at -196 °C and heated at 165 °C for 2.2 h. The product was checked by infrared spectroscopy, but it was found that no reaction took place. The reaction mixture was again heated at 250-260 °C for 5 h. The products were separated by trap-to-trap distillation through a trap at  $-196$  °C which retained the compound found to be  $(CF_3)_2C=N$ -N=C(CF3)2 **(0.46** mmol).

 $\equiv$ C(CF<sub>3</sub>)<sub>2</sub> (0.46 mmol).<br>**Reaction of CF<sub>2</sub>CF<sub>2</sub>CF<sub>2</sub>CF<sub>2</sub>S==NLi with CF<sub>2</sub>CF<sub>2</sub>CF<sub>2</sub>CF<sub>2</sub>S==NC-F((CF,),).** Into a 100-mL reaction vessel which retained **2.0** mmol product and  $-196$  °C which retained the compound found to be  $(CF_3)_2C=N$ .<br>  $N=C(CF_3)_2$  (0.46 mmol).<br>
Reaction of  $CF_2CF_2CF_2CF_2S=NLi$  with  $CF_2CF_2CF_2CF_2S=NC$ -<br>  $F((CF_3)_2)$ . Into a 100-mL reaction vessel which retained 2.0 mmol<br>  $CF_2CF_2S=NCF(CF_3)_2$  at -196 °C. The reaction mixture was gradually warmed to 25 °C.<sup>3</sup> After 6 h, the product was separated by trap-to-trap distillation. However, only starting material  $(\text{CF}_2\text{CF}_2\text{CF}_2\text{S}=\text{NCF}(\text{CF}_3)_2, 1.76 \text{ mmol})$  and small amounts of  $CF_2CF_2CF_2S$  were found. **2104**<br>
at -196 °C and heated at 165 °C for<br>
by infrared spectroscopy, but it was<br>
The reaction mixture was again heap<br>
products were separated by trap-tc<br>
at -196 °C which retained the com<br>
N=C(CF<sub>3</sub>)<sub>2</sub> (0.46 mmol).<br>
Re

**Acknowledgment** is expressed to the donors of the Petroleum Research Fund, administered by the American Chemical Society, and to the National Science Foundation (Grant CHE-7727395) for support of this research. We thank Charles Barinaga and Dennis Gage for mass and <sup>19</sup>F NMR spectral data.

**Registry No.**  $CF_2CF_2CF_2CF_2S=NSi(CH_3)_3$ , 77110-78-2; CHE-7727395) for support of this research. We thank Charles<br>Barinaga and Dennis Gage for mass and <sup>19</sup>F NMR spectral<br>data.<br>**Registry No.** CF<sub>2</sub>CF<sub>2</sub>CF<sub>2</sub>CF<sub>2</sub>CF<sub>2</sub>S=NS(CH<sub>3</sub>)<sub>3</sub>, 77110-78-2;<br>CF<sub>2</sub>CF<sub>2</sub>CF<sub>2</sub>CF<sub>2</sub>CF<sub>2</sub>CF<sub>2</sub>C 77110-79-3;  $CF_2CF_2CF_2CF_2S$ NSO<sub>2</sub>CF<sub>3</sub>, 77110-80-6; CF<sub>2</sub>CF<sub>2</sub>CF<sub>2</sub>CF<sub>2</sub>S=NSO<sub>2</sub>Cl, 77110-81-7; <br>  $\overline{ }$ CF<sub>2</sub>CF<sub>2</sub>CF<sub>2</sub>CF<sub>2</sub>S=NC(O)N=SCF<sub>2</sub>CF<sub>2</sub>CF<sub>2</sub>CF<sub>2</sub>, 77110-82-8;  $CF_2CF_2CF_2CF_2S-NC(O)C(O)N=SCF_2CF_2CF_2CF_2$ , 77110-83-S<del>=</del>N(CF<sub>3</sub>)<sub>2</sub>, 77110-85-1; bis(trifluoromethyl)diazirine, 3024-50-8; CF,CF2CF2CF2S=NLi, **771 10-86-2;** (CF3)2S=NLi, **61097-18-5;**  (CH3),SiC1, **75-77-4;** CF\$(O)CI, **354-32-5;** CICN, **506-77-4;**  9;  $CF_2CF_2CF_2CF_2S=NCN$ , 77110-84-0;  $(CF_3)$ <sub>2</sub>N=SC(O)C(O)- $CF_2CF_2CF_2CF_2S$ , 706-76-3;  $(CF_3)_2C=NN=CC(F_3)_2$ , 1619-84-7; **r**. CF<sub>2</sub>CF<sub>2</sub>CF<sub>2</sub>CF<sub>2</sub>S=NH, 77110-80-2; (CF<sub>3</sub>)<sub>2</sub>S=NH, 60646-40-4; *CF<sub>2</sub>CF<sub>2</sub>CF<sub>2</sub>CF<sub>2</sub>S=NH, 17110-87-3;* (CF<sub>3</sub>)<sub>2</sub>S=NH, 60646-40-4; CFjSOZF, **335-05-7;** ClS02F, **13637-84-8;** COC12, **75-44-5;** FC- (O)C(O)F, **359-40-0;** FN=C(CF3)2, **2802-70-2.** 

Contribution from Rocketdyne, a Division of Rockwell International Corporation, Canoga Park, California **91 304** 

# **Syntheses and Properties of FOIF40, CIOIF40, HOIF40, and Tetrafluoroperiodates**

KARL 0. CHRISTE,\* RICHARD D. WILSON, and CARL J. SCHACK

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Mixtures of *cis-* and trans-CsIF<sub>4</sub>O<sub>2</sub> were prepared by the interaction of CsIO<sub>4</sub> with either anhydrous HF, BrF<sub>5</sub>, ClF<sub>3</sub>, ClF<sub>3</sub>, or  $F_2$ . The vibrational spectra of these mixtures were recorded, and partial assignments are given for *cis-* and trans-IF<sub>4</sub>O<sub>2</sub>. The assignments for trans-IF<sub>4</sub>O<sub>2</sub><sup>-</sup> were supported by a normal-coordinate analysis. The CsIF<sub>4</sub>O<sub>2</sub> salt dissolves in CH<sub>3</sub>CN with the formation of  $IF_4O_2^-$  anions but undergoes solvolysis in anhydrous HF with formation of  $HOIF_4O$ . An improved synthesis of HOIF<sub>4</sub>O from CsIF<sub>4</sub>O<sub>2</sub> and BiF<sub>5</sub> in anhydrous HF is reported, and its Raman and <sup>19</sup>F NMR spectra were recorded. The interaction of CsIF<sub>4</sub>O<sub>2</sub> with NF<sub>4</sub>SbF<sub>6</sub> in anhydrous HF results in solutions containing NF<sub>4</sub><sup>+</sup>, HF<sub>2</sub><sup>-</sup>, and HOIF<sub>4</sub>O. When standing or when pumped to dryness, these mixtures decompose to yield  $NF_3$  and the new compound FOIF40 in high yield. The latter compound, the first known example of an iodine hypofluorite, was thoroughly characterized and shown by vibrational and NMR spectroscopy to **be** a mixture of the cis and trans isomers. For comparison, the vibrational spectra of IF<sub>5</sub>O have also been recorded. The reaction of CsIF<sub>4</sub>O<sub>2</sub> with ClOSO<sub>2</sub>F was shown to yield the novel compound CIOIF<sub>4</sub>O. The fluorination reactions of CsIO<sub>4</sub>, CsIF<sub>4</sub>O<sub>2</sub>, IF<sub>5</sub>O, and HOIF<sub>4</sub>O with elementary fluorine were also studied.

### **Introduction**

The number of elements known to form stable hypofluorites is very limited.' Thus stable hypofluorites are known only for carbon-, nitrogen-, sulfur-, selenium-; fluorine-, and chlorine-containing compounds. In addition, the unstable hypofluorous acid, HOF, has been prepared.<sup>2</sup> Since recent work in our laboratory had shown that the thermal decomposition of certain  $NF_4^+$  salts of oxyanions such as  $NF_4ClO_4^3$ and  $NF_4SO_3F^4$  produces the corresponding hypofluorites in high yield, it was interesting to apply this method to the synthesis of novel hypofluorites. Preliminary results<sup>5</sup> showed that FOIF40, the first known example of an iodine hypofluorite, can be prepared in this manner. In this paper, detailed information is given on the synthesis, properties, and reaction chemistry of this interesting compound and of related iodine oxyfluoride derivatives such as  $ClOIF<sub>4</sub>O$ ,  $HOIF<sub>4</sub>O$ , and the  $IF_4O_2^-$  anion.

The literature on the synthesis and properties of salts containing the  $IF_4O_2^-$  anion is scant. The first report on the existence of  $IF_4O_2^-$  salts was published in 1971 by Engelbrecht

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and co-workers<sup>6</sup> but was limited to a one-sentence statement that HOIF40 interacts with either alkali-metal fluorides or trifluoroacetates to yield the corresponding salts. In a subsequent paper,<sup>7</sup> this statement was repeated, but again no data were given. In 1975, Aubke and co-workers reported<sup>8</sup> that CsF combines with an excess of  $IF_3O_2$  to give  $Cs^+IF_4O_2^-$ . A melting point, elemental analysis, and incomplete vibrational spectra were given, which were incorrectly interpreted in terms of a cis isomer. In 1976, Selig and Elgad reported<sup>9</sup> that partial hydrolysis of  $IF<sub>7</sub>$  produces  $IF<sub>5</sub>O$ , HOIF<sub>4</sub>O, and, with increased water addition, the IF<sub>4</sub>O<sub>2</sub><sup>-</sup> anion, which was identified by <sup>19</sup>F NMR and vibrational spectroscopy as the cis isomer. Although Selig and Elgad reported only solution data, their vibrational spectra strongly disagreed with those reported by Aubke for solid  $Cs^+IF_4O_2^-$ . In 1977, Gillespie and Krasznai reported<sup>10</sup> that solutions of  $KIO<sub>4</sub>$  in  $IF<sub>5</sub>$  contain a mixture of IO<sub>2</sub>F, IOF<sub>3</sub>, and *cis*- and *trans*-IF<sub>4</sub>O<sub>2</sub><sup>-</sup>. On cooling solutions of KIO<sub>4</sub> dissolved in boiling IF<sub>5</sub>, they isolated a KIF<sub>4</sub>O<sub>2</sub>.2IF<sub>5</sub> adduct, which could be converted into  $KIF<sub>4</sub>O<sub>2</sub>$ . Both compounds were shown by <sup>19</sup>F NMR and Raman spectroscopy to

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contain in the solid state and in  $CH<sub>3</sub>CN$  solution trans-IF<sub>4</sub>O<sub>2</sub><sup>-</sup>. In IF<sub>5</sub> solution, however,  $IF_4O_2$ <sup>-</sup> was shown to exist in a cistrans equilibrium, with the cis isomer being favored.

