COMPONENTS :	ORIGINAL MEASUREMENTS:
<ul> <li>(1) Holmium fluoride; HoF<sub>3</sub>; [13760-78-6]</li> <li>(2) Alcohols</li> </ul>	Kirmse, E.M. Wiss. Hefte, Paed. Inst. Koethen <u>1978</u> , 2, 85-90.
VARIABLES:	PREPARED BY:
Room temperature	T. Mioduski and M. Salomon
EXPERIMENTAL VALUES:	

			HoF <sub>3</sub> solubi	ility <sup>a,b</sup>
solvent			mass %	mol kg <sup>-1</sup>
methanol	сн <sub>4</sub> 0;	[67-56-1]	0.01	$4.5 \times 10^{-4}$
ethanol	с <sub>2</sub> н <sub>6</sub> 0;	[64-17-5]	0.01	$4.5 \times 10^{-4}$

<sup>a</sup>Molalities calculated by the compilers.

 $^{\rm b}{\rm Solid}$  phases were dried in a desiccator over  ${\rm P_40}_{10}$  and the Ho:F ratio found to equal almost 1:3.

ATTYTTTARY	INFORMATION
VOVIDIUVI	THIOWHITOW

METHOD/APPARATUS/PROCEDURE: Isothermal method. About 100 mg of HoF <sub>3</sub> was	SOURCE AND PURITY OF MATERIALS: $Ho_2O_3$ (source and purity not specified) was dissolved in HCL and the fluoride precipi-
ture mechanically agitated at room tempera- ture for 100 h. 5-10 g of saturated solu- tion were removed by decanting or by centri- fuging, and the solution evaporated to dry- ness. The residue was heated with about 10	tated by addition of aq HF. The solid pro- duced was HoF3.0.5H <sub>2</sub> O and was dehydrated by washing with acetone followed by drying at 310°C for 120 hours.
$cm^3$ of 10% KOH solution for 1-2 h to obtain solid Ho(OH) <sub>3</sub> and a basic F solution. The precipitate was washed, dissolved in aq HCl, and Ho determined several times by complexo- metric titration with potentiometric end-	The solvents were dried and purified by "standard methods."
point detection (1). The fluoride content in the filtrate was determined photometri- cally using Al-Eriochrome cyanine color lake indicator.	ESTIMATED ERROR: Soly: results with relative errors exceeding 50% were rejected. Temp: unknown.
The reported solubility is a mean of	
"numerous parallel determinations," or "at least two parallel determinations."	REFERENCES: 1. Schilbach, U.; Kirmse, E.M. Z. Chem. <u>1974</u> , 14, 484.
	<ol> <li>Schilbach, U.; Hetze, I.; Kirmse, E.M. Chemia Analityczna <u>1975</u>, 20, 33.</li> </ol>

COMPONENTS:	ORIGINAL MEASUREMENTS -
(1) Holmium fluoride: HoFa:	Dressler, H.
[13760-78-6]	
	Dissertationschrift. Paed. Inst. Koethen.
(2) Ethers	GDR. <u>1980</u> .
VARIABLES:	PREPARED BY:
	T. Mioduski and M. Salomon
EXPERIMENTAL VALUES:	
	solubility
1	301001111y
solvent	mass % mol/100 g sin
l-methoxydecane; n-C <sub>1</sub>	$_{1H_{2}}0;$ [7289-52-3] 0.03 <sup>a</sup> 1.35 x 10 <sup>-4</sup>
	1 24
	v and (and (and )) a and a contract
1-(chloromethoxy)butane; n-C5	$H_{11}^{C10}$ [2351-69-1] 0.02 9.0 x 10
a In the solid phase the Ho:F:ether	
In the borra phase the norricther	2 1410 15 1.2.5510.02.0.40
b. In the solid phase the Ho:F:ether	ratio is 1:2.89:0.06.
	AUXILIARY INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS;
Method analogous to that described	in (1). It appears that the fluoride was prepared
No other information available.	as in (1). In spite of drying the
	Ho:F:H=0 ratio was 1:3.04:0.70
	No other information available.
	ESTIMATED ERROR:
	Nothing specified.
	Nothing opecities.
1	REFERENCES:
	1 Kirmse F M
	Wiss. Hefte. Paed. Inst. Koethen.
	<u>1978,</u> 2, 85.
1	

