Thermal behavior and other properties of Pr(III), Sm(III), Eu(III), Gd(III), Tb(III) complexes with 4,4′-bipyridine and trichloroacetates

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Abstract A novel mixed-ligand complexes with empirical formulae: Ln(4-bpy)1.5(CCl3COO)3·nH2O (where Ln(III) = Pr, Sm, Eu, Gd, Tb; n = 1 for Pr, Sm, Eu and n = 3 for Gd, Tb; 4-bpy = 4,4′-bipyridine) were prepared and characterized by chemical, elemental analysis and IR spectroscopy. Conductivity studies (in methanol, dimethylformamide and dimethylsulfoxide) were also described. All complexes are crystalline. The way of metal–ligand coordination was discussed. The thermal properties of complexes in the solid state were studied under non-isothermal conditions in air atmosphere. During heating the complexes decompose via intermediate products to the oxides: Pr6O11, Ln2O3 (for Sm, Eu, Gd) and Tb4O7. TG-MS system was used to analyze principal volatile thermal decomposition and fragmentation products evolved during pyrolysis of Pr(III) and Sm(III) compounds in air.

Keywords Lanthanide complexes · 4,4′-bipyridine · Trichloroacetates · Thermal decomposition

Introduction

This work is a continuation of our previous studies on synthesis, properties, and thermal decomposition of metal complexes with bipyridine isomers and carboxylates [1–7]. Lanthanide compounds are curious for many research workers in last years because of their variety applications [8–13]. Halogenoacetates complexes show varied types of coordination, therefore they are interesting ligands.

Authors [14–18] isolated compounds type: Ln(CCl2HCOO)3·2H2O (where Ln(III) = Pr, Er), Ln(CCl3COO)3·2H2O (where Ln(III) = Pr, Nd, Eu, Dy, Yb), [Nd(CCl3HCOO)6(H2O)3]·mH2O and Nd,Ln1−x(CCl3COO)2·2H2O. In 2003 year [Tb2(CH3COO)6(H2O)4] complexes were produced [19]. Rohde and Urland [20] obtained monocrystals Ln(CClF2COO)3·3H2O (where Ln(III) = Gd, Dy, Ho and Er). There is report about crystal structure of neodymium(III) compounds with monochloroacetate [21]. Imai et al. [22] synthesized and characterized [Ln(CCl3HCOO)3(H2O)]n (where Ln(III) = La, Pr, Nd, Eu) and their properties were measured.

There are scant information about lanthanide complexes with bipyridine isomers and halogenoacetates. Authors [23, 24] synthesized and determined crystal structure of compounds type: [Ln(CCl2HCOO)3(2-bpy)]n (where Ln(III) = Nd, Pr; 2-bpy = 2,2′-bipyridine). Single crystals of [Gd2(CClF2COO)6(H2O)2(2-bpy)]2·C7H2OH have been obtained [25]. Crystal structure and magnetic properties of the gadolinium complexes type: Gd2(CClH2COO)6(2-bpy)2 [26] and Gd2(CCl3COO)6(2-bpy)2(H2O)2·4(2-bpy) [27] were measured. Authors [27] described also compounds Ln(CCl3COO)3(2-bpy)2 (Ln(III) = Pr, Nd). Spacu and Antonescu [28] reported thermal stability and some properties of complexes [LnA3(N-donors)]·nH2O (where Ln(III) = La→Eu (expect Pm), Dy and Er; A = chloroacetates; N-donors = 1,10-phenantroline, 2,2′-bipyridine and 4,7-diphenyl-1,10-phenantroline). Hart and Laming [29] described synthesis of complexes Ln(2-bpy)X3·nH2O where Ln(III) = La, Pr, Ce, Nd, Sm, Eu, Dy, Er; X = chloroacetates. Kokonov et al. [30] characterized thermal studies of compounds neodymium and erbium with 2-bpy and dibromoacetates.
Owing to two nitrogen donor atoms, 4,4′-bipyridine is used as a potential ligand. This isomer forms polymeric species [31–34]. In combination with carboxylate groups, 4-bpy makes up interesting structures [35–39]. Our studies presented here complete series of lanthanides metal complexes with 4,4′-bipyridine and halogenoacetates [35, 40–44].