**Tetrafluoroorthoperiodic** acid, HOIF40, was first prepared Ferrallitorial co-workers<sup>11</sup> according to<br>by Engelbrecht and co-workers<sup>11</sup> according to <br> $Ba_3H_4(IO_6)_2 + 14HSO_3F \rightarrow$ 

$$
Ba_3H_4(IO_6)_2 + 14HSO_3F →
$$
  
2HOIF\_4O + 8H\_2SO\_4 + 3Ba(SO\_3F)\_2  
CU 1GCP 2C 11.2H 16. J = 15.6 F1. J = 15

Since  $HOIF_4O$  could not be separated from  $HSO_3F$  by dissince HOIF<sub>4</sub>O could not be separated from HSO<sub>3</sub>F by dillation, it was converted into the more volatile  $IF_3O_2$ <br>HOIF<sub>4</sub>O + SO<sub>3</sub> → IF<sub>3</sub>O<sub>2</sub> + HSO<sub>3</sub>F

$$
IOIF4O + SO3 \rightarrow IF3O2 + HSO3F
$$

which was distilled off and then reconverted to HOIF<sub>4</sub>O by<br>HF addition according to<br> $IF_3O_2 + HF \rightarrow HOIF_4O$ HF addition according to

$$
IF3O2 + HF \rightarrow HOIF4O
$$

The acid was characterized<sup>6,11</sup> by its physical constants and mass and NMR spectra, which showed the compound to be a mixture of the cis and trans isomers, with the cis isomer **being**  more abundant. Selig and Elgad found<sup>9</sup> that solutions of  $NaIO<sub>4</sub>$  in anhydrous HF contain HOIF<sub>4</sub>O, as well as other unidentified fluorine species, and reported the 19F NMR spectra of both isomers in HF solution. Gillespie and Krasznai<sup>10</sup> also listed chemical shifts for cis- and trans-HOIF40, without specifying the solvent, and gave a coupling constant of 21 Hz for the cis isomer, which disagrees with the values of about 220 Hz reported by others. $6,9,11$ 

## **Experimental Section**

**Caution.** Two explosions were encountered in reactions involving FOIF,O. Most hypofluorites are shock-sensitive materials' and appropriate precautions should therefore be taken when one is working with larger amounts of FOIF<sub>4</sub>O.

**Materials.** Literature methods were used for the syntheses of  $NF_4SbF_6$ <sup>12</sup> IF<sub>5</sub>O,<sup>13,14</sup> and ClOSO<sub>2</sub>F.<sup>15</sup> CsIO<sub>4</sub> was prepared by slowly combining, with stirring, stoichiometric amounts of concentrated aqueous solutions of CsCl and NaI04. The mixture was cooled to 0 **"C,** and the CsI04 precipitate was filtered off, washed three times with ice water, and dried for 16 h in an oven at 110 °C. Its vibrational **spectra** showed no detectable **impurities.** Bismuth pentafluoride (Ozark Mahoning Co.) was used as received.  $BrF<sub>5</sub>$  (Matheson) was treated with 35 atm of F<sub>2</sub> at 200 °C for 24 h and then purified by fractional condensation through traps kept at **-64** and **-95** "C, with the material retained in the latter being used. Hydrogen fluoride (Matheson) was dried by treatment with 20 atm of  $F_2$  at room temperature, followed by storage over  $BiF_5$  to remove the last traces of  $H_2O^{16}$  ClF<sub>3</sub> (Matheson) and  $CIF_5$  (Rocketdyne) were purified by fractional condensation prior to their use.

**Apparatus.** Volatile materials used in this work were handled in either a Monel-Teflon FEP, a stainless steel-Teflon FEP, or a Teflon PFA vacuum line. The last was constructed exclusively from injection-molded PFA fittings and valves (Fluoroware, Inc.). The anhydrous HF was preferentially handled in the PFA or Monel line, whereas the halogen fluorides were handled mainly in a steel line. All lines were well passivated with  $CIF_3$  and, if HF was to be used, with HF. Nonvolatile materials were handled in the dry nitrogen atmosphere of a glove box. Metathetical reactions were carried out in HF solution using an apparatus consisting of two FEP U-traps interconnected through a coupling containing a porous Teflon filter **(see** Figure **1** of ref **17).** For NMR or low-temperature vibrational spectra, the second FEP U-trap, which served as a receiver, was replaced by either a 4-mm Teflon FEP or a thin-walled Kel-F tube.

Infrared spectra were recorded in the range **4000-200** cm-' on a Perkin-Elmer Model **283** spectrophotometer. Room-temperature

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spectra of solids were obtained by using dry powders pressed between AgCl disks. Spectra of gases were obtained by using a Teflon cell of 5-cm path length equipped with AgCl windows. The spectra of matrix-isolated FOIF<sub>4</sub>O and IF<sub>5</sub>O were obtained at 6 K with an Air Products Model **DE202S** helium refrigerator equipped with CsI windows. Research grade Ne (Matheson) was used as a matrix material in a mole ratio of **1000:l.** The spectrometer was calibrated by comparison with standard gas calibration points,<sup>18,19</sup> and the reported frequencies are believed to be accurate to  $\pm 2$  cm<sup>-1</sup>.

The Raman spectra were recorded on a Cary Model **83** spectrophotometer using the 4880-A exciting line and a Claassen filter<sup>20</sup> for the elimination of plasma lines. Sealed quartz, Teflon FEP, or Kel-F Tubes were used as sample containers in the transverse-viewing, transverse-excitation technique. Polarization measurements were carried out according to method VIII listed by Claassen et al.<sup>20</sup> Lines due to the Teflon or Kel-F sample tubes were suppressed by the use of a metal mask.

The I9F NMR spectra were recorded at **84.6** MHz on a Varian Model EM **390** spectrometer equipped with a variable-temperature probe. Chemical shifts were determined relative to external CFCl<sub>3</sub> with positive shifts being downfield from CFCl<sub>3</sub>.<sup>21a</sup>

The mass spectra were recorded with an EA1 Quad **300** quadrupole spectrometer at an ionization potential of **70** eV.

Preparation of CsIF<sub>4</sub>O<sub>2</sub>. In a typical experiment, CsIO<sub>4</sub> (31.44 mmol) was placed in a  $\frac{3}{4}$ -in. o.d. Teflon FEP ampule equipped with a stainless-steel valve. Anhydrous HF **(20 mL** of liquid) was condensed into the ampule, and the mixture was stirred with a magnetic stirring bar for **4 days** at ambient temperature. Volatile products were pumped off overnight at ambient temperature and for an additional **2** h at **50** "C. The solid residue **(1 1.402 g,** weight calculated for **31.44** mmol of CsIF402 **11.564 g)** was shown by Raman spectroscopy to still contain some unreacted CsI04. It was treated again, as described above, with fresh anhydrous HF (15 mL of liquid). After the residue was pumped to dryness, the Raman spectrum of the solid residue **(1 1.532 g)** showed **cis-** and trans-CsIF402 as the principal products and only a trace of unreacted CsI04.

A total of eight preparations were carried out in a similar manner, with use of shorter reaction times, slightly higher reaction temperatures  $(\sim 50 \text{ °C})$ , and rapid HF removal at elevated temperature. The conversion of  $CsIO<sub>4</sub>$  to  $CsIF<sub>4</sub>O<sub>2</sub>$  after the first HF treatment was generally in the range of **75-90%,** and the Raman spectra showed the presence of some unreacted  $CsIO<sub>4</sub>$ . This unreacted  $CsIO<sub>4</sub>$  was readily converted to  $CsIF_4O_2$  by repeated treatment with anhydrous HF; however in most *cases,* repeated HF treatments resulted in a slight weight decrease and the appearance of bands due to HF<sub>2</sub><sup>-</sup> (infrared **1435 s,** br, **1228 ms** cm-I; Raman complex band at **790-740** *cm-'* with maximum at 759 cm<sup>-1</sup>). This is caused by the solvolysis of CsIF<sub>4</sub>O<sub>2</sub> in anhydrous HF and the volatility of the resulting HOIF40 (see below). The ratio between the cis and trans isomers of  $CsIF_4O_2$  varied somewhat for the different preparations, with the trans isomer being slightly favored at the lower and the cis isomer being somewhat favored at the higher reaction temperatures.

The **CsI04-BrFS** System. Cesium periodate **(2.453** mmol) was placed in a passivated sapphire reactor equipped with a stainless-steel valve and a magnetic stirring bar. Bromine pentafluoride **(14.99** mmol) was added at **-196** "C, and the mixture was allowed to react during warm-up to room temperature. A fast reaction with gas evolution occurred, which was moderated by intermittent cooling with liquid **N2.** After completion of the warm-up cycles, the mixture was stirred at **20** "C for **24** h, resulting in a clear, pale yellow solution. The Raman spectrum of this solution showed the presence of  $BrF_5$ ,  $BrF_3O$ , and  $IF_4O_2^-$  (mainly trans with a small amount of cis isomer). The solution was kept at 22 °C for 4 days and then cooled to -196 °C. The materials volatile at **-196** "C consisted of **1.92** mmol of oxygen. The materials volatile at **22** "C were separated by fractional condensation and identified by Raman spectroscopy. They consisted of unreacted

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 $BrF<sub>5</sub>$  (9.9 mmol) and a mixture of  $BrF<sub>3</sub>$  and  $BrF<sub>3</sub>O$  (found 682 mg, calcd for 3.84 mmol of  $BrF_3 + 1.06$  mmol of  $BrF_3O$  688 mg). The solid residue (found 804 mg, weight calcd for 2.453 mmol of  $CsIF<sub>4</sub>O<sub>2</sub>$ 787 mg) was shown by Raman spectroscopy to consist mainly of *trans*-CsIF<sub>4</sub>O<sub>2</sub>, CsBrF<sub>4</sub>, and smaller amounts of *cis*-CsIF<sub>4</sub>O<sub>2</sub>, and possibly some solvated  $BrF_3O$ . Vacuum pyrolysis at 90 °C resulted in a solid residue consisting again of trans-CsIF<sub>4</sub>O<sub>2</sub>, CsBrF<sub>4</sub> and a small amount of cis  $CsIF<sub>4</sub>O<sub>2</sub>$  and also in the evolution of some IF<sub>5</sub>  $(\sim 8$  weight %).

