

Delft University of Technology

Exploring next-generation scintillation materials

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DOI 10.4233/uuid:d3f4e239-a40d-4857-8781-160851ea8b51

Publication date 2017 **Document Version**

Final published version

Citation (APA) Awater, R. (2017). *Exploring next-generation scintillation materials*. [Dissertation (TU Delft), Delft University of Technology]. https://doi.org/10.4233/uuid:d3f4e239-a40d-4857-8781-160851ea8b51

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Exploring next-generation scintillation materials

Proefschrift

ter verkrijging van de graad van doctor aan de Technische Universiteit Delft, op gezag van de Rector Magnificus prof. ir. K.C.A.M. Luyben, voorzitter van het College voor Promoties, in het openbaar te verdedigen op maandag 2 oktober 2017 om 15:00 uur

door

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Samenstelling promotiecommissie:

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Onafhankelijke leden: Prof. dr. L. D. A. Siebbeles, Prof. dr. ir. H. van der Graaf, Prof. dr. F. J. Beekman, Prof. dr. C. R. Ronda, Prof. dr. K. W. Krämer, Dr. V. Ouspenski,

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This research is supported by the Dutch Technology Foundation STW, which is part of the Netherlands Organisation for Scientific Research (NWO) and partly funded by the Ministry of Economic Affairs (project number 12799). This work was partly funded by Saint Gobain Research, France.

The research described in this thesis was carried out at the Luminescent Materials section, part of the Radiation Science & Technology department, faculty of Applied Sciences, Delft University of Technology.

Keywords:Scintillator, CeBr3, VRBE, Bi3+, Bi2+, Pb2+, Tl+Printed by:ProefschriftMaken.nl || Vianen

Cover design: Maikel Awater

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ISBN 978-94-6295-750-3

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I wish to dedicate this thesis to my parents and to the love of my life Elise ۱__

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List of abbreviations

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AE	Alkaline earth
APD	Avalanche photo diode
CB	Conduction band
CCD	Charge coupled device
CERN	European Organization for Nuclear Research
CMS	Compact Muon Solenoid
CS	Chemical shift
CT	Charge transfer
IVCT	Intervalence charge transfer
LHC	Large hadron collider
LY	Light yield
MMCT	Metal-to-metal charge transfer
MRI	Magnetic resonance imaging
NDE	Near defect exciton
NUV	Near ultraviolet
PET	Positron emission tomography
PL	Photoluminescence
PMT	Photo multiplier tube
R	Energy resolution
RE	Rare earth
RET	Resonant energy transfer
TL	Thermoluminescence
TM	Transition metal
UV	Ultraviolet
VB	Valence band
VIS	Visible
VRBE	Vacuum referred binding energy
VUV	Vacuum ultraviolet
XEL	X-ray excited luminescence
XRD	X-ray diffraction

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General introduction

1.1. Radiation detection

In order to protect ourselves from the harmful effects of ionizing radiation, it is essential that we can accurately measure the incoming ionizing radiation. For the detection of ionizing radiation a scintillator can be used. In short, a scintillator material converts the energy of the incoming radiation into multiple photons with lower energy, usually in the visible part of the spectrum. The emitted scintillation light, the amount of which is a direct measure of the energy of the ionizing radiation, is then detected using a photodetector. There is a wide variety of scintillation materials used for radiation detection, *e.g.* single crystals, plastics, glasses, liquids and gases. This thesis focuses exclusively on inorganic scintillation crystals for the detection of high-energy photons (X- and γ -rays).

Fig. 1.1a schematically illustrates the detection of ionizing radiation using a scintillator crystal coupled to a photomultiplier tube (PMT). γ -rays emitted by a radioactive source are absorbed by the scintillator material. As a result of photoelectric absorption or Compton scattering of the γ -radiation by the scintillation material, an electron is knocked out of its orbit around the nucleus and gains the absorbed energy in the form of kinetic energy. This ionization process is shown in more detail in Fig. 1.1b. The highenergy electron travels through the scintillator and forms an ionization track as it slows down while losing kinetic energy. The now mobile electrons and holes of the ionization track lose energy via thermalization until reaching the bottom of the conduction band (CB) and the top of the valence band (VB), respectively. The electrons and holes migrate to a luminescence recombination center, which is intentionally added to the scintillator in the form of a dopant/activator ion to enhance the probability of photon emission. An inorganic scintillation material is an insulator with typically a band gap of 4–12 eV. The ground state (GS) and excited state (ES) of the dopant ion are located inside the band gap of the host compound. This allows the electron and hole to recombine on the dopant ion, releasing the energy in the form of a visible photon. The wavelength of the emitted photon strongly depends on the dopant ion and its chemical environment provided by the host lattice. Due to the high-energy of the incoming γ -ray (keV–GeV), the resulting scintillation pulse may contain several hundreds of thousands of visible photons.