COMPO	NENTS:	ORIGINAL MEASUREMENTS:
(1)	Holmium fluoride; HoF <sub>3</sub> ; [13760-78-6]	Kirmse, E.M.
(2)	Tributyl phosphate; C <sub>12</sub> H <sub>27</sub> O <sub>4</sub> P; [126-73-8]	Wiss. Hefte, Paed. Inst. Koethen <u>1978</u> , 2, 85-90.
VARIA	BLES:	PREPARED BY:
Room	temperature	T. Mioduski

## EXPERIMENTAL VALUES:

The solubility of  $HoF_3$  in  $[CH_3(CH_2)_3]_3P(0)$  at room temperature was given as

0.02 mass %

The corresponding molality calculated by the compiler is

 $9.0 \times 10^{-4} \text{ mol kg}^{-1}$ 

The solid phase was dried in a desiccator over  ${\rm P_40}_{10}$  and the Ho:F ratio determined to be almost 1:3.

AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE: Isothermal method. About 100 mg of HoF <sub>3</sub> was added to 10-20 cm <sup>3</sup> of solvent, and the mixture mechanically agitated at room temp- erature for 100 h. 5-10 g of saturated sol- ution were removed by decanting or by cen- trifuging, and the solution evaporated to dryness. The residue was heated with about 10 cm <sup>3</sup> of 10% KOH solution for 1-2 h to obtain solid Ho(OH) <sub>3</sub> and a basic F <sup>-</sup> solu- tion. The precipitate was washed, dissolved in aq HCl, and Ho determined several times by complexometric titration with potentio- metric end-point detection (1). The fluoride content in the filtrate was determined photometrically using Al-Eriochrome cyanine color lake indicator (2).	<pre>SOURCE AND PURITY OF MATERIALS: Ho203 (source and purity not specified) was dissolved in HCl and the fluoride precipi- tated by addition of aq HF. The solid pro- duced was HoF3.0.5H20 and was dehydrated by washing with acetone followed by drying at 310°C for 120 hours. The solvent was dried and purified by "standard methods."</pre> ESTIMATED ERROR: Soly: results with relative errors exceed- ing 50% were rejected. Temp: unknown.
"numerous parallel determinations," or "at least two parallel determinations."	<ul> <li>REFERENCES:</li> <li>1. Schilbach, U.; Kirmse, E.M. Z. Chem. <u>1974</u>, 14, 484.</li> <li>2. Schilbach, U.; Hetze, I.; Kirmse, E.M. Chemia Analityczna <u>1975</u>, 20, 33.</li> </ul>

.

COMPO	NENTS:	ORIGINAL MEASUREMENTS:
(1) (2)	Holmium fluoride; HoF <sub>3</sub> ; [13760-78-6] Dimethylsulfoxide; C <sub>2</sub> H <sub>6</sub> OS; [67-68-5]	Kirmse, E.M. Wiss. Hefte, Paed. Inst. Koethen <u>1978</u> , 2, 85-90.
VARIA	BLES :	PREPARED BY:
Room	Temperature	T. Mioduski
EXPER	RIMENTAL VALUES:	com temperature was given as
		0.02 mass %
The c	orresponding molality calculated b	y the compiler is
		9.0 x $10^{-4}$ mol kg <sup>-1</sup>
The se almost	olid phase was dried in a desiccat t 1:3.	for over $P_4O_{10}$ and the Ho:F ratio found to be

SOURCE AND PURITY OF MATERIALS: Ho <sub>2</sub> O <sub>3</sub> (source and purity not specified) was dissolved in HCl and the fluoride pre- cipitated by addition of aq HF. The solid produced was HoF <sub>3</sub> .0.5H <sub>2</sub> O and was dehydrated by washing with acetone followed by drying at 310°C for 120 hours. The solvent was dried and purified by "standard methods."
ESTIMATED ERROR: Soly: results with relative errors exceed- ing 50% were rejected. Temp: nothing specified.
<pre>REFERENCES: 1. Schilbach, U.; Kirmse, E.M. Z. Chem. <u>1974</u>, 14, 484. 2. Schilbach, U.; Hetze, I.; Kirmse, E.M. Chemia Analityczna <u>1975</u>, 20, 33.</pre>