The CsIO<sub>4</sub>-ClF<sub>3</sub> System. A well-passivated (with ClF<sub>3</sub>) sapphire tube equipped with a stainless-steel valve and containing a Tefloncoated stirring bar was loaded with **CsIO,** (1.14 mmol), followed by  $CIF<sub>3</sub>$  (10.6 mmol). The liquid  $CIF<sub>3</sub>$  and solid periodate were stirred magnetically overnight at  $0-20$  °C. This resulted in a clear, very pale yellow solution. Upon removal of the volatile material and several hours of pumping at ambient temperature, a white powder (0.493 g) remained in the tube, which was identified by vibrational spectroscopy as a mixture of  $CsF·3IF_5$  and  $CsCIF_4$  (weight calculated for the conversion of 1.14 mmol of  $CsIO<sub>4</sub>$  to 0.38 mmol of  $CsF·3IF<sub>5</sub>$  and 0.76 mmol of  $CsClF_4$  was 0.497 g). The volatile materials consisted of CIF,  $FCIO<sub>2</sub>$ , and unreacted CIF<sub>3</sub>.

**The CsIO<sub>4</sub>-CIF<sub>5</sub> System.** When CsIO<sub>4</sub> was allowed to interact with a large excess of  $CIF_5$  in a stainless-steel reactor at room temperature, the composition of the solid reaction product depended on the reaction time. After short reaction times (about several hours) the solid consisted, on the basis of its weight change and Raman spectra, mainly of unreacted  $CsIO<sub>4</sub>$  and smaller amounts of trans-CsIF<sub>4</sub>O<sub>2</sub>. After longer reaction times (in excess of 1 month), the solid consisted mainly of  $CsIF_8$  and *trans*-CsIF<sub>4</sub>O<sub>2</sub> and some CsIF<sub>4</sub>O.

**The CsIO<sub>4</sub>-F<sub>2</sub> System.** The fluorination of CsIO<sub>4</sub> with elemental fluorine in a static system at temperatures up to 60  $^{\circ}$ C resulted in a solid product, which, on the basis of its vibrational spectra, was a mixture of mainly  $CsIF_8$ ,  $CsIF_6$ , and  $CsIO_4$  with smaller amounts of CsIF<sub>4</sub>O and *cis*- and *trans*-CsIF<sub>4</sub>O<sub>2</sub> also being present.

**Synthesis of HOIF<sub>4</sub>O.** In a typical experiment, CsIF<sub>4</sub>O<sub>2</sub> (2.0 mmol) and BiF, (2.0 mmol) were placed in a passivated Teflon FEP U-trap containing a magnetic stirring bar. One arm of the trap was closed off by the stainless-steel valve, while the other one was connected through a porous Teflon filter to a second Teflon U-trap, which was capped off by another valve. Anhydrous HF *(5* mL of liquid) was condensed into the U-trap, and the  $CsIF_4O_2-BiF_5-HF$  mixture was stirred at 25 °C for 1 h. The double U-trap assembly was cooled to -78 **"C** and inverted, and the HOIF,O-containing HF solution was separated from the CsBiF<sub>6</sub> precipitate by pressure filtration. The HF solvent was pumped off at  $-45$  and  $-13$  °C. The residue was allowed to warm to ambient temperature, and the material volatile at 25 °C was collected at  $-78$  °C in a 4-mm o.d. external Teflon U-trap. This trap was shown to contain HOIF<sub>4</sub>O ( $\sim$ 2 mmol), which was identified by its Raman and <sup>19</sup>F NMR spectra. The filter cake (0.9 g) was identified by its Raman spectrum as  $CsBiF<sub>6</sub>$ .

**Synthesis of FOIF<sub>4</sub>O.** In a typical experiment, CsIF<sub>4</sub>O<sub>2</sub> (5.0 mmol) and  $NF_4SbF_6$  (5.0 mmol) were placed in the Teflon FEP metathesis apparatus (see above), and anhydrous HF (5 mL of liquid) was condensed in at  $-78$  °C. The mixture was stirred for 1 h at room temperature. The apparatus was cooled to  $-78$  °C and inverted, and the white precipitate was separated from the solution by pressure filtration. Most of the HF solvent was pumped off over several hours at temperatures ranging from  $-64$  to  $-30$  °C. The resulting white solid residue was allowed to decompose during slow warm-up from -30 "C to ambient temperature. The volatile products were passed through a Teflon U-trap containing passivated NaF pellets, followed by a series of cold traps kept at  $-78$ ,  $-95$ ,  $-112$ , and  $-210$  °C. The -78 "C trap contained a small amount of unidentified material, which was discarded, the -95  $\degree$ C fraction consisted of pure FOIF<sub>4</sub>O (2.36) mmol), the  $-112$  °C trap had 1.69 mmol of  $\overline{FOIF}_4O$  containing a small amount of IF<sub>5</sub>O as an impurity, and the -210  $\degree$ C trap contained  $NF<sub>3</sub>$  (4.0 mmol). A small amount of white solid residue, which was left behind after the thermal decomposition of the filtrate, was shown by vibrational spectroscopy to consist mainly of trans-CsIF<sub>4</sub>O<sub>2</sub>. The filter cake (1.8 g) was identified by Raman spectroscopy as  $CsSbF<sub>6</sub>$ . The  $-95$  °C fraction was used for the characterization of  $FOIF_4O$ and was shown by vibrational and <sup>19</sup>F NMR analysis to be free of  $IF<sub>s</sub>O$ .

For the elemental analysis, 278.7 mg of the material was condensed at  $-196$  °C into an ampule containing 12 mL of frozen 1 N NaOH. The mixture was warmed to ambient temperature for 12 h and then

analyzed for total iodine by energy-dispersive X-ray fluorescence spectrometry, for **IO4-** by iodometric titration, for base consumption by back-titration with 0.1 N HC1 using a pH electrode and for fluoride by titration using  $La(NO<sub>3</sub>)<sub>3</sub>$  and an Orion specific-ion electrode. Anal. Calcd for  $FOIF_4O: I$ , 49.98; F, 37.42; OH<sup>-</sup> consumed, 6.0 equiv/mol; iodometric titration, 8.0 equiv/mol, with the assumption of the hydrolysis reaction  $FOIF_4O + 6OH^- \rightarrow IO_4^- + 5F^- + 0.5O_2(g) + 3H_2O$ . Found: I, 50.0; F, 36.0; OH- consumed, 6.1 equiv/mol; iodometric titration, 7.8 equiv/mol.

Synthesis **of C101F40.** A 30-mL stainless-steel cylinder was loaded with 2.32 mmol of  $CsIF_4O_2$ , and 2.12 mmol of  $CISO_3F$  was added at -196 °C. After the cylinder was kept for 5 days at -78 °C, the volatile products were removed from the cylinder. The solid residue was identified by vibrational spectroscopy as  $CsSO<sub>3</sub>F$ . The volatiles were fractionated through traps cooled to  $-45$ ,  $-78$ , and  $-196$  °C. The lowest temperature fraction (0.77 mmol) was mainly  $\text{Cl}_2$  together with some FCIO<sub>2</sub>, while the -45 °C trap contained a white solid, which melted above 0 °C and which was identified by its infrared spectra as IF<sub>5</sub>. The -78 °C trap contained a yellow-orange solid, which on slight warming melted to an orange liquid. Its gas-phase infrared spectrum was recorded at 25 °C and showed the following bands (cm<sup>-1</sup>; relative intensity, assignment): 912 m, I=O stretch; 763 mw, O-Cl stretch; 678 vs, 635 s, 532 mw, I-F and **1-0** stretching. The compound was found to be thermally unstable and very difficult to handle. It readily decomposed to  $IF_5$ , and its synthesis required careful temperature control. When the synthesis was carried out for example at -45 °C, only decomposition products were obtained. Attempts to isolate fluorocarbon derivatives of C101F40 by adding it across the C= $\text{C}$  double bond of C<sub>2</sub>F<sub>4</sub> resulted at -78 °C in no reaction and at room temperature in the oxygenation, fluorination, and decomposition products  $COF_2$ ,  $CF_3COF$ ,  $ClCF_2COF$ ,  $C_2F_3Cl$ , and  $IF_5$ .

#### **Results and Discussion**

**Synthesis of**  $CsIO<sub>2</sub>F<sub>4</sub>$ **.** In our work the known<sup>9</sup> equilibrium

$$
IO_4^- + 4HF \rightleftarrows IF_4O_2^- + 2H_2O
$$

was utilized for a convenient synthesis of  $CsIF<sub>4</sub>O<sub>2</sub>$ . So that this equilibrium can be shifted to the right, a large excess of HF must be used and the HF treatment must be repeated at least once. The resulting  $CsIF<sub>4</sub>O<sub>2</sub>$  consists of a mixture of the cis and trans isomers, as shown by  $^{19}$ F NMR and vibrational spectroscopy (see below). The ratio of cis to trans isomer varies somewhat with the reaction conditions used, but the formation of the cis isomer appears to be slightly favored. The 19F NMR and Raman spectra were recorded for solutions of  $CsIF<sub>4</sub>O<sub>2</sub>$ in anhydrous HF and CH<sub>3</sub>CN. Whereas the CH<sub>3</sub>CN solution spectra show the presence of the  $IF_4O_2^-$  anion, the spectra of the HF solutions are characteristic (see below) for those of HOIF40. This finding is in excellent agreement with the previous report by Selig and Elgad<sup>9</sup> that solutions of NaIO<sub>4</sub> in anhydrous HF contain  $HOIF<sub>4</sub>O$  and the report by Engelbrecht and co-workers<sup>6,7</sup> that HOIF<sub>4</sub>O interacts with alkalimetal fluorides to form  $IF_4O_2^-$  salts. Consequently, the above equilibrium reaction involves at least two reactions, the first being

$$
IO_4^-
$$
 + 6HF $\xrightarrow{-HF}$  HOIF<sub>4</sub>O + HF<sub>2</sub><sup>-</sup> + 2H<sub>2</sub>O

and, upon HF and  $H_2O$  removal, the second being

$$
-HF
$$
  
on HF and H<sub>2</sub>O removal, the second being  
HOIF<sub>4</sub>O + CsHF<sub>2</sub>  $\frac{-HF}{+HF}$  CsIF<sub>4</sub>O<sub>2</sub> + 2HF

The intermediate formation and the slight volatility of  $HOIF<sub>4</sub>O$  also explain why in some of our  $CsIF<sub>4</sub>O<sub>2</sub>$  preparations, when the HF was rapidly pumped off at elevated temperature, a weight loss accompanied by some  $CsHF_2$  formation was observed.