The photons emitted by the scintillator are directed to a photodetector, which is most commonly a PMT. Inside the PMT, the visible photons are absorbed by the photocathode material and, due to the low work function of the cathode material, photoelectrons are released. The liberated photoelectrons accelerate due to an applied electric field and are directed by the focusing electrode to the first dynode. The photoelectrons, which have gained considerable kinetic energy while accelerating, impact the first dynode. Upon impact of the photoelectrons, two or more electrons are ejected from the dynode per incoming photoelectron which are accelerated towards the second dynode. The number of electrons are multiplied again at the second dynode. By using multiple dynode stages, the original amount of photoelectrons can be multiplied more than a million times. Eventually, the electrons reach the anode where the electrical signal is readout and this signal can be analyzed. More information on radiation detection and measurement can be found in the literature [1].



Figure 1.1: Schematic illustration of a) a scintillator detector and b) the scintillation process.

1.2. Brief history of scintillation materials research

In 1895, Röntgen observed luminescence from a $BaPt(CN)_4$ phosphor placed next to a discharge tube. He recognized that the luminescence was caused by a new type of radiation which he named X-rays [2]. Since these phosphors were inefficient at converting X-rays into visible light, the search for better scintillators started. Already in 1896, CaWO₄ powder was discovered to emit blue light much more efficiently when exposed to X-rays. Together with ZnS-based powders discovered a few years later, these scintillator compounds were used in X-ray photography for more than 75 years [3]. ZnS was also used in visual scintillation counting until the introduction of the PMT in the 1940s.

In the late 1940s, Hofstadter discovered the single-crystal scintillator NaI:Tl which is still the most commonly used scintillator for γ -ray spectroscopy [4]. This was also the start of a new boost in scintillation materials research, resulting in the discovery of many new scintillators, *e.g.* CdWO₄, CsI:Tl and glass scintillators activated with cerium. Steadily over the next decades, new scintillator materials were discovered of which the extraordinary fast BaF₂ and very dense Bi₄Ge₃O₁₂ are most notable. In the 1990s, driven by the need for denser scintillators for high-energy physics and higher light yield scintillators for medical imaging, a new uplift in scintillation material research occurred. This led to the development of the high density and very fast PbWO₄ scintillation material for the CMS detector in the LHC at CERN. For medical PET scanners, Lu₂SiO₅:Ce³⁺ was found to excel Bi₄Ge₃O₁₂, the most commonly used material at that time, in terms of higher light yield and faster decay time.

Since 2000, scintillation materials research focused mainly on Ce³⁺ and Eu²⁺ activated halide scintillators. LaBr₃:Ce³⁺ and SrI₂:Eu²⁺ are two examples of materials that were discovered with excellent scintillation properties, such as a record high light yield of up to 120000 photons emitted per MeV of absorbed ionizing radiation energy and excellent energy resolution of down to 3.0% at 662 keV γ -excitation. Later, the energy resolution of LaBr₃:Ce³⁺ crystals was even further improved to a value of only 2.0% by co-doping the crystal with a small amount of Sr²⁺ [5].

The extensive research on scintillation materials resulted in a large variety of commercially available scintillators, each with their own characteristic material properties tailored to a specific application. Table 1.1 lists some of the scintillation properties of the most commonly applied scintillators that are discovered so far. Table 1.1 is not extensive and a more complete list of the scintillation properties of inorganic scintillation materials is available on the internet [6]. The search for new, better performing scintillation materials is still a very active field of research. The ever-increasing material demands set by various applications drive current research and development efforts in finding the next-generation of scintillation materials.

	Light yield	R (% at	Density	Decay time	λ_{\max}	
Scintillator	(ph/MeV)	662 keV)	(g/cm ³)	(ns)	(nm)	Ref.
LaBr ₃ :Ce,Sr	78000	2.0	5.1	18 + slow	370	[5]
LaBr ₃ :Ce	64000	3.1	5.1	16	360	[7]
CeBr ₃	60000	4.1	5.2	17	370	[8]
SrI ₂ :Eu	120000	3	4.6	1200	435	[9]
Lu ₂ SiO ₅ :Ce	27000	7.9	7.4	40	420	[10]
Cs ₂ LiLaBr ₆ :Ce	45000	3.1	4.2	180 + 1140	420	[11]
NaI:Tl	40000	6.7	3.7	230	415	[12, 13]
CsI:Tl	65000	6.6	4.5	980	530	[12, 14]
Bi ₄ Ge ₃ O ₁₂	9000	9	7.1	300	480	[13, 15]
PbWO ₄	300	30 - 40	8.3	2 - 3	410	[3]
CdWO ₄	16000	8.3	7.9	13000	475	[16]
$CaWO_4$	15000	7.8	6.1	8000	425	[17]
BaF ₂ (fast)	2000	10	4.9	0.8	220	[18]

Table 1.1: Physical properties of common inorganic scintillation materials for γ -ray detection.

