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A Terbium Chlorobismuthate(III) Double Salt: Synthesis, Structure, and Photophysical Properties

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Abstract

We report on the structure and luminescence of a double salt trivalent rare earth ion acceptor, Tb$^{3+}$, with octahedral [BiCl$_6$]$^{3-}$ donor clusters. The novel TbBiCl$_6$$\cdot$14H$_2$O (1) was prepared from aqueous BiOCl and TbCl$_3$$\cdot$6H$_2$O. The crystal structure of compound 1 exhibits isolated [BiCl$_6$]$^{3-}$ and [Tb(OH)$_2$)$_8$]$^{3+}$ clusters. Luminescence data show energy transfer from octahedral chlorobismuthate(III) clusters to rare earth metal ions. Density Functional Theory (DFT) calculations show distinctly different emission pathways at high and low excitation energies.

1. Introduction

Rare earth metal ions produce sharp emission bands making them particularly useful in a variety of applications, including optoelectronics, biosensors, and photovoltaic compounds.$^{1-7}$ The challenge in developing viable materials using these ions centers on the difficulty of excitation of the lanthanide $f$ electrons. The $f$ orbitals are held close to the nucleus below the $d$ orbitals due to lanthanide contraction and $f$-$f$ transitions are symmetrically forbidden.$^{8}$ This results in weak emission intensities from these ions. Much research has focused on improving the emission intensity of solid state crystals.$^{9-11}$ Energy transfer to rare earth metal centers is a possible avenue
of approach in achieving bright emission.\textsuperscript{12,13} As an example, work by our group and Leznoff \textit{et al.} have shown that rare earth ion emission can be tuned via changes in the bonding of coordinated d\textsuperscript{10} Au(CN)\textsubscript{2}\textsuperscript{−} ligands.\textsuperscript{1,2,5,14,15} More specifically, we have discovered that the presence or absence of metallophilic Au-Au interactions has an effect on the emission intensity of the rare earth center. This change in emission occurs due to changes to the charge-transfer pathway whereby energy is transferred from the Au(CN)\textsubscript{2}\textsuperscript{−} anion to the rare earth ion.

Currently, our group is interested in rare earth excitation via post-transition halobismuthate(III) ions because of the large number of crystal motifs that halobismuthate(III) complexes are able to form. Complex halobismuthate(III) anions such as [Bi\textsubscript{2}Cl\textsubscript{10}]\textsuperscript{2−}, [Bi\textsubscript{4}Cl\textsubscript{18}]\textsuperscript{6−}, and {[Bi\textsubscript{2}Cl\textsubscript{8}]\textsuperscript{2−}}\textsubscript{∞} with various organic cations have been reported.\textsuperscript{16–26} These anions are composed of [BiCl\textsubscript{6}]\textsuperscript{3−} octahedra with various degrees of µ-Cl bridging. We have reported on simple alkali chlorobismuthate double salts.\textsuperscript{13} These double salts form highly networked structures of chloride-bridged metal cations whose structures are dependent on the choice of alkali metal cation. Based on these findings we have been interested in the capacity of halobismuthate(III) complexes to form network structures with photophysically active metal ions whereby energy transfer occurs from the [BiCl\textsubscript{6}]\textsuperscript{3−} anion to the neighboring ion. We believe this behavior is possible due to the extensive metal-metal bridging that occurs within the crystal lattice. We recently investigated possible energy transfer in mixed iodobismuthate(III)/iodocuprate(I) crystals coordinated to organic ligands which also contain halide bridge metal centers.\textsuperscript{16} Theoretical calculations indicated that these absorption bands were primarily the result of a halide/metal-to-metal charge transfer between the Bi(III) and Cu(I) centers. Based on these findings it seems possible to induce photoluminescence in photophysically active centers via energy transfer between halobismuthate(III) anions and lanthanide(III) cations.