Experimental

Materials

All the chemicals and solvents used were of analytical grade. Trichloroacetic acid was obtained from Reanal–Budapest, methanol (MeOH, anhydrous) from Scan–Lab, SmCl3·6H2O from Aldrich, Tb2O7 Koch Light Laboratory England and Pr6O11 Rare Earth Products Limited. 4,4′-Bipyridine, Eu3O3, Gd2O3 and other products were obtained from POCh–Gliwice. Solutions of lanthanide(III) trichloroacetates as hydrated products were prepared by dissolving freshly prepared trihydrated products of lanthanides metal complexes with 4,4′-bipyridine and halogenoacetates [35, 40–44].

Methods and instruments

The contents of Ln(III) ions in obtained solutions of trichloroacetates were complexometrically (EDTA) determined. The mixed-ligand complexes were prepared by mixing 10 mmol of 4-bpy in 96% v/v ethanol (31.25 mL) with the freshly obtained solution of 5 mmol metal trichloroacetates in 8.75 mL of water at room temperature. The equation for synthesis of complexes:

\[
\text{Ln(OH)}_3 + 3\text{CCl}_3\text{COOH} \rightarrow \text{Ln(4-bpy)}_1.5(\text{CCl}_3\text{COO})_3 \cdot n\text{H}_2\text{O}
\]

where Ln(III) = Pr(III), Sm(III), Eu(III), Gd(III), Tb(III).

Synthesis and analysis

During several days the compounds crystallized. The obtained complexes were filtered off; washed with 40% v/v ethanol and then with EtOH and Et2O mixture (1:1) and air dried at room temperature. The contents of N, H, and C in prepared complexes were determined by a Carbo-Erba analyzer with V2O5 as an oxidizing agent; metals(III) in mineralized samples complexometrically.

Methods and instruments

IR spectra were recorded with a NICOLETT 6700 Spectrometer (4000–400 cm⁻¹ with accuracy of recording 1 cm⁻¹) using KBr pellets. Molar conductance was measured on a conductivity meter of the OK-102/1 type equipped with an OK-902 electrode at 298 ± 0.5 K, using 1 × 10⁻⁴ mol L⁻¹ solutions of complexes in methanol (MeOH), dimethylsulfoxide (DMSO) and dimethylformamide (DMF). The thermal properties of complexes were studied by TG, DTG and DTA techniques; TG, DTG and DTA curves were recorded on derivatograph type Q-1500, samples of 100 mg, α-Al2O3 served as reference material in air static atmosphere. From TG, DTA and DTG curves the solid intermediate products decomposition were determined and were confirmed by the IR spectra of sinters. In sinters (prepared during heating of complexes up to temperatures defined from TG curves) the vibration modes of 4-bpy and trichloroacetates were analyzed as well as the presence of anions Cl⁻ was also stated. The TG-MS system, consisted of a TG/DTA-SETSYS-16/18 coupled to a mass spectrometer (QMS-422; ThermoStar from Balzers Instruments, an ion source of ca 423 K by using 70 eV electron impact ionization and flow rate 1 L/h), used to monitored principal volatile species produced during pyrolysis of the Pr(III) and Sm(III) complexes (with samples of mass 4.64 and 4.82 mg, respectively) in dynamic air atmosphere. The m/z values are given based on H, 12C, 14N, 16O, and 35Cl (additionally 13C and 18O in case of CO2). All thermal investigations were carried out in the range of temperature 293–1273 K at a heating rate of 10 K min⁻¹. For all complexes and final solid decomposition products, X-ray diffractograms (D-5000 diffractometer, Ni-filtered CuKα radiation) were done in the range of 2θ angles 2–80°. Obtained results were analyzed using the Powder Diffraction File [45].