The above synthesis of  $CsIF_4O_2$  from  $CsIO_4$  and HF appears more convenient than the previously reported methods involving either the difficult to obtain  $IF_3O_2$  as a starting material or the isolation and recrystallization of  $MIF_4O_2.2IF_5$ from  $IF_5$ , followed by its pyrolysis. However, the latter method produces almost exclusively the trans isomer and might be the



**Figure 1.** Vibrational spectra of a CsIF<sub>4</sub>O<sub>2</sub> sample containing mainly *trans*-IF<sub>4</sub> $O_2^-$  (bands marked by their frequency values) and smaller amounts of  $cis$ -IF<sub>4</sub>O<sub>2</sub>. Trace A is an infrared spectrum of the solid as a dry powder pressed between AgCl disks; the broken line indicates absorption due to the window material. Trace B is a Raman spectrum of the solid. Inserts C are Raman bands of the  $CH<sub>3</sub>CN$  solution, recorded with parallel and perpendicular polarization.

preferred method if pure trans-IF<sub>4</sub>O<sub>2</sub><sup>-</sup> is desired. The fact that the cis-trans isomer ratio strongly depends on the nature of the reactants suggests that this ratio is kinetically and not thermodynamically controlled. This conclusion is in excellent agreement with those reached by Toetsch and Sladky for the closely related  $\text{TeF}_4(\text{OH})_2$  system.<sup>21b</sup>

An alternate method for the formation of  $CsIF_4O_2$  involves the reaction of CsIO<sub>4</sub> with BrF<sub>5</sub>. The main reaction can be described by  $CSIO_4 + 2BrF_5 \rightarrow CsIF_4O_2 + 2BrF_3O$ described by

$$
CsIO4 + 2BrF5 \rightarrow CsIF4O2 + 2BrF3O
$$

This reaction is analogous to that<sup>10</sup> previously reported for<br> **KIO<sub>4</sub>** + IF<sub>5</sub>, i.e.<br> **KIO<sub>4</sub>** + 2IF<sub>5</sub> → **KIF<sub>4</sub>O<sub>2</sub>** + 2IF<sub>3</sub>O  $KIO<sub>4</sub> + IF<sub>5</sub>, i.e.$ 

$$
XIO_4 + 2IF_5 \rightarrow KIF_4O_2 + 2IF_3O
$$

and produces almost entirely the trans isomer. Compared to the IF<sub>5</sub> reaction, the BrF<sub>5</sub> reaction offers the advantage that the  $BrF_3O$  and  $BrF_3$  byproducts are volatile and can easily be pumped off. However, the resulting product was contaminated by nonvolatile  $CsBrF<sub>4</sub>$ , which could not be readily separated from the  $CsIF<sub>4</sub>O<sub>2</sub>$ .

The reactions of  $CsIO<sub>4</sub>$  with chlorine fluorides were also briefly studied. With ClF<sub>5</sub>, trans-CsIF<sub>4</sub>O<sub>2</sub> was formed in low conversion according to<br>CsIO<sub>4</sub> + ClF<sub>5</sub>  $\rightarrow$  CsIF<sub>4</sub>O<sub>2</sub> + FClO<sub>2</sub> conversion according to

$$
CsIO4 + ClF5 \rightarrow CsIF4O2 + FCIO2
$$

Attempts to achieve higher conversions by the use of longer reaction times failed due to the formation of  $CsIF<sub>s</sub>$  as the main product and of CsIF40 as a minor product.

When ClF<sub>5</sub> was replaced by the more reactive ClF<sub>3</sub>, complete conversion of the  $CsIO<sub>4</sub>$  was obtained; however, all the



**Figure 2.** Vibrational spectra of a  $CsIF<sub>4</sub>O<sub>2</sub>$  sample containing mainly **cis-IF402-** (bands marked by frequency values) and smaller amounts of trans-IF<sub>4</sub>O<sub>2</sub><sup>-</sup> (marked by t). Trace A is an infrared spectrum of the solid as a dry powder pressed between AgCl disks. The weak bands at 815 and 470  $cm^{-1}$  probably do not belong to  $IF_4O_2^-$ . Trace B is a Raman spectrum of the solid. Traces C are Raman spectra of a  $CH<sub>3</sub>CN$  solution. Solvent bands are marked by an asterisk.

oxygens in  $IO<sub>4</sub>^-$  were exchanged for fluorine, and the solid product consisted of a mixture of  $CsIF_{6}^22IF_{5}^{22}$  and  $CsCIF_{4}^{23}$ Based on the observed material balance, the following reaction<br>occurred:<br> $3CsIO<sub>4</sub> + 11ClF<sub>3</sub> \rightarrow$ occurred:

$$
sIO4 + 11CIF3 \rightarrow
$$
  
6FCIO<sub>2</sub> + 3CIF + 2CsCIF<sub>4</sub> + CsIF<sub>6</sub>·2IF<sub>5</sub>

The formation of CIF and of half of the  $FCIO<sub>2</sub>$  can be readily explained by the well-known<sup>24</sup> disproportionation of the expected unstable FClO intermediate:

$$
2FCIO \rightarrow FCIO_2 + CIF
$$

The fluorination of  $CsIO<sub>4</sub>$  by elemental fluorine at temperatures up to 60 °C in a static system was also studied. The main products were  $CsIF_8$  and  $CsIF_6$ , with  $CsIF_4O$  and cisand trans-CsIF<sub>4</sub>O<sub>2</sub> as minor products.

In view of the fact that the fluorination reactions of CSIO, with  $CIF_5$ ,  $BrF_5$ ,  $CIF_3$ , or  $F_2$  do not result in pure compounds, they are less attractive synthetic methods for the preparation of  $CsIF<sub>4</sub>O<sub>2</sub>$ .

**Vibrational Spectra of CsIF,O2.** The vibrational spectra of solid  $CsIF_4O_2$  were recorded for samples that differed in their cis and trans isomer content. The observed spectra are given

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Table **11.** Vibrational Spectra of cis-IF<sub>4</sub>O<sub>7</sub>

		obsd freq. $cm^{-1}$ , and rel intens <sup>a</sup>			
solid		CH <sub>3</sub> CN soln	assignt for	approx description	
IR	Raman	Raman	point group $C_{2\nu}$	of mode	
875 vs	875	870 sh	$\nu_{12}$ (B <sub>2</sub> )	$\nu_{\rm as}({\rm IO}_2)$	
855 vs	856 (10)	851 (9) p	$\nu, (A_1)$	$\nu_{\rm s}({\rm IO}_2)$	
600 vs, br			$v_a$ (B <sub>1</sub> )	$\nu_{\text{as}}(\text{IF}_2)_{\text{ax}},$ $\nu_{\text{as}}(IF_{2})_{\text{eq}}$	
600 vs. br	605(9.8)	$609(10)$ p	$\nu_{2}(A_{1})$	$\nu_{\rm s}(\rm IF_2)_{\rm eq}$	
550 mw	552 sh	540 (10) p	$\nu$ , $(A_1)$	$\nu_{\rm s}(\text{IF}_2)_{\rm ax}$	
395 sh	394 (3.4)		$\nu_{\bullet}$ (A <sub>1</sub> )	$\delta_{\text{sciss}}(IO_2)$	
364s	365 sh	355 sh			
328 mw	332(6.5) 235(0.2) 210 (0.5)	335 sh			

*a* Uncorrected Raman intensities (peak height).

in Figures 1 and 2, and the observed frequencies and their assignments in point group  $D_{4h}$  and  $C_{2v}$ 



are summarized in Tables I and 11, respectively. The bands belonging to the trans isomer could be readily distinguished from those of the cis isomer due to the fact that only the trans isomer has a center of symmetry, which causes the infrared and Raman bands to be mutually exclusive. Furthermore, the 19F NMR spectrum (see below) clearly distinguished the trans from the cis isomer and established which isomer was more abundant in a given sample.