Herein, we report the synthesis, structural analysis, and photophysical properties of the double salt TbBiCl\textsubscript{6}•14H\textsubscript{2}O, along with DFT and TD-DFT calculations to help interpret our findings. Solid state absorption measurements were carried out and showed significant overlap of the broad [BiCl\textsubscript{6}]\textsuperscript{3−} anion band with that of the sharp Tb(III) absorption bands. This is in agreement with luminescence measurements which reveal enhanced Tb(III) emission at 77 K. Experimental findings with theoretical support indicate that rare earth emission is the result of energy transfer from the Bi(III) metal center via the bridging chloride ions.
2. Experimental

2.1 Synthesis and Analysis of TbBiCl₆•14H₂O (1). All reagents were purchased from Aldrich or Acros and used without purification. Terbium(III) hexachlorobismuthate(III) tetrakaidecahydrate (1) was synthesized by combining BiOCl (5 mL 0.25 M in conc. aq. HCl) and aq. TbCl₃•6H₂O (5 mL 0.25 M). The filtered solution was slowly evaporated to yield pale yellow plates (1, 88.5%). Thermogravimetric analyses (TGA) (Figure S1 (SI)) were conducted using a TA Instruments Q500 in the dynamic (variable temp.) mode with a maximum heating rate of 50 °C/min. to 300 °C under 60 mL/min. N₂ flow. Compound 1 was analyzed for its Bi content via flame atomic absorption spectroscopy (AAS) using a PerkinElmer AAnalyst 700 instrument, confirming the formula derived from the X-ray crystallography. Samples were dissolved in 7% nitric acid (by weight) and then diluted in water to a concentration of 0.100 g/L. Blank solutions were made and their signal was subtracted from the sample scans. All scans were run three times with identical results. Experimental Bi content 26.2% (Theo. 25.1%).

2.2 X-ray Analysis. X-ray quality crystals were produced by partial evaporation of the aqueous solutions described above. The single crystal determination was carried out using a Bruker SMART Apex II diffractometer using graphite-monochromated Mo Kα radiation. Data for the complexes were collected at 100 K. The data were corrected for Lorentz and polarization effects and absorption using SADABS. The structure was solved by use of direct methods. Least squares refinement on F² was used for all reflections. Structure solution, refinement and the calculation of derived results were performed using the SHELXTL package of software. The non-hydrogen atoms were refined anisotropically. All hydrogen atoms were placed in theoretical positions.

2.3 Photophysical Measurements. Steady-state luminescence scans were run on 1 at 77 K. Compound 1 does not emit at temperatures above 90 K, so no room temperature scans were included. Spectra were taken with a Quantamaster-1046 photoluminescence spectrometer from Photon Technology International. This device uses a 75W xenon arc lamp combined with two excitation monochromators and one emission monochromator to adjust the bandwidth of light hitting the sample and detector, respectively. Signal intensity was measured using a photomultiplier tube. The samples were mounted on a copper plate using non-emitting copper-dust-high vacuum grease. Low-temperature scans were run on this system coupled to a Janis ST-
100 optical cryostat. Liquid nitrogen was used as coolant. Diffuse reflectance spectra (DRS) were collected on solid samples at 298 K. The light source was a Mikropack DH-2000 deuterium and halogen light source coupled with an Ocean Optics USB4000 detector. Scattered light was collected with a fiber optic cable. Spectra were referenced with PTFE. Data were processed using SpectraSuite 1.4.2_09.

2.4 Theoretical Calculations. DFT calculations were performed using the Gaussian ‘09 Software (Gaussian Inc.) hosted by the University of Maine Advanced Computing Group.† Ground state geometries and molecular orbital calculations were performed using the hybrid density functional theory due to Becke’s 3-parameter nonlocal exchange functional with the nonlocal correlation functional of Lee, Yang and Parr, B3LYP and the basis set SDD for all atoms. Molecular orbital calculations were performed on the ground state. The Avogadro software 1.2.0 was utilized for molecular orbital visualization. To best model the electronic properties of 1 we have performed calculations on a single cluster composed of a [BiCl$_6$]$^{3-}$ octahedron and [Tb(OH)$_2$)$_8$]$^{3+}$ square antiprism center as shown in Scheme 1 with the overall charge of the model being neutral. The X-ray structure coordinates were used as input for the initial geometry.

Scheme 1. A neutral model cluster of 1 composed of a [BiCl$_6$]$^{3-}$ octahedron and [Tb(OH)$_2$)$_8$]$^{3+}$ square antiprism center.