1.3. Applications of scintillation materials

Each application in which scintillators are used has its own set of specific material requirements. Some of the most important requirements for scintillation materials used for γ -ray detection are (not necessarily in this order):

- 1. High light yield
- 2. Excellent energy resolution
- 3. Proportional scintillation response
- 4. Low cost
- 5. High density
- 6. High effective atomic number
- 7. Fast decay time
- 8. Short rise time
- 9. No self-absorption
- 10. No afterglow
- 11. Emission wavelength matches photodetector sensitivity
- 12. Refractive index matches photodetector window material
- 13. No internal radioactivity
- 14. High temperature stability
- 15. Excellent radiation hardness
- 16. High mechanical strength
- 17. Able to grow large crystals
- 18. Non hygroscopic

The following paragraphs provide a brief overview of the material requirements for each application.

Medical diagnostics Scintillation materials are used in several medical diagnostics and imaging techniques such as X-ray computed tomography, single-photon emission computed tomography (SPECT) and positron emission tomography (PET). In order to expose a patient to the lowest amount of radiation as possible, a high detection efficiency is crucial. This requires scintillation materials with a high light yield, a high density and high effective atomic number. A short rise time and fast scintillation decay time are essential for time-of-flight PET measurements and for short acquisition times. For X-ray tomography, Gd₂O₂S:Pr,Ce,F and (Y,Gd)₂O₃:Eu,Pr ceramic scintillators are the most widely used. NaI:Tl and CsI:Tl are currently favored as scintillator in SPECT scanners. Due to its high density and reasonably fast decay time, Bi4Ge3O12 used to be the material of choice in PET scanners. However, Lu₂SiO₅:Ce offers an even faster decay time and much higher light yield and is therefore replacing Bi₄Ge₃O₁₂ in these scanners. Drawbacks of Lu₂SiO₅:Ce are the difficulties in growing large, good quality crystals and the high price of lutetium. LaBr₃:Ce has been considered to replace the current scintillators for SPECT and PET, however, high production costs and too low density are limiting its commercialization for these applications. A more detailed discussion of scintillator materials for medical diagnostics can be found in Ref. [19].

High-energy physics High-energy physics experiments, such as the particle accelerators at CERN, require very high density scintillators in order to efficiently stop and detect the extremely high-energy (GeV - TeV) radiation. It also requires the scintillation material to have an excellent radiation hardness. For the L3 detector in the Large Electron-Positron Collider at CERN, nearly 12000 $Bi_4Ge_3O_{12}$ crystals of 24 cm were used [20]. While for the CMS detector in the LHC, more than 75000 PbWO₄ crystals of 23 cm were used [21]. Both scintillation materials have high densities and have a high effective atomic number which makes them ideal candidates for use in high-energy physics experiments. The large quantities of material used in these detectors require that the scintillator can be grown in large volumes at low cost. On the other hand, the amount of energy deposited in the scintillation material is so large that a relatively low light yield of for example only 300 ph/MeV for PbWO₄ is already acceptable. Furthermore, a fast scintillation decay time is required that is at least faster than the collision rate in the particle accelerator. More information on scintillation materials and their requirements for high-energy physics can be found in Refs. [22] and [23].

Astrophysics and space exploration Scintillation materials are used in astrophysics experiments, for example to detect the characteristic X-rays emitted by newly formed stars or to study the radiation coming from supernova and other astrophysical events. γ -ray spectrometers containing scintillation materials are also put on-board spacecraft for planetary remote sensing. This provides valuable information regarding the planet's elemental surface composition. To accurately monitor and differentiate between the various emitted energies coming from the surface, a high energy resolution is necessary. Furthermore, the amount of radiation reaching the spacecraft is very low, which requires a high sensitivity. Essential for these low count-rate applications is to have an as low as possible background noise, which requires the scintillator to have sufficiently low internal radioactivity. Previously, only high-purity germanium detectors met the requirements to be used as γ -ray spectrometers for space applications. Since 2000, LaBr₃:Ce scintillators provide excellent energy resolution, high sensitivity and room temperature operability that can compete with the semiconductor detectors [24]. In the last decade, LaBr₃:Ce scintillator detectors were developed for the European Space Agency's Bepi-Colombo mission to Mercury [25]. Unfortunately, LaBr₃:Ce contains a small amount of the naturally occurring radioactive ¹³⁸La isotope, which spoils its capability in low noise applications. Despite a slightly worse energy resolution, CeBr₃ emerged as a promising alternative to LaBr₃:Ce as it has an almost 30× lower internal activity.

