluble in chloroform and toluene.

#### Conclusion

Two new fluorocompounds of PbX<sub>2</sub> was synthesized simply. (C<sub>4</sub>H<sub>9</sub>)<sub>4</sub>N[PbCl<sub>2</sub>F] was prepared by the reaction of (C<sub>4</sub>H<sub>9</sub>)<sub>4</sub>NF and PbCl<sub>2</sub> and (C<sub>4</sub>H<sub>9</sub>)<sub>4</sub>N[PbI<sub>2</sub>F] was prepared by the reaction of (C<sub>4</sub>H<sub>9</sub>)<sub>4</sub>NF and PbI<sub>2</sub> in a 1:1 ratio in MeCN solvent. Electronic and vibrational spectra of these new Fluoro-complexes were studied. These compounds were characterized by elemental analysis, IR. UV/Visible, and <sup>13</sup>C-NMR, <sup>1</sup>H-NMR and <sup>19</sup>F-NMR techniques. Production of this compound shows the ability of tetrabutylammonium fluoride in fluoride addition to main group elements compounds.

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