3. Results and Discussion

3.1 Structural Results. The crystal structure of 1 (Fig. 1) revealed isolated [BiCl$_6$]$^{3-}$ octahedra (Bi−Cl range = 2.571(5)-2.775(4) Å, cis-Cl−Bi−Cl range = 82.69(12)-98.9(2)°) and Tb(OH$_2$)$_8$ square antiprisms (Tb−O range = 2.318(13)-2.470(12) Å, cis-O−Tb−O angle range = 70.0(5)-81.3(3)°). These bismuth octahedra appear similar to those seen in previous reports of bismuth-chloride complexes. Three of the Cl and four of the bound H$_2$O ligands show site disorder. In addition, six non-coordinated water molecules were present. A series of O−H···Cl and O−H···O hydrogen bonds produces a 3-D network (Fig 2). This bridging of the bismuth-chloride center to
the terbium-H$_2$O center may allow for energy transfer from the bismuth cluster to the terbium ion. Additional crystallographic details are given in Tables S1-6. Thermogravimetric analysis of 1 revealed dehydration producing TbBiCl$_6$$\cdot$7H$_2$O and TbBiCl$_6$ phases (Fig. S1); TbBiCl$_6$$\cdot$7H$_2$O was produced on vacuum drying of 1. The expected Bi content for 1 was confirmed via AAS.

![Diagram](image1)

**Fig. 1** Thermal ellipsoid (50%) drawing of the crystallographically independent unit in 1 (secondary locations of disordered O and Cl atoms omitted for clarity).

![Diagram](image2)

**Fig. 2** Hydrogen bonding in 1 viewed along the $b$-axis (orange = Bi, green = Cl, turquoise = Tb, red = O, white = H).

**3.2 Diffuse Reflectance Spectroscopy.** Diffuse reflectance spectra of 1 were measured from crystals at room temperature (Fig. 3). A strong absorption band was observed at wavelengths less than ~380 nm with a weaker absorption noted between 380 nm and 530 nm. The sharp drop in absorption indicates an optical band edge at 388 nm. For comparison we measured the reflectance
of terbium(III) nitrate hexahydrate, which showed the characteristic rare earth emission bands at 542 nm and absorption bands from 325 nm to 390 nm. Interestingly, the reflectance of 1 does not display these $f-f$ absorption peaks and instead reveals only a single broad absorption band. We have assigned this broad band to the $[\text{BiCl}_6]^{3-}$ octahedral center as a Cl $3p \rightarrow \text{Bi} 6s$ transition. This assignment is in agreement with other studies of $[\text{BiCl}_6]^{3-}$ complexes which describe these bands as chloride to bismuth(III) ligand-to-metal charge transfer (LMCT) and bismuth(III) metal-centered transition (MC).\textsuperscript{39–41} The absorption band of the chlorobismuthate ion easily overlaps those of the terbium(III) nitrate, indicating possible energy transfer from the $[\text{BiCl}_6]^{3-}$ anion to the rare earth $[\text{Tb(OH}_2)_8]^{3+}$ cluster.

![Diffuse reflectance UV-vis spectra of 1 and terbium(III) nitrate hexahydrate.](image)