Security and environmental monitoring Machines for luggage screening at airports and container scanning at naval ports contain scintillation materials. Most of the modern scanners operate in the same way as X-ray computed tomography scanners for medical diagnostics to image the contents inside a bag or container. A high light yield and fast decay time are the most important requirements for fast and accurate image acquisition. Also the protection from terrorism and monitoring of radiation from environmental disasters (like Chernobyl and Fukushima) requires accurate detection of dangerous neutron and/or gamma-emitting materials. In both cases, the amount of radiation is low which requires high light yield scintillators for a high sensitivity. A high energy resolution and good neutron/gamma discrimination are required for accurate isotope identification. Furthermore, low cost scintillation materials are needed for mass production of these radiation detectors. Scintillation materials that meet these requirements and are currently under development are Cs₂LiYCl₆:Ce, Cs₂LiLaBr₆:Ce and SrI₂:Eu.

Geophysical and well logging Gamma-ray detectors are used to remotely measure the natural radioactivity of the different soil layers of a drilled well or borehole. When the scintillator detector is lowered into a drilled well, it records the geophysical data as function of the well depth in a gamma-ray log. These gamma-ray logs are used in the search for oil and mineral sources but also to gather information on the age and formation of rock and soil layers. With temperatures over 200 °C and shock and vibrations encountered during operation, the scintillation materials used for well logging should have a high temperature stability and should be mechanically rugged. Furthermore, these scintillation materials should also have a high density, high effective atomic number, fast decay and should be non-hygroscopic to avoid the need of special packaging [26]. So far no scintillation material exists which meets all of these requirements. For most well logging applications NaI:Tl is used, however, $Bi_4Ge_3O_{12}$ and BaF_2 are in some cases favored.

1.4. Understanding the performance of scintillators

Essential in finding the next-generation of scintillation materials is to understand and predict their scintillation properties. The electronic structure of a host compound, together with the location of the impurity states of a dopant ion within the band gap, provide crucial insight in the performance of scintillation materials [27]. Based on lanthanide spectroscopy, Dorenbos developed the redshift, centroid shift and charge transfer models which can be used to locate the lanthanide 4f and 5d electron binding energies relative to the valence band [28]. This was followed by the chemical shift model, which relates all the lanthanide levels and host bands to the vacuum energy [29, 30]. The vacuum energy is defined as the energy of an electron at rest in the vacuum and is set as energy zero. The obtained absolute electron binding energies are conveniently visualized in a so-called Vacuum Referred Binding Energy (VRBE) diagram. Fig. 1.2 shows as an example the VRBE diagram of LaBr₃, including the 4f and 5d electron binding energies of the di- and trivalent lanthanide ions. The lowest 4f and 5d states of the di- and trivalent lanthanide ions. The lowest 5d state for Tb³⁺ until Lu³⁺ is split into a high spin (HS) and low spin state (LS).

To achieve efficient 4f-5d luminescence, the CB bottom should be at least higher in energy than the VRBE in the lowest 5d state. Another requirement is that the emission energy is smaller than the band gap energy. This information can be derived directly from a VRBE diagram. Furthermore, it was found that the lanthanide impurity levels change rather predictably when changing the chemical environment [31]. A small change in the location of the impurity states as compared to the host states can have a major effect on the scintillation performance. Such knowledge allows deliberate engineering of the electronic structure of scintillation materials by changing the chemical composition of the host compound or using different host-dopant combinations [32].



Figure 1.2: The vacuum referred binding energies of the divalent (red) and trivalent (blue) lanthanide 4f and 5d electrons in $LaBr_{q}$. E_X is the binding energy in the host exciton.

1.5. Research objective and thesis outline

With all the stable elements of the periodic table at our disposal to synthesize host compounds, there are an uncountable number of scintillator host lattices available. Combined with a suitable dopant ion, which is not limited to the lanthanides but also comprise the transition metal and post-transition metal ions, this provides an almost infinite research field on scintillation materials development. The general aim of the research described in this thesis is to explore the next-generation of scintillation materials. Supported by the VRBE diagrams described in section 1.4, new avenues of scintillation materials research and development are investigated.