**Assignments and Normal-Coordinate Analysis for** *trans-*IF<sub>4</sub>O<sub>2</sub><sup>-</sup>. The trans-IF<sub>4</sub>O<sub>2</sub><sup>-</sup> anion of symmetry  $D_{4h}$  should possess 11 fundamental modes classified as  $2 \overrightarrow{A}_{1g} + 2 A_{2u} +$  $B_{1g} + B_{2g} + B_{2u} + E_g + 3 E_u$ . Of these, the A<sub>1g</sub>, B<sub>1g</sub>, B<sub>2g</sub>, and  $E_{g}$  modes should be Raman active only and the  $A_{2u}$  and  $E_{u}$ modes should be infrared active only, whereas the  $B_{2u}$  mode should be inactive in both spectra. Of the 10 active modes, all five Raman-active modes and three out of the five infrared-active modes have been observed and can be readily assigned on the basis of their activity, polarization data, and comparison with the closely related species IF<sub>4</sub><sup>-25</sup> IF<sub>4</sub>O<sup>-26</sup>  $IF_{5}^{27,28}$  and  $IF_{5}O^{14,29}$  (see Table I). The correctness of these assignments was confirmed by a normal-coordinate analysis using the symmetry coordinates and *G* matrix elements previously published<sup>30</sup> by Beattie and co-workers. The bond distances were estimated to be  $r_{IF} = 1.92$  Å and  $d_{IO} = 1.72$ **A,** on the basis of the known structures and stretching frequencies of the related  $IF_5O^{14,29,31}$  and  $IF_5^{27,28,32}$  molecules and the  $IF_4O^-$  anion.<sup>26,33</sup> The force constants of the  $B_{1g}$ ,  $B_{2g}$ , and  $E_s$  species are uniquely determined. In the  $A_{1g}$  block, the  $G_{12}$ element is zero, and therefore  $F_{12}$  can be ignored. For the  $A_{2u}$ block, the extremal solution<sup>34</sup>  $F_{44}$  = minimum was used, which has previously been shown<sup>25</sup> for the closely related  $IF_{4}^-$  anion

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Table **III.** Symmetry Force Constants<sup>a</sup> of trans-IF<sub>4</sub>O<sub>2</sub>

A<sub>18</sub> 
$$
\nu_1
$$
 569  $F_{11} = f_r + 2f_{rr} + f_{rr'}$  3.623  
\n $\nu_2$  824  $F_{22} = f_{\alpha} + f_{\alpha\alpha}$  6.400  
\nA<sub>21</sub>  $\nu_3$  885  $F_{33} = f_d - f_{dd}$  5.903  
\n $\nu_4$  349  $F_{44} = f_{\beta} + 2f_{\beta\beta} + f_{\beta\beta'} - f_{\beta\beta''} - 2f_{\beta\beta}^{\circ} - f_{\beta\beta}^{\dagger}$  1.700  
\nA<sub>21</sub>  $F_{34} = 2f_{d\beta} - 2f_{d\beta'}$  6.400  
\nA<sub>22</sub>  $F_{34} = 2f_{d\beta} - 2f_{d\beta'}$  6.400  
\nA<sub>23</sub> 6.400  
\nA<sub>24</sub> 349  $F_{44} = f_{\beta} + 2f_{\beta\beta} + f_{\beta\beta'} - 2f_{\beta\beta''} - 2f_{\beta\beta}^{\dagger}$  6.400

$$
B_{1g} \quad \nu_s \quad 555 \quad F_{ss} = f_r - 2f_{rr} + f_{rr'} \tag{3.447}
$$
\n
$$
B_{r\sigma} \quad \nu_s \quad 255 \quad F_{rs} = f_{\infty} - 2f_{\infty} + f_{\infty'} \tag{0.671}
$$

$$
E_g \nu_s \quad 380 \ F_{ss} = f_\beta - f_{\beta\beta}' - f_{\beta\beta}'' + f_{\beta\beta} + \nu_{\beta\beta} \tag{1.201}
$$

$$
E_{\mathbf{u}}^{\mathbf{v}} \quad \nu_{\mathbf{y}} \quad 590 \quad F_{\mathbf{y}\mathbf{y}}^{\mathbf{y}} = f_{r} - f_{rr}^{\mathbf{y}} \tag{2.998}
$$

**a stretching constants in mdyn/& deformation constants in**  mdyn A/rad<sup>2</sup>, and stretch-bend interaction constants in mdyn/rad.

to be an excellent approximation to a general valence force field for these weakly coupled systems. For the **E,,** block, only the frequency value of the stretching mode is experimentally known. A comparison with the force field of the related  $IF_4$ anion<sup>25</sup> showed that, due to the heavy iodine central atom, the approximation  $F_{99} = \lambda_9/G_{99}$  yields an almost exact value for the stretching force constant in the **E,** block and was therefore used for  $IF_4O_2^-$ . The resulting force field is listed in Table I11 and strongly supports our assignments. Table IV gives a comparison of the internal stretching force constants of *trans*-IF<sub>4</sub>O<sub>2</sub><sup>-</sup> with those of the closely related species IF<sub>4</sub><sup>-25</sup> IF<sub>4</sub>O<sup>-</sup>,<sup>26</sup> IF<sub>5</sub>,<sup>35</sup> IF<sub>5</sub>O,<sup>29</sup> and IF<sub>6</sub><sup>+</sup>.<sup>36</sup> As previously discussed<sup>37</sup> for chlorine oxyfluorides, the IF stretching force constants increase in the sequence anions < neutral molecules < cations and within a given group with increasing oxidation state of the iodine central atom. The IO stretching force constants are in the range expected for I=0 double bonds and demonstrate that, even in the anions, the formal negative charge is located mainly on the more electronegative fluorine ligands rather than on the oxygen ligand. Consequently, contributions from resonance structures such as I and I1 are more important



than those from III and IV to explain the bonding in  $IF_4O_2^-$ .



Resonance structures such as I and I1 also account for the decrease of the IF stretching force constants with increasing formal negative charges and also with decreasing oxidation state of the central atom. Both effects increase the  $I^{\delta+}-F^{\delta-}$ polarity of the IF bonds, thereby causing the bonds to become more ionic, longer, and therefore weaker. This weakening of the IF bonds can be very significant as is demonstrated by the low value of f, in IF<sub>4</sub>-, which is only 41% of that in IF<sub>6</sub><sup>+</sup>. The fact that the IO stretching force constant drops from  $IF_4O^-$ (6.56 mdyn/Å) toward  $IF_4O_2^-(6.15 \text{ mdyn/A})$ , in spite of an increase in the oxidation state of the iodine atom, is interesting and parallels the trends previously noted<sup>37</sup> for chlorine oxyfluorides, i.e., the electron-releasing effect of oxygen ligands

in highly electronegative compounds.

**Assignments for cis-IF<sub>4</sub>O<sub>2</sub>.** Our assignments for cis-IF<sub>4</sub>O<sub>2</sub>. have been limited to the stretching modes because only 10 of the 15 fundamentals expected for point group  $C_{2v}$  have been observed and because no reliable assignments have been published for similar  $XF_4O_2$  species. The assignment of the two  $IO<sub>2</sub>$  stretching modes is straightforward on the basis of their high frequencies, relative intensities, and the previously published <sup>18</sup>O spectra.<sup>9</sup> The symmetric IF<sub>2</sub> axial and the symmetric  $IF<sub>2</sub>$  equatorial stretches must belong to the two intense polarized Raman bands at **540** and 609 cm-', respectively, with the axial mode resulting in a weak and the equatorial mode resulting in a strong infrared counterpart. The antisymmetric axial and the antisymmetric equatorial  $IF<sub>2</sub>$ stretches should both be very intense in the infrared spectrum and therefore are assumed to coincide at about  $600 \text{ cm}^{-1}$ , resulting in a very strong, broad band.

**Comparison with Previous**  $IF_4O_2^-$  **Assignments. Disregarding** some solvent-induced shifts, we find the above assignments for cis-IF<sub>4</sub>O<sub>2</sub><sup>-</sup> agree well with those previously reported<sup>9</sup> by Selig and Elgad for an aqueous solution. The only minor discrepancy is the assignment of the antisymmetric axial  $IF<sub>2</sub>$ stretch. For trans-IF<sub>4</sub>O<sub>2</sub><sup>-</sup>, the assignments proposed by Gillespie and Krasznai for **six** of the modes have been revised for three of them. The vibrational spectra reported by Carter et al.<sup>8</sup> show that their sample contained mainly *trans*-IF<sub>4</sub>O<sub>2</sub> but was incorrectly interpreted in terms of the cis isomer.

<sup>19</sup>F NMR Spectra of IF<sub>4</sub>O<sub>2</sub><sup>-</sup> and **HOIF<sub>4</sub>O.** The presence and the relative amounts of *cis*- and of *trans*-IF<sub>4</sub> $O_2$ <sup>-</sup> in the above samples were verified by <sup>19</sup>F NMR spectroscopy. The spectra were recorded in CH<sub>3</sub>CN solution at  $-70$  °C and showed a narrow singlet at  $\phi$  65.1 for the trans isomer and a broader  $A_2B_2$  pattern at  $\phi$  66.0 and 112.8 with  $J_{FF}$  = 204 Hz for the cis isomer. The observed shifts and coupling constant are in fair agreement with the value previously reported for solutions in CH<sub>3</sub>CN (trans  $\phi$  62.0),<sup>10</sup> IF<sub>5</sub> [cis  $\phi$  68.5, 102.1 *(J<sub>FF</sub>* = 202) Hz); trans  $\phi$  70.6], and aqueous HF [cis  $\phi$  64, 105  $(J_{\text{FF}} = 196$  $Hz$ ].<sup>9</sup>

Solutions of  $CsIF_4O_2$  in anhydrous HF at  $-75$  °C resulted in a sharp singlet at  $\phi$  62.0 and a broadened  $A_2B_2$  pattern at  $\phi$  61.8 and 85.9 with  $J_{FF}$  = 220 Hz. At room temperature, the  $A_2B_2$  pattern was broadened to the extent that it could barely be detected. Although these spectra are similar to those of  $IF_4O_2$ , it was conclusively shown (see below) by Raman spectroscopy that they are due to *cis-* and trans-HOIF40 and not to  $IF_4O_2^-$ . This finding is in excellent agreement with the conclusion9 reached by Selig and Elgad that their signals **1**  and 2 observed for solutions of NaI04 in HF are due to *cis*and trans-HOIF<sub>4</sub>O. The observed chemical shifts and coupling constant are in fair agreement with previous reports, $6,7,9,10$ considering the different solvents and conditions used for recording the spectra. The coupling constant of 21 Hz previously reported by Gillespie and Krasznai<sup>10</sup> appears to be a typographical error.