Results and discussion

The analytical data are shown in Table 1. These results establish the stoichiometry of these compounds with the general empirical formulae: Ln(4-bpy)₁.₅(CCl₃COO)₃·nH₂O (where Ln(III) = Pr, Sm, Eu) and Ln(4-bpy)₁.₅(CCl₃COO)₃·nH₂O (Ln(III) = Gd, Tb). The observed molar conductivities values in MeOH, DMSO, and DMF are given also in Table 1. Conductivity measurements suggested that in all solutions of obtained compounds have behavior non-electrolytes (they dissociate in limited degree in these solutions) [46]. This indicates that trichloroacetate anions are located in the coordination sphere. All complexes are stable in air at room temperature for ca 5 months. The X-ray diffraction powder patterns (Fig. 1) show that all the compounds have crystalline structure and the crystallinity of these complexes follows the order: Pr > Sm > Gd ≈ Tb > Eu. Difference in the crystallinity of these compounds probably depends on the velocity of the evaporation which
was not controlled. They are isostructural in the groups: Pr, Sm, Eu and Gd, Tb. Some complexes in solid state show fluorescence at room temperature: Eu (intensive pink) and Tb (light green). This fluorescence was observed in the light of an LS/58 quartz lamp at room temperature.

IR spectra

IR spectra of the all obtained complexes exhibit several absorption bands characteristic for 4-bpy and –COO groups. The fundamental vibration modes of 4-bpy and –COO groups for complexes are reported in Table 2. The IR spectrum of free 4,4′-bipyridine undergoes a modification when coordination with a lanthanide. The most characteristic ring vibration modes \( \nu(CC) \), \( \nu(CN) \), \( \nu(CC_{\infty}) \)-\( A_1 \) symmetry and \( \nu(CC) \), \( \nu(CN)-B_1 \) symmetry appear at 1588 and 1530 cm\(^{-1}\) in the free ligand [47]. In the IR spectra of complexes they are observed at 1602–1600 and 1534–1533 cm\(^{-1}\), respectively. The ring deformation modes are between 1002 and 1000 cm\(^{-1}\) shifted to higher frequencies in comparison with free 4-bpy (988 cm\(^{-1}\)). The bathochromic shifts of principal absorption bands suggest that 4-bpy is coordinated to Ln(III) ions [47].

The IR spectra of Ln(4-bpy)\(_{1.5}\)(CCl\(_3\)COO)\(_3\)·nH\(_2\)O (Ln(III) = Pr, Sm, Eu) show asymmetric \( \nu_{as}(COO) \) and symmetric \( \nu_{s}(COO) \) vibration of –COO groups in the range 1671–1646 and 1371–1367 cm\(^{-1}\), respectively. On the grounds of spectroscopic criteria [44, 48–51] it can be stated, that in case of Pr(III), Sm(III) and Eu(III) complexes carboxylate groups are bonded as bidentate chelating ligand (the values of \( \Delta \nu_{as} = \nu_{as} – \nu_{s} \) of these complexes are smaller than for sodium salt \( \Delta \nu_{Na} = 324 \)).

In the case of complexes Ln(4-bpy)\(_{1.5}\)(CCl\(_3\)COO)\(_3\)·3H\(_2\)O (Ln(III) = Gd, Tb) the \( \nu_{s} \) is splitted into doublet. Therefore, we may suppose that non-completely equivalent bands (probably chelating and bridging or tridentate chelating-bridging [52]) between Ln(III) (Gd, Tb) and carboxylate ligands of trichloroacetate ligands are formed. The different types of bonds between lanthanide(III) and carboxylate ligands within one molecule in reported in the literature [19, 21, 32, 33]. A broad band in the water