**Fig. 3** Diffuse reflectance UV-vis spectra of 1 and terbium(III) nitrate hexahydrate.

### 3.3 Photophysical Results

Double salt 1 was examined via steady state luminescence spectroscopy at 77 K. The compound did not emit at temperatures above 90 K. Fig. 4 shows the steady state luminescence spectra for 1. The annotations with arrows indicate the excitation/emission energy corresponding to the emission/excitation peaks, respectively. For Tb/Bi double salt 1 four emission peaks are observed. Each of these corresponds with a known emission of the Tb$^{3+}$ ion.\textsuperscript{1,2} However, a distinction exists between direct excitation of Tb$^{3+}$ and the energy transfer from Bi$^{3+}$ to Tb$^{3+}$. Under 351 nm excitation, Bi$^{3+}$ may become excited and transfer energy to Tb$^{3+}$. The Tb$^{3+}$ ion can also be excited directly at energies of 305, 358, 369 or 378 nm. The peak
centered at 584 nm is also partially the result of relaxation of the Bi\textsuperscript{3+} excited lattice via phonon-phonon interactions, in agreement with similar studies.\textsuperscript{42,43} The observed emission band is broad since bismuth has a high-energy excited state and low-energy excited states corresponding with the high and low energy emissions, respectively. The energy transfer pathway is favored at 352 nm because \textit{f} orbitals are difficult to excite directly, in contrast to \textit{d} orbitals.\textsuperscript{1,2,5,14} Octahedral bismuth-chloride centers undergo excitation of the chloride 2\textit{p} to the bismuth 6\textit{p}/chloride 3\textit{s}, which constitutes a HOMO-LUMO transition.\textsuperscript{36} This mechanism was first reported by Pelle \textit{et al.} in Cs\textsubscript{2}NaBiCl\textsubscript{6} crystal clusters.\textsuperscript{36-38} Addition of the terbium ions offers an alternative emission pathway. Due to the forbidden nature of \textit{f}-\textit{f} transitions, we expect the excitation of 1 at these wavelengths to occur via bismuth. Such transitions are also reported by Pelle \textit{et al.} who assigned the transition in the bismuth-chloride centers from the HOMO to the cation LUMO.\textsuperscript{36-38} The bridging of the chlorobismuthate octahedron to the [Tb(OH\textsubscript{2})\textsubscript{8}]\textsuperscript{3+} center in 1 provides an important interface which may facilitate energy transfer by shifting the emission of the Bi\textsuperscript{3+} slightly so as to increase the spectral overlap between bismuth and terbium. This increased overlap allows for greater Förster energy transfer.\textsuperscript{44} Table 1 shows state assignments for 1.

\begin{figure}
\centering
\includegraphics[width=0.8\textwidth]{Fig4}
\caption{Steady state luminescence spectra for 1 at 77 K. The annotations indicate the excitation/emission energy that corresponds with the selected emission/excitation peaks, respectively.}
\end{figure}
Table 1 Observed emission maxima for 1 at 77 K. Note: Tb^{3+} can also be directly excited at 305, 358, 369 and 378nm.

<table>
<thead>
<tr>
<th>$\lambda_{em}$ (nm), [$\lambda_{ex}$ (nm)]</th>
<th>Stokes Shifts (cm$^{-1}$)</th>
<th>Assignment</th>
</tr>
</thead>
<tbody>
<tr>
<td>490, [351]</td>
<td>8,080</td>
<td>$^5D_4 \rightarrow ^7F_6$ energy transfer</td>
</tr>
<tr>
<td>542, [351]</td>
<td>10,000</td>
<td>$^5D_4 \rightarrow ^7F_5$ energy transfer</td>
</tr>
<tr>
<td>584, [351]</td>
<td>11,400</td>
<td>$^5D_4 \rightarrow ^7F_4$ energy transfer</td>
</tr>
<tr>
<td>619, [351]</td>
<td>12,300</td>
<td>$^5D_4 \rightarrow ^7F_3$ energy transfer</td>
</tr>
</tbody>
</table>

3.4 DFT Calculations. DFT calculations have been performed on a single [Tb(OH$_2$)$_8$][BiCl$_6$] unit (Fig. 5) to explore the electronic transitions responsible for the observed luminescence behavior at different excitation energies. Ground state calculations produce a structure that is in general agreement with experimental X-ray studies (Bi–Cl range = 2.562-2.818 Å, cis-Cl–Bi–Cl range = 89.2-93.7°, Tb–O range = 2.398-2.541 Å, cis-O–Tb–O angle range = 63.8-79.9°). In this ground state both the octahedral [BiCl$_6$]$^{3-}$ and square antiprismatic [Tb(OH$_2$)$_8$]$^{3+}$ geometries are preserved. Also preserved in this ground state calculation are the close Cl–H distances (2.047-2.482 Å) which are important in our consideration of energy transfer from the chlorobismuthate anion to the terbium ion center.

Fig. 5 B3LYP/SDD calculated ground state of [Tb(OH$_2$)$_8$][BiCl$_6$]. Close Cl–H interactions shown in gray.
Fig. 6 Molecular orbitals diagram (left) and isodensity representations (right) of the frontier molecular orbitals in 1.