Synthesis and Properties of HOIF<sub>4</sub>O. The above described experiments involving CsIF<sub>4</sub>O<sub>2</sub> in HF solutions indicate the existence of the equilibrium<br>
CsIF<sub>4</sub>O<sub>2</sub> + 2HF  $\frac{+HF}{-HF}$  CsHF<sub>2</sub> + HOIF<sub>4</sub>O existence of the equilibrium

$$
CsIF_4O_2 + 2HF \xrightarrow{-HF} CsHF_2 + HOIF_4O
$$

which, in the presence of a large excess of HF, is shifted all the way to the right side. In view of the lack of a convenient synthesis of  $HOIF_4O,$ <sup>6,7</sup> this reaction was utilized to prepare HOIF40. Since during HF removal the above equilibrium is shifted back to the left, the  $CsHF_2$  was converted into an insoluble  $BiF_6^-$  salt according to DIF<sub>4</sub>O,<sup>6,7</sup> this reaction was utility<br>the during HF removal the above to the left, the CsHF<sub>2</sub> was consalt according to<br>SHF<sub>2</sub> + BiF<sub>5</sub>  $\xrightarrow{\text{HF}}$  CsBiF<sub>6</sub><sup>1</sup>

$$
CsHF_2 + BiF_5 \xrightarrow{HF} CsBiF_6\downarrow
$$

which can be filtered off at  $-78$  °C. The resulting mixture

**<sup>(35)</sup> Christe, K. 0.; Curtis, E. C.; Wilson, R. D.** *J. Inorg. Nucl. Chem.* 

*Suppl.* **1976, 437. (36) Christe, K. 0.; Sawodny, W.** *Inorg. Chem.* **1967,6, 1783. (37) Christe, K. 0.; Schack, C. J.** *Ado. Inorg. Chem. Radiochem. 1976, 18,*  **331.** 

Table IV. Stretching Force Constants (mdyn/A) of trans-IF<sub>4</sub>O<sub>2</sub><sup>-</sup> Compared to Those of Similar Molecules and Ions

	anions			molecules		cation	
	IF $^+$ (+III) <sup><math>\alpha</math></sup>	$IF_4O^-(+V)^b$	IF <sub>4</sub> O <sub>7</sub> $(+VII)$	IF, $(+V)^c$	IF, $O (+VII)^d$	IF $^+$ (+VII) <sup>e</sup>	
$f_r$ (IF)	2.22	2.46	3.27	3.77	4.42	5.42	
Jrr	0.18	0.16	0.04	0.04	0.00	$-0.07$	
$_{Irr^{'}}$	0.47	0.45	0.27	0.38	0.18	0.19	
$f_d(IO)$		6.56	6.15		6.99		
Jdd			0.25				

<sup>*a*</sup> Data from ref 25; iodine oxidation state in parentheses for all species. <sup>*b*</sup> Data from ref 26. *<sup>c</sup>* Data from ref 35. *<sup>d</sup>* Data from ref 29. **e** Data from ref **36.** 

of HOIF40 and HF can be easily separated by fractional condensation or distillation.

The <sup>19</sup>F NMR spectrum of  $HOIF<sub>4</sub>O$  in HF solution was recorded at -78 °C and was identical with that of the product obtained by dissolving  $CsIF_4O_2$  in HF (see above). The ratio of cis to trans isomer in the HOIF40 sample appeared to be similar to that in the  $CSIF_4O_2$  starting material. It should be pointed out that at room temperature the signal due to the cis isomer can be so broad that it is difficult to detect, thereby giving the false impression of dealing with samples containing exclusively the trans isomer.

The Raman spectra of liquid  $HOIF<sub>4</sub>O$  showed some variation. Freshly prepared samples and HF solutions exhibited spectra similar to that of trace A of Figure 3. After the solutions were allowed to stand, the **872** cm-' band decreased in intensity and bands at **828** and **799** cm-' started to grow. In addition the bands in the **600-700** cm-' region became broader and shifted to slightly lower frequencies, as shown by trace B of Figure 3. On the basis of its <sup>19</sup>F NMR spectrum, a sample of HOIF40 in HF solution, which showed a Raman spectrum very similar to that of trace A of Figure 3, consisted mainly of the cis isomer. Whether the change from Raman spectrum A to spectrum B involves a change in the isomer ratio or is caused by association effects was not clearly established.

Synthesis of FOIF<sub>4</sub>O. Previous studies have shown that unstable  $NF_4$ <sup>+</sup> salts containing oxyanions such as  $ClO_4^{-3}$  or  $SO_3F^{-4}$  can be prepared by metathesis in anhydrous HF solution according to thesis of FOIF<sub>4</sub>O. Previous studies have shown<br>le NF<sub>4</sub><sup>+</sup> salts containing oxyanions such as ClO<sub>4</sub><br><sup>4</sup> can be prepared by metathesis in anhydrous H<br>according to<br>NF<sub>4</sub>SbF<sub>6</sub> + CsXO<sub>4</sub>  $\xrightarrow{\text{HF}}$  CsSbF<sub>6</sub> + + NF<sub>4</sub>XO<sub>4</sub><br>

$$
NF_4SbF_6 + CsXO_4 \xrightarrow{HF} CsSbF_6 + NF_4XO_4
$$

Thermal decomposition of these  $NF_4$ <sup>+</sup> salts provided a new high-yield synthetic route to hypofluorites.<sup>3,4</sup> Since no examples of iodine hypofluorites had previously been known, it was interesting to examine the applicability of this method to periodates.

Since the  $IO_4^-$  anion is fluorinated to  $IF_4O_2^-$  in anhydrous HF, as shown by the above studies and the previous report by Selig and Elgad,<sup>9</sup> the metathetical reaction of  $IO<sub>4</sub>$  itself could not be studied. However, when  $IO_4^-$  was replaced by  $IF_4O_2^-$ , the following metathetical reaction occurred:

$$
NF4SbF6 + CsIF4O2 \xrightarrow[-.78 \text{ } ^{+}C_5SbF6]+ HOIF4O + NF4HF2
$$

The CsSbF<sub>6</sub> precipitate could be easily filtered off at  $-78$  °C, and Raman and <sup>19</sup>F NMR spectroscopy of the filtrate showed the presence of  $NF_4+38$  and HOIF<sub>4</sub>O (see above) with no evidence for the  $IF_4O_2^-$  anion. This is in agreement with the above results for CsIF<sub>4</sub>O<sub>2</sub>, which demonstrated that MIF<sub>4</sub>O<sub>2</sub> salts undergo solvolysis in anhydrous HF according to<br>MIF<sub>4</sub>O<sub>2</sub> + 2HF  $\rightarrow$  MHF<sub>2</sub> + HOIF<sub>4</sub>O

$$
MIF_4O_2 + 2HF \rightarrow MHF_2 + HOIF_4O
$$

Raman and <sup>19</sup>F NMR spectra showed that these  $NF_4HF_2$ -HOIF40-containing HF solutions are unstable at room tem-





**Figure 3.** Raman spectra of **liquid** HOIF,O, recorded in Teflon **FEP**  tubes at room temperature.

perature and slowly decompose to  $NF<sub>3</sub><sup>39,40</sup>$  and a new compound identified (see below) as a mixture of cis- and trans-FOIF<sub>4</sub>O. At the same time, the relative intensities of the  $NF_4^+$ and  $HOIF<sub>4</sub>O$  signals decreased accordingly. When the HF solvent was pumped off at  $-30$  °C from a freshly prepared  $NF_4HF_2-HOIF_4O$  solution, a white solid residue was obtained. The low-temperature Raman spectrum of this solid showed the presence of the  $NF_4$ <sup>+</sup> cation, but the remaining bands were too broad to permit a positive distinction among  $IF_4O_2$ , HOIF<sub>4</sub>O, and possibly some  $HF_2^- \cdot nHF$ <sup>3</sup>. The new compound FOIF40 was obtained in high yield by decomposing at room temperature this thermally unstable solid, with the byproduct being NF<sub>3</sub>. Since the same products were obtained from HF solutions that, on the basis of their <sup>19</sup>F NMR and Raman spectra, contained only  $HOIF_4O$  but not  $IF_4O_2^-$ , it appears

**<sup>(39)</sup> Rose, W. B.; Nebgen, J. W.; Metz, F.** I. *Rev. Sci. Instrum.* **1966,37,** 

**<sup>238.</sup>  (40) Shamir, J.; Hyman, H. H.** *Specrrochim. Acta, Part A* **1967,234 1899.** 

that  $FOIF_4O$  is formed by fluorination of  $HOIF_4O$  by either NF4+ or nascent fluorine formed during the thermal decomposition of the marginally stable  $NF_4 + HF_2 - nHF^3$  Consequently, it was interesting to investigate whether FOIF<sub>4</sub>O could also be obtained by the fluorination of HOIF<sub>4</sub>O with elemental fluorine. However, fluorination reactions carried out at **25**  <sup>o</sup>C with the use of either neat or HF solutions of HOIF<sub>4</sub>O, **2** atm of F2 pressure, and a shaker for agitation did not result in any fluorination of  $HOIF<sub>4</sub>O$ , and only unchanged starting materials were recovered.

Since the fluorination reactions of alkali-metal salts such as  $CsSF<sub>5</sub>O$ ,  $CsCF<sub>3</sub>O$ ,  $CsClO<sub>4</sub>$ , or  $KNO<sub>3</sub>$  with elemental fluorine yield the corresponding hypofluorites,<sup>1</sup> it was interesting to study the analogous fluorination reaction of  $CsIF<sub>4</sub>O<sub>2</sub>$ . In static systems up to  $60 °C$  slow reactions between  $CsIF_4O_2$ and  $F_2$  were observed, producing IF<sub>5</sub>O in low yield as the only volatile product. Since  $IF<sub>5</sub>O$  is the primary decomposition product of FOIF40 (see below), the intermediate formation of some FOIF40 in this reaction cannot be ruled out. Similarly, the fluorination of  $CsIO<sub>4</sub>$  with  $F<sub>2</sub>$  under comparable conditions produced small amounts of  $IF<sub>5</sub>O$  as the only volatile product. The Raman spectra of the solid residues from both reaction systems showed the presence of  $CsIF<sub>8</sub>$ <sup>41</sup>  $CsIF<sub>6</sub>$ <sup>22</sup> *cis*and trans-CsIF<sub>4</sub>O<sub>2</sub>, and CsIF<sub>4</sub>O.<sup>26</sup> The low reactivity of the I= $O$  double bond in IF<sub>5</sub>O was further demonstrated by separate experiments, showing that  $F_2$  is not added across the  $I=O$  double bond, even in the presence of CsF as a catalyst, at temperature between -196 and +25 °C with the use of an excess of  $F_2$ .