| Table 1 | Analytical data and molar conductivity in MeOH, DMF, and DMSO for the Ln(4-bpy)\(_{1.5}\)(CCl\(_3\)COO)\(_3\)·nH\(_2\)O |
|---------|--------------------------------------------------|
| Compound (color) | Analysis: found (calculated)% | \( \Lambda_{\text{m}} \) (\( \Omega^{-1} \) cm\(^2\) mol\(^{-1}\); \( c = 1 \times 10^{-3} \) mol L\(^{-1}\)) |
| | Ln | C | N | H | | MeOH | DMF | DMSO |
| (I) Pr(4-bpy)\(_{1.5}\)(CCl\(_3\)COO)\(_3\)·H\(_2\)O (light green) | 16.36 | 28.73 | 5.05 | 1.57 | 58.6 | 19.8 | 14.7 |
| (II) Sm(4-bpy)\(_{1.5}\)(CCl\(_3\)COO)\(_3\)·H\(_2\)O (light yellow) | 17.72 | 28.41 | 4.72 | 1.55 | 56.6 | 20.2 | 9.7 |
| (III) Eu(4-bpy)\(_{1.5}\)(CCl\(_3\)COO)\(_3\)·H\(_2\)O (white) | 16.60 | 28.39 | 4.98 | 1.52 | 55.1 | 17.4 | 7.3 |
| (IV) Gd(4-bpy)\(_{1.5}\)(CCl\(_3\)COO)\(_3\)·3H\(_2\)O (white) | 17.18 | 27.11 | 4.39 | 1.91 | 59.6 | 20.4 | 27.7 |
| (V) Tb(4-bpy)\(_{1.5}\)(CCl\(_3\)COO)\(_3\)·3H\(_2\)O (white) | 17.59 | 27.08 | 4.91 | 1.89 | 55.0 | 18.0 | 14.1 |

Fig. 1 X-ray diffraction patterns for complexes Ln(4-bpy)\(_{1.5}\)(CCl\(_3\)COO)\(_3\)·H\(_2\)O (I–III) and Ln(4-bpy)\(_{1.5}\)(CCl\(_3\)COO)\(_3\)·3H\(_2\)O for (IV) and (V)
stretching region (3550–3200 cm\(^{-1}\)) and only shoulder in the water bending region (ca 1630–1600 cm\(^{-1}\)) are observed for all complexes. In addition, in obtained compounds there are bands in the ranges: 837–683 cm\(^{-1}\) for Pr(III), 839–681 cm\(^{-1}\) for Sm(III), 838–680 cm\(^{-1}\) for Eu(III), 837–680 cm\(^{-1}\) for Gd(III) and 838–681 cm\(^{-1}\) for Tb(III). These absorptions are associated with several vibrations: \(\nu_{CCl}\) (800–550 cm\(^{-1}\)), \(\nu_{as(CCl_3)}\) (849, 833 cm\(^{-1}\)), \(\nu_{s(CCl_3)}\) (746, 685 cm\(^{-1}\)), \(\gamma_{(CH)}\) 4-sub pyridine (810, 745, 733, 672 cm\(^{-1}\)) [47–49].

Table 2  Principal IR bands (cm\(^{-1}\)) for 4-bpy and COO group in obtained complexes

| Assignment of bands | 4-bpy [47] | CC\(_3\)COONa [48] | Complexes of Ln(III) |
|---------------------|-------------|----------------------|---------------------|
|                     |             |                      | Pr                  |
| Coordinated 4,4'-bipyridine modes |             |                      | Sm                  |
| \(\nu(CC, CN, C_{12})\) \(A_1\) | 1588        | –                    | 1602                |
| \(\nu(CC, CN)\) \(B_1\) | 1530        | –                    | 1533                |
| Ring deformation    | 988         | –                    | 1001                |
| Carboxylate group modes |             |                      | Eu                  |
| \(\nu_{as} (COO)\) | –           | 1677                 | 1646                |
| \(\nu_{s} (COO)\) | –           | 1353                 | 1371                |
| \(\Delta\nu = \nu_{as} (COO) - \nu_{s} (COO)\) | –           | 324                  | 275                 |