Isodensity representations of the frontier molecular orbitals in 1 are presented in Fig. 6. The higher-lying occupied molecular orbitals are primarily composed of the Cl 3p atomic orbitals, with significant contribution of the Bi 6s to the HOMO. A small contribution from the Tb 4f atomic orbital to the HOMO is also observed. The LUMO has significant contributions from the Bi 6s, Cl 3p, and Tb 4f. The remaining low-lying unoccupied molecular orbitals are composed of the Tb 4f atomic orbitals. In the luminescence spectra at low energies only rare earth f-f peaks are observed. In this case electrons most likely are promoted from the HOMO to unoccupied MOs other than the LUMO. In the case of high energy excitation, we observed both rare earth bands and a broad chlorobismuthate band which can only result from a Cl 3p → Bi 6s transition. According to the MO calculations, only the LUMO contains atomic orbitals for each of these transitions. Thus, at high energy excitation we expect that low-lying occupied MO electrons to be excited to the LUMO band. Since these low-lying MOs are composed only on the Cl 3p, this transition constitutes a Cl 3p → Tb$^{3+}$ energy transfer via a chlorobismuthate(III)/rare earth cluster-cluster interaction. With this in mind, we have developed an energy transfer mechanism which we believe describes the energy transfer processes at high and low energy excitation, shown in Fig. 7.
**Fig. 7** Energy diagram of 1 at high and low energy excitation. At high energy excitation (<350 nm) energy absorbed by the chlorobismuthate cluster is reemitted at 584 nm and transferred to the rare earth cluster competitively. At low energy excitation (>370 nm) energy absorbed by the chlorobismuthate cluster is transferred to the rare earth cluster and undergoes non-radiative relaxation. (ISC = Intersystem system crossing).

### 4.1 Conclusions

The novel bismuth(III)/terbium(III) double salt TbBiCl$_6$$\cdot$14H$_2$O (1) was synthesized and its crystal structure was determined. Compound 1 contains isolated [BiCl$_6$]$^{3-}$ and [Tb(OH)$_2$)$_8$]$^{3+}$ units built within a network structure. Structural measurements reveal close Cl$^{-}$H interactions between the bismuth and terbium ions, indicating the potential for energy transfer between centers. The broad absorption band observed in diffuse reflectance spectra are assigned to a Cl 3$p$$\rightarrow$Bi 6$s$ transition within the chlorobismuthate cluster. These transitions have been previously proposed by Pelle *et al.*$^{36-38}$ This [BiCl$_6$]$^{3-}$ absorption band strongly overlaps the sharp absorption bands of terbium(III). Luminescence experiments show enhanced rare earth ion emission from energy transfer between the excited [BiCl$_6$]$^{3-}$ anion and the [Tb(OH)$_2$)$_8$]$^{3+}$ cation. We have supported our findings with DFT calculations.

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Notes and references

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References

(1) Ahern, J. C.; Roberts, R. J.; Follansbee, P.; Lezno, D. B.; Patterson, H. H. Structure and Emissive Properties of Heterobimetallic Ln–Au Coordination Polymers: Role of Tb and Eu in Non-Aurophilic \([\text{nBu_4N}]_2[\text{Ln(NO_3)_4Au(CN)_2}]\) versus Aurophilic \(\text{Ln[Au(CN)_2]}_3\cdot3\text{H}_2\text{O/3D}_2\text{O Chains. Inorg. Chem. 2014, 53 (3), 7571–7579.}


(4) Wang, Y.; Yang, Z.; Ma, Y.; Chai, Z.; Qiu, J.; Song, Z. Upconversion Emission


(28) *SAINT Plus, Data Reduction Software*, version 7.34a; Bruker AXS Inc.: Madison, WI, 2005.

(29) Sheldrick, G.M. SADABS; University of Göttingen: Göttingen, Germany, 2005.


(44) Baril-Robert, F.; Guo, Z.; Patterson, H. H. Study of the Energy Transfer Process in the Highly Luminescent Heterometallic Dimers of Ce\(^{3+}\) and d\(^{10}\) [Ag(CN)]\(^{−}\) or d\(^{8}\) [Pt(CN)\(_4\)]\(^{2−}\) Ions. *Chem. Phys. Lett.* **2009**, *471*(4–6), 258–263 DOI: 10.1016/j.cplett.2009.02.039.