**Properties of** FOIF40. As shown by NMR and vibrational spectroscopy (see below),  $FOIF<sub>4</sub>O$  exists in the form of two isomers, one in which the two oxygens are cis and one in which they are trans to each other. Attempts were unsuccessful to separate the two isomers by gas chromatography at 25 °C using a 30-ft, 3/16-in. 0.d. stainless-steel column containing **50%**  Halocarbon oil No. **4-llV** on Kel-F **300 (70-80** mesh).42 Consequently, the physical properties could only be determined for a mixture of both isomers. On the basis of their  $^{19}$ F NMR peak areas, the ratio of cis to trans isomers in the sample used for the physical property measurements was 1.92:1. FOIF<sub>4</sub>O is colorless as a gas, pale yellow as a liquid, and white in the solid state. The given sample melted at -33.1 °C. Vapor pressures were fitted by the method of least squares to the equation

log *[P* (mm)] = **7.62925** - **1432.0/[T** (K)]

the index of correlation being **0.99991.** The extrapolated boiling point is 28.37 °C. Measured vapor pressures at the noted temperatures are as follows  $[T(^{\circ}C), P^{(mm)}]: -45.3$ , **22; -33.1, 47; -23.0, 80; -13.7, 129;** 0, **244.** The latent heat of vaporization of FOIF40 **is655** kcal/mol and the derived Trouton constant is **21.73,** indicating little association in the liquid phase. This is in agreement with the relatively low boiling point and the small changes between the vibrational spectra of the gas and the liquid (see below). The molecular weight was determined from the vapor density and found to be **254.5** (calcd for FOIF40 **253.9).** The good agreement indicates little or no association in the gas phase at the pressure used  $(P \sim 1 \text{ atm})$ .

FOIF4O is marginally stable at room temperature and can be handled in well-passivated metal and Teflon equipment without rapid decomposition. The fact that  $IF<sub>5</sub>O$  was frequently observed as an impurity in the vibrational and NMR spectra suggests the primary decomposition mode<br>  $FOIF_4O \rightarrow IF_5O + \frac{1}{2}O_2$ 

$$
FOIF_4O \rightarrow IF_5O + \frac{1}{2}O_2
$$

When a sample of  $FOIF_4O$  has heated in a stainless-steel cylinder to 120 °C for 388 h, decomposition to IF<sub>5</sub> and O<sub>2</sub> was observed. This is not surprising in view of a previous report<sup>13</sup> that IF,O readily decomposed to IF, and *02.* **As** expected for a hexacoordinated iodine species, FOIF40 is neither a good fluoride ion acceptor nor a good donor. Thus, it does not form stable adducts at room temperature with either the strong Lewis acid  $SbF_5$  or the strong Lewis base CsF. Attempts to add FOIF<sub>4</sub>O across the C=C double bond in  $C_2F_4$  were unsuccessful. Fluorination and oxygenation of  $C_2F_4$  occurred with  $\text{COF}_2$ , CF<sub>3</sub>CFO, and C<sub>2</sub>F<sub>6</sub> being the principal reaction products.

different temperatures. The signals due to the **0-F** fluorines are given at a 10 times wider scale than those due to fluorines on iodine. Positive

shifts are downfield from the external CFCl<sub>3</sub> standard.

I9F **NMR Spectra of** FOIF40. The 19F NMR spectra of FOIF40 were recorded for the neat material and HF solutions and were essentially identical. The spectra of the neat liquid and solid are shown in Figure **4,** together with the observed chemical shifts and coupling constants. Peak-area measurements showed that the **202** and **67** ppm signals belong to an  $AX_4$  and the 176, 175, and 77 ppm signals to an  $A_2BCX$ system. The AX<sub>4</sub> system is readily assigned to the trans isomer









**Figure 5.** Vibrational spectra of a mixture of *cis-* and *trans-FOIF<sub>4</sub>O* in a ratio of about 1.9:l. Traces **A** and B are infrared spectra **of** the gas, recorded at pressures of 10 and 95 mm, respectively, in a 5-cm path length cell equipped with **AgCl** windows. Most of the absorption below **400** cm-' is due to the window material. Trace C is an infrared spectrum of FOIF<sub>4</sub>O isolated in a neon matrix [mole ratio (MR) loOO:l] and recorded at *6* **K.** Traces D-F are Raman spectra of liquid FOIF<sub>4</sub>O recorded in 4-mm o.d. quartz tubes at  $-20$  °C for two samples containing somewhat different ratios of cis (c) to trans (t) isomers, with the incident polarization parallel and perpendicular.

is similar to those of other hypofluorites such as  $O<sub>3</sub>CIOF(219)$ is similar to those of other hypofluorites such as  $O_3CIOF$  (219 = 14 Hz, and  $J_{cx}$  being too small to be resolved. When the ppm),<sup>43</sup> SF<sub>3</sub>OF (189 ppm),<sup>44</sup> or *trans*-SeF<sub>4</sub>(OF)<sub>2</sub> (179 ppm).<sup>45</sup> solution is cooled, th The fluorine-fluorine coupling constant,  $J_{FF} = 34$  Hz, is in good agreement with the value of 27 **Hz** previously reported for the coupling constant of the four equatorial fluorines to the two hypofluorite fluorines in trans-Se $F_4(OF)_2$ .<sup>45</sup> The broadened fluorine on iodine resonance and the lack of observable fine structure of  $FOIF<sub>4</sub>O$  is attributed to unresolved IF coupling  $(I_1 = \frac{5}{2})$  due to decreased quadrupole relaxation caused by the approximately spherically symmetric electric



Figure 6. Infrared spectra of IF<sub>5</sub>O and FOIF<sub>4</sub>O in a Ne matrix (MR) 1OOO:l) at *6* K recorded at 20-fold scale expansion. The bands due to  $IF<sub>5</sub>O$  in the FOIF<sub>4</sub>O spectrum are marked by an asterisk.



Figure 7. Infrared spectra of IF<sub>5</sub>O and FOIF<sub>4</sub>O in a Ne matrix (MR 1000:1).

field about the iodine.<sup>6</sup> The  $A_2BCX$  system is assigned to the cis isomer



and also shows an unresolved IF signal (at about 77 ppm) and a resolved hypofluorite signal (at about 202 ppm), which exhibits a pronounced temperature dependence. At 40 °C, the OF signal approximates a first-order doublet  $(J = 88Hz)$ of triplets  $(J = 14Hz)$ . Since coupling should be stronger to the cis fluorines than to the trans fluorine,<sup>44</sup> this spectrum could then be interpreted as being due to the cis isomer with free rotation around the I-O single bond and  $J_{BX} = 88$  Hz,  $J_{AX}$ solution is cooled, the line width becomes smaller and the CX coupling becomes observable. **As** can be seen from Figure **4,**  the spectrum exhibits pronounced second-order effects, and a computer-aided analysis will be required to obtain precise coupling constants. The fact that the unresolved fluorine on iodine signal has a significantly larger line width for the cis than for the trans isomer is not surprising because the cis isomer possesses three similar but nevertheless nonequivalent types of fluorine on iodine.

**Vibrational Spectra of FOIF<sub>4</sub>O and IF<sub>5</sub>O.** The infrared spectra of the gas and of the neon matrix isolated solid and the Raman spectra of liquid and solid FOIF40 were recorded (see Figures 5-7), and the observed frequencies are summa-

**<sup>(43)</sup> Christe, K. O., unpublished results.** 

**<sup>(44)</sup> Harris, R. K.; Packer, K. J.** *J. Chem.* **SOC. 1962, 3077. (45) Smith, J. E.; Cady, G. H.** *Inorg. Chem.* **1970,** *9,* **1293.** 





 $a$  Mixture of cis and trans isomers.  $b$  Uncorrected Raman intensities based on trace E of Figure **5.** 

rized in Table V. The studied samples were mixtures of cisand trans-FOIF<sub>4</sub>O with a cis: trans ratio of about 1.9 based on the NMR spectra and, in the matrix study, also contained a small amount of  $IF<sub>5</sub>O$ , formed during manipulation of the sample. Since the vibrational spectra of *cis*- and *trans*-FOIF<sub>4</sub>O and of  $IF<sub>5</sub>O$  (see Figure 8) are all very similar, the gas-phase infrared spectra are only of limited value for distinguishing the three compounds. However, the Raman spectra of the liquid and solid and particularly the infrared spectra of the matrix-isolated samples definitely confirm the presence of the two FOIF<sub>4</sub>O isomers established by the <sup>19</sup>F NMR study. Some distinction of the cis from the trans isomer bands was possible from a comparison of spectra of samples having different cis to trans ratios (see for example traces D and **E**  of Figure *5).* 

Tentative assignments for the stretching modes of *cis-* and trans-FOIF40 are given in Table V and were made by comparison with those established for  $IF_5O^{14,29}$  and  $IF_4O_2^-$  (see above), relying mainly on the observed relative infrared and Raman intensities.

The vibrational spectra observed for  $IF<sub>5</sub>O$  are in excellent agreement with those previously reported, $^{14,29}$  except for the fact that our spectra do not show a strong infrared band at  $640 \text{ cm}^{-1}$ . As previously suggested,<sup>29</sup> this band is due to IF<sub>5</sub>, the principal decomposition product of  $IF<sub>5</sub>O$ .

Mass Spectra. The recording of the mass spectrum of FOIF40 presented difficulties due to reaction of the compound with the inlet system, resulting in the formation of some  $IF<sub>5</sub>O$ . Furthermore,  $I_2$  has almost the same mass  $(253.8)$  as the parent FOIF40 **(253.9)** thus making a distinction of the two



**Figure 8.** Vibrational spectra of IF<sub>5</sub>O. Trace A is infrared spectra of the gas, trace B is an infrared spectrum in a Ne matrix, and trace C is a Raman spectrum of the liquid. All were recorded under conditions identical with those of Figure *5.* 

Table VI. Mass Spectrum<sup>a</sup> of a Mixture of *cis*- and **trans-FOIF,O** 

m/e	rel intens	ion	m/e	rel intens	ion
219	72	$IF_4O^+$	165	18	$IF_2$ <sup>+</sup>
203	100	$IF_4^+$	162	38	$IOF^*$
200	32	$IF3O+$	146	15	$IF+$
184	18	IF,	143		IO <sup>+</sup>
181	73		127	38	Ţ÷
178		$IF2O+$ $IO2F+$			

Recorded with an ionization potential of **70** eV, with the use of a **1: 1** mixture of **FOIF,O** and **CIF,.** Peaks due to **ClF,** and IF,O have been subtracted from the pattern.

molecules difficult. These problems were overcome by recording the spectra of pure  $IF<sub>5</sub>O$  under the same conditions and subtracting the IF<sub>5</sub>O pattern from that of the FOIF<sub>4</sub>Ocontaining sample. The interference from  $I_2$  was eliminated by recording spectra of 1:1 mixtures of  $CIF_3$  and  $FOIF_4O$ . The  $CIF<sub>3</sub>$  oxidized I<sub>2</sub> rapidly to iodine fluorides but did not appear to interact with  $FOIF_4O$ . The mass cracking pattern obtained in this manner for  $FOIF_4O$  is listed in Table VI and agrees with the expectations<sup>46</sup> for a hypofluorite. The I-OF single bond is readily broken to yield an intense  $IF_4O^+$  fragment, which can undergo additional oxygen and/or fluorine loss.