Table 3  Thermal decomposition data of obtained complexes in air; mass sample 100 mg

| No. | Complex | Range of decomposition/K | DTA peaks/K | Mass loss/% | Intermediate and residue solid products |
|-----|---------|--------------------------|-------------|-------------|----------------------------------------|
| (I) | Pr(4-bpy)\(_{1.5}\)(CCl\(_3\)COO)\(_3\)/C\(_2\)H\(_2\)O | 333–391 | 388 endo | 2.5 | Pr(4-bpy)\(_{1.5}\)(CCl\(_3\)COO)\(_3\) |
|     |         | 391–513 | 503 exo | 43.0 | Pr(4-bpy)\(_{1.5}\)Cl\(_3\) |
|     |         | 513–1053 | 978 exo | 35.0 | Pr\(_3\)O\(_{11}\) |
| (II) | Sm(4-bpy)\(_{1.5}\)(CCl\(_3\)COO)\(_3\)/C\(_2\)H\(_2\)O | 333–418 | 378 endo | 13.0 | Sm(4-bpy)\(_{1.5}\)Cl\(_3\) |
|      |         | 418–523 | 453 exo | 33.0 | Sm(4-bpy)\(_{1.5}\)Cl\(_3\) |
|      |         | 523–668 | 593 endo | 8.5 | Sm\(_2\)O \(^3\) |
|      |         | 668–1053 | 723, 783, 1023 exo | 24.0 | SmOCl |
|      |         | 1053–1213 | 1083 endo | 2.5 | Sm\(_2\)O \(^3\) |
| (III) | Eu(4-bpy)\(_{1.5}\)(CCl\(_3\)COO)\(_3\)/C\(_2\)H\(_2\)O | 333–408 | 373 endo | 19.5 | Eu(4-bpy)\(_{1.5}\)(CCl\(_3\)COO)\(_1.75\)Cl\(_{0.25}\) |
|      |         | 408–468 | 463 exo | 24.0 | Eu(4-bpy)\(_{1.5}\)Cl\(_3\) |
|      |         | 468–718 | 653 endo | 17.0 | Eu\(_2\)O \(^3\) |
|      |         | 718–893 | 703, 838 exo | 15.0 | EuOCl |
|      |         | >893 | 703, 838 exo | 2.5 | Eu\(_2\)O \(^3\) |
| (IV) | Gd(4-bpy)\(_{1.5}\)(CCl\(_3\)COO)\(_3\)/C\(_3\)H\(_2\)O | 333–413 | 368 endo | 21.0 | Gd(4-bpy)\(_{1.5}\)(CCl\(_3\)COO)\(_2\)Cl |
|      |         | 413–483 | 463 exo | 27.0 | Gd(4-bpy)\(_{1.5}\)Cl\(_3\) |
|      |         | 483–763 | 733 exo | 13.0 | Gd\(_2\)O \(^3\) |
|      |         | 763–1043 | 813, 1023 exo | 17.0 | GdOCl |
|      |         | >1123 | 813, 1023 exo | 3.0 | Gd\(_2\)O \(^3\) |
| (V)  | Tb(4-bpy)\(_{1.5}\)(CCl\(_3\)COO)\(_3\)/C\(_3\)H\(_2\)O | 333–403 | 353 endo | 20.0 | Tb(4-bpy)\(_{1.5}\)(CCl\(_3\)COO)\(_2\)Cl |
|      |         | 403–573 | 453 exo | 39.0 | Tb(4-bpy)\(_{1.5}\)Cl\(_3\) |
|      |         | 573–753 | 723 exo | 5.0 | Tb\(_2\)O \(^7\) |
|      |         | 753–988 | 808, 953 exo | 14.0 | TbOCl |
|      |         | >988 | 808, 953 exo | 3.0 | Tb\(_2\)O \(^7\) |

\(a\) Probably via Ln(4-bpy)\(_{1.5}\)-m Cl\(_3\)
Thus, the interpretation of IR spectra in these regions is difficult to discuss.

Thermal decomposition

Thermal decomposition of analyzed complexes in air is a multistage process. The solid intermediate products of pyrolysis were determined from TG and DTG curves. The thermal decomposition data are collected in Table 3. Figure 2 presents, as an example, the thermoanalytical curves of Eu(4-bpy)$_{1.5}$(CCl$_3$COO)$_3$·H$_2$O and Tb(4-bpy)$_{1.5}$(CCl$_3$COO)$_3$·3H$_2$O. All the compounds are stable to 333 K. DTA curves present several endo- and exothermic peaks. The very strong and broad exothermic effects correspond to

Fig. 2 Thermoanalytical curves of complexes Eu(III) and Tb(III) in air; mass sample 100 mg

Fig. 3 Some profiles of ion current detected in the mass spectrometer versus time for complexes Pr(4-bpy)$_1$·(CCl$_3$COO)$_3$·H$_2$O (mass sample 4.64 mg) (I) and Sm(4-bpy)$_{1.5}$ (CCl$_3$COO)$_3$·3H$_2$O (mass sample 4.82 mg) (II), in air, for m/z values; heating rate 10 K min$^{-1}$
oxidation of organic ligands and combustion of remaining organic fragments (Fig. 2).

The complex of Pr(4-bpy)$_{1.5}$(CCl$_3$COO)$_3$·H$_2$O loses water molecule in first stage (333–391 K). The dehydration process is accompanied by small endothermic effects (388 K). The anhydrous compound is stable up to 391 K. Next, total decomposition of trichloroacetates takes place and intermediate specie Pr(4-bpy)$_{1.5}$Cl$_3$ (391–513 K) is formed. DTA curve presents exo peak at 503 K. On temperature elevation (513–1053 K) Pr(4-bpy)$_{1.5}$Cl$_3$ directly decomposes to Pr$_6$O$_{11}$.

The dehydration process of complex Sm(4-bpy)$_{1.5}$(CCl$_3$COO)$_3$·H$_2$O is connected with partial decomposition of trichloroacetate ligands (333–418 K), which are demonstrated by endothermic peak on DTA curve (378 K). Probably Sm(4-bpy)$_{1.5}$(CCl$_3$COO)$_2$Cl$_{0.75}$ is formed. When the temperature raises it converts to Sm(4-bpy)$_{1.5}$Cl$_3$ (exo peak at 453 K). Next the 4-bpy is lost (probably via intermediate species Sm(4-bpy)$_{1.5-\cdots}$Cl$_3$), together with anions (2Cl$^-$), and SmOCl occurs. A constant mass level for pure Sm$_2$O$_3$ appears above 1053 K.

The pyrolysis of Eu(III) compound is multistage. In the ranges of temperature: 333–408, 408–468, and 468–893 K intermediate species Eu(4-bpy)$_{1.5}$(CCl$_3$COO)$_{1.75}$Cl$_{1.25}$, Eu(4-bpy)$_{1.5}$Cl$_3$ and EuOCl (probably via Eu(4-bpy)$_{1.5-\cdots}$Cl$_3$) are created, respectively. On the DTA curve endo- and exothermic peaks exist. Formation of pure Eu$_2$O$_3$ begins above 893 K.

Thermolysis of Gd(4-bpy)$_{1.5}$(CCl$_3$COO)$_3$·3H$_2$O complex starts with dehydration and partial decomposition of trichloroacetates Gd(4-bpy)$_{1.5}$(CCl$_3$COO)$_2$Cl (333–413 K). It converts to Gd(4-bpy)$_{1.5}$Cl$_3$. These processes are accompanied with endo and exo effects at 368 and 463 K, respectively. When the temperature rises, further decomposition of organic ligands is similar to Eu(III) compound. In the range 763–1043 K GdOCl is formed. On DTA curve exo peaks at 813 and 1023 K appear. Above 1123 K pure Gd$_2$O$_3$ exists.