Synthesis of CIOIF<sub>4</sub>O. Since FOIF<sub>4</sub>O was found to be stable, the synthesis of the analogous hypochlorite,  $CIOIF<sub>4</sub>O$ , appeared feasible. Using  $CsIF_4O_2$  and  $CIOSO_2F$ , a generally

useful reagent for the syntheses of hypochlorites, $47$  the synthesis of ClOIF4O was accomplished according to Inorg. Chen<br>
ul reagent for the syntheses of hypochlorites,<sup>47</sup> the synth<br>
CIOIF<sub>4</sub>O was accomplished according to<br>
CsIF<sub>4</sub>O<sub>2</sub> + ClOSO<sub>2</sub>F  $\frac{-78 \text{ °C}}{1000}$  CsSO<sub>3</sub>F + ClOIF<sub>4</sub>O<br>
resulting ClOIF<sub>4</sub>O appears to be highl

$$
CsIF_4O_2 + CIOSO_2F \xrightarrow{-78 \text{ °C}} CsSO_3F + CIOIF_4C
$$

The resulting CIOIF<sub>4</sub>O appears to be highly reactive, difficult to handle, and thermally unstable. Consequently, the compound could not be well characterized. The main evidence for its existence is the infrared spectrum of the gas, which is similar to that of  $FOIF_4O$  except that the  $O-F$  stretch is replaced by a band at  $763 \text{ cm}^{-1}$ , characteristic of an  $O$ —Cl stretch,<sup>47</sup> and the I=O, IF, and I-O stretching modes are shifted to slightly lower frequencies. The compound decomposes to  $IF_5$ , and attempts to add it across the  $C=C$  double bond of  $C_2F_4$  did not result in stable adducts.

**Conclusion.** Although the isolation of  $NF_4$ <sup>+</sup> salts of either  $IO_4^-$  or  $IF_4O_2^-$  was not possible, solutions containing  $NF_4^+$  and  $HOIF<sub>4</sub>O$  were found to decompose to produce  $FOIF<sub>4</sub>O$  in high yield. This is in marked contrast to the similar syntheses of FOClO<sub>3</sub> and FOSO<sub>2</sub>F where the corresponding  $NF_4$ <sup>+</sup>ClO<sub>4</sub><sup>-3</sup>

(47) Schack, C. J.; Christe, K. 0. *Isr. J. Chem.* 1978, *17,* 20.

and  $NF_4$ <sup>+</sup>SO<sub>3</sub>F<sup>-4</sup> salts were shown to be the actual intermediates. FOIF<sub>4</sub>O is the first known example of an iodine hypofluorite and exists as cis and trans isomers. It is a stable compound and was thoroughly characterized. The analogous hypochlorite, ClOIF<sub>4</sub>O, was also prepared for the first time but, as expected, is considerably less stable than FOIF40. The reaction of  $CsIO<sub>4</sub>$  with HF was found to be a convenient synthesis of  $CsIF<sub>4</sub>O<sub>2</sub>$ , which, by reaction with  $BIF<sub>5</sub>$  in HF, can readily be converted into HOIF<sub>4</sub>O, thus providing easy access to tetrafluoroperiodates. The bonding in trans-IF<sub>4</sub>O<sub>2</sub><sup>-</sup> was studied by vibrational spectroscopy, and the results of a normal-coordinate analysis are in excellent agreement with the trends previously established<sup>37</sup> for chlorine oxyfluorides.

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**Registry No.** trans-CsIF<sub>4</sub>O<sub>2</sub>, 77224-44-3; cis-CsIF<sub>4</sub>O<sub>2</sub>, 55188-51-7; trans-HOIF<sub>4</sub>O, 25685-16-9; cis-HOIF<sub>4</sub>O, 25685-15-8; trans-FOIF<sub>4</sub>O, 72151-31-6; cis-FOIF,O, 72123-55-8; C101F40, 77224-34-1; CsIO,, 13478-04-1; HF, 7664-39-3; BrF<sub>5</sub>, 7789-30-2; ClF<sub>3</sub>, 7790-91-2; ClF<sub>5</sub>, 13637-63-3; F<sub>2</sub>, 7782-41-4; BiF<sub>5</sub>, 7787-62-4; NF<sub>4</sub>SbF<sub>6</sub>, 16871-76-4;  $CISO<sub>3</sub>F$ , 13997-90-5; IF<sub>5</sub>O, 16056-61-4.

Contribution from the Department of Chemistry, University of Missouri, Columbia, Missouri 65201, and the Chemistry Division, Argonne National Laboratory, Argonne, Illinois 60439

## **Oxidation by Aqueous Fluoroxysulfate: Catalysis by Silver(1)** '\*

RICHARD C. THOMPSON<sup>1b</sup> and EVAN H. APPELMAN\*<sup>1c</sup>

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The oxidations of the ions  $Cr^{3+}$ ,  $Co^{2+}$ , and  $VO^{2+}$  by the fluoroxysulfate ion,  $SO_4F$ , in aqueous solution are catalyzed by Ag<sup>+</sup>. The rate-determining step for all three catalyzed reactions is the bimolecular oxidation of Ag<sup>+</sup> by SO<sub>4</sub>F<sup>-</sup>, which has a rate constant of (1.3  $\pm$  0.2)  $\times$  10<sup>3</sup> M<sup>-1</sup> s<sup>-1</sup> at 17 °C. Activation parameters for this reaction are  $\Delta H^* = 6.1 \pm 0.5$  kcal/mol and  $\Delta S = -23 \pm 2$  cal/(mol deg). In the absence of Ag<sup>+</sup>,  $\text{Co}^{2+}$  and VO<sup>2+</sup> react very slowly with  $\text{SO}_4\text{F}$ <sup>-</sup>, while  $\text{Cr}^{3+}$  does not react at all. Despite its high thermodynamic oxidizing power, the fluoroxysulfate ion acts as a very selective oxidant.

## **Introduction**

The fluoroxysulfate ion, **S04F,** is the only known ionic hypofluorite.2 It is also one of the most powerful of all aqueous oxidants, with a standard electrode potential of 2.46 **V.3** The rubidium and cesium fluoroxysulfates are rather easily made and are sufficiently stable that they can be conveniently stored prior to use. These salts are therefore very suitable substances with which to explore and take advantage of the chemical properties of a hypofluorite, which we may expect to act as a vigorous oxidizing and fluorinating agent.

In the course of our initial survey of the reaction chemistry of aqueous fluoroxysulfate,<sup>2,4</sup> we have observed that  $SO_4F^$ readily oxidizes most reducing substrates. However, some reducing agents such as  $Cr^{3+}$  are not oxidized at all, while others such as **Mn2+,** Ce3+, Co2+, and **V02+** react rather sluggishly. On the other hand, Ag<sup>+</sup> is very rapidly oxidized by **SO4F,** and since higher oxidation states of silver are known to be powerful oxidants, it appeared to us that Ag<sup>+</sup> should catalyze oxidation by **S04F** and might thereby extend the application of the latter as an oxidizing agent. We have found that such catalysis does indeed take place, and we have un-

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- (4) Thompson, R. C.; Appelman, E. H. *Inorg. Chem.* 1980, *19,* 3248.

dertaken to study the Ag+-catalyzed reactions of **S04F** with Cr3+, Co2+, and **V02+.** For the sake of comparison, we have also examined the slow uncatalyzed reactions of **S04F** with  $Co<sup>2+</sup>$  and  $VO<sup>2+</sup>$ .

## **Experimental Section**

**Materials.** Cesium fluoroxysulfate was prepared and purified by techniques described previously.<sup>2,4</sup> Solutions of VO(ClO<sub>4</sub>)<sub>2</sub> in 1 M HC104 were prepared from commercial vanadyl sulfate by ion exchange. Stock solutions of lower acidity were obtained by partial neutralization of the acid with lithium carbonate. Zinc perchlorate was prepared by the reaction of Johnson-Matthey high-purity ZnO with HCIO<sub>4</sub>, followed by two recrystallizations. Other starting materials such as G. F. Smith  $Zn(C1O<sub>4</sub>)<sub>2</sub>$ , reagent grade ZnO, and reagent grade Zn metal resulted in  $Zn(C1O<sub>4</sub>)<sub>2</sub>$  solutions containing impurities that rapidly reduced or decomposed fluoroxysulfate. The preparations of  $Co(CIO_4)_2$ ,  $Cr(CIO_4)_3$ , and  $LiClO_4$  have been described elsewhere.<sup>5</sup> Other materials were commercial products of reagent grade. Deionized water was distilled before use first from acid dichromate and then from alkaline permanganate.

Solutions of  $HSO<sub>5</sub>$  in which the terminal peroxide oxygen was enriched in *'\*O* were prepared by allowing cesium fluoroxysulfate to decompose in a 1 M HCIO, solution that was enriched in **lSO.** The hydrogen peroxide formed was decomposed by addition of a stoichiometric amount of Ce(IV). The yield of peroxymonosulfate in 1 **M** HClO, was of the order of 40% of the fluoroxysulfate. This is considerably higher than the yield obtained at lower acidity.<sup>4</sup>

**Analytical Procedures.** Solutions of  $Co^{2+}$ ,  $Co^{3+}$ ,  $VO^{2+}$ ,  $VO_2^+$ , and  $HCrO<sub>4</sub>$  were analyzed spectrophotometrically, with use of extinction

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<sup>(2)</sup> Appelman, E. H.; Basile, L. J.; Thompson, R. C. *J. Am. Chem. SOC.*  1979, *101,* 3384.

*<sup>(5)</sup>* Sullivan, J. C.; Thompson, R. C. *Inorg. Chem.* 1967, 6, 1795.