The Tb(4-bpy)$_{1.5}$(CCl$_3$COO)$_3$·3H$_2$O complex decomposes in the temperature range 333–403 K. Dehydration is connected with partial pyrolysis of trichloroacetate ligands and forms intermediate Tb(4-bpy)$_{1.5}$Cl$_3$. This process is characterized by endo effect at 353 K. Next, the step-wise decomposition of organic ligands takes place (453 and 723 K exo peaks). The final solid product of decomposition is Tb$_4$O$_7$ (via TbOCl).
Mass spectrometry

A coupled TG-MS system has been used to study volatile species and fragments evolved during the dynamic thermal decomposition of Ln(4-bpy)_{1.5}(CCl_3COO)_3·H_2O (where Ln(III) = Pr, Sm, Eu, Gd, Tb) in air atmosphere. MS data of Pr(III) and Sm(III) complexes are very similar. They detected several ion signal intensities. Figure 3 presents some profiles of ion current for m/z values detected in the mass spectrometer versus time for these complexes in air. Generally, many signals are observed in the range 373–593 K. The m/z values are given for \(^1\)H, \(^{12}\)C, \(^{14}\)N, \(^{16}\)O (additionally \(^{13}\)C and \(^{18}\)O). The profiles observed for OH\(^+\) and H_2O\(^+\) are ample. They show the mixture a sufficient dehydration by the release of hydrogen in form of crystalline (or coordination of H_2O in first step and the large amount attributed to oxidation of the organic matter above. In gaseous pyrolysis products dominate ion C\(^+\) and CO\(_2\)\(^+\) (m/z=12, 44) with centers at 448, 576, 756, 971 K (Pr(III)) and 402, 486, 733, 998 K (Sm(III)). The first maxima of these species coincide with beginning of trichloroacetate ligands decomposition. Further peaks are connected with the total decomposition of ligands and the burning of organic residues. Probably maximum rate of forming of NO\(^+\) (or CH_2O\(^+\) m/z = 30) at 649 K for Pr(III) and 651 K for Sm(III) complexes (traces of N_2O_3 with m/z = 76 at 590 K for Pr(III) and Sm(III)) are observed. The mass spectrometer was also monitoring of species containing halogens: Cl\(^+\), HCl\(^+\), CCl\(_3\)\(^+\), CH_2Cl\(^+\), CH_3Cl\(^+\), CH_2Cl_2\(^+\) (m/z = 35, 36, 47, 49, 50, 84). Additionally, ion currents with m/z = 27, 29, 43, 70, 72 and 74 (HCN\(^+\), CHO\(^+\), CHNO, \(^{35}\)Cl\(^+\), \(^{35}\)Cl\(^{17}\)Cl\(^+\), \(^{37}\)Cl\(^+\)) were monitored. The ion currents for different fragments are illustrated together with the corresponding TG (Fig. 4).

Conclusions

Now, the new fine-crystalline compounds with stoichiometry: Ln(4-bpy)_{1.5}(CCl_3COO)_3·nH_2O (where Ln(III) = Pr, Sm, Eu, Gd, Tb; n = 1 for Pr, Sm, Eu and n = 3 for Gd, Tb) were isolated. The IR spectra give us the information about different coordination of organic ligands. Resting on \(\Delta v\), we can see that Pr(III), Sm(III) and Eu(III) complexes possess bidentate chelating \(-\text{COO}\) groups. In the case of Gd(III) and Tb(III) probably non-completely equivalent bands between these ions and carboxylate groups are formed. Conductivity data show, that trichloroacetate ligands are inside coordination sphere. All obtained complexes are stable at room temperature. During heating they decompose progressively. Only a monohydrated Pr(III) compound loses water molecule over the range 333–391 K, and next decomposes via Pr(4-bpy)_{1.5}Cl_3 to Pr_6O_11. For the other complexes dehydration is united with partial decomposition of trichloroacetates. When the temperature rises total decomposition of organic ligands takes place. The final solid products are the oxides: Ln_2O_3 (Ln(III) = Sm, Eu, Gd) and Tb_2O_3.

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