Currently, most research is focused on halide crystals with lanthanide dopants and Ce^{3+} in particular. However, extensive research efforts on these type of materials have already been performed over the last decades. These Ce^{3+} -doped halide materials have almost reached their theoretical maximum light yield, which allows only little room for finding new, better performing Ce^{3+} -doped halides. An alternative to finding new materials is to improve the scintillation properties of existing scintillation materials. Chapter 2 discusses such an approach by means of aliovalent doping of $CeBr_3$. The effects of Na⁺, Mg²⁺, Ca²⁺, Sr²⁺ and Ba²⁺ doping on the scintillation properties of CeBr₃ are investigated.

Another direction in the field of scintillation materials is the search for longer wavelength (red-to-infrared) emitting scintillators. The red-to-infrared emission wavelengths match with the sensitivity of modern photodiodes. Unlike PMTs, scintillator photodiode detectors are small, operate at low voltage, are rugged and are insensitive to magnetic fields, which makes them ideally suited as radiation detectors on-board spacecraft and in PET scanners combined MRI. Furthermore, by engineering red-emitting scintillation materials with a smaller bandgap it is possible to go to even higher light yields. Chapters 3 and 4 are focused on Bi^{2+} , a relatively new dopant ion in the field of scintillation materials with red-to-infrared emission wavelengths. Chapter 3 discusses the X-ray induced valence change of Bi^{3+} to Bi^{2+} and the luminescence and scintillation properties of bismuth doped $Li_2BaP_2O_7$. Chapter 4 deals with the luminescence and charge carrier trapping in YPO₄:Bi.

New scintillation materials with higher densities and higher effective atomic number are of particular interest for high-energy physics experiments. However, with halide crystals it is not possible to achieve densities higher than 6 g/cm³. With scintillation materials based on oxides it is possible to go to densities beyond 8 g/cm³. Even more attractive would be oxides containing high effective atomic number elements such as Tl, Pb or Bi. In order to evaluate if Ce^{3+} can luminescence efficiently in these type of host compounds, knowledge on the impurity levels of these post-transition metal ions relative to the host conduction and valence bands is required. This is the topic of chapters 5, 6 and 7. Chapter 5 deals with the electron binding energies in the impurity levels of Bi³⁺ and how these change in relation to the chemical environment. Based on the electron binding energies in the impurity levels in Bi³⁺ of chapter 5, together with electron binding energies in the impurity levels in Bi²⁺, a concentration quenching model of the Bi³⁺ luminescence via election transfer between bismuth pairs is introduced in chapter 6. Chapter 7 reviews the electron binding energies in the impurity levels of Pb²⁺ and Tl⁺ in different inorganic compounds and compares the obtained results with those of Bi³⁺ from chapter 5.

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2

Effects of Na⁺, Mg²⁺, Ca²⁺, Sr²⁺ and Ba²⁺ doping on the scintillation properties of CeBr₃

The effects of Na⁺, Mg²⁺, Ca²⁺, Sr²⁺ and Ba²⁺ doping on the scintillation properties of CeBr₃ are evaluated in this chapter. Ca²⁺ and Sr²⁺ doped CeBr₃ show different scintillation properties as compared to undoped CeBr₃, i.e. an additional perturbed Ce³⁺ site with emission maxima red-shifted by 30 nm compared to unperturbed Ce³⁺ sites as evidenced by X-ray excited emission and photoluminescence measurements. Also a nearly constant light yield at temperatures below 300 K, an increased decay time and the appearance of multiple glow peaks in thermoluminescence measurements are observed for Ca and Sr doped CeBr₃. These effects on the scintillation properties of CeBr₃ are explained using a single trap model where Br vacancies, formed as charge compensation for the aliovalent dopants, act as electron traps.

This chapter is based on the publication: **R. H. P. Awater**, K. W. Krämer and P. Dorenbos, "Effects of Na⁺, Mg²⁺, Ca²⁺, Sr²⁺ and Ba²⁺ doping on the scintillation properties of CeBr₃", *IEEE Transactions on Nuclear Science* **62** (2015), 2343.

2.1. Introduction

Since the invention of the photomultiplier tube and the discovery of NaI:Tl⁺ crystal scintillators in the 1940's, many new scintillator materials have been discovered and investigated for the detection of gamma radiation. However, even at present time NaI:Tl⁺ is still the most commonly used scintillator worldwide. With the discovery of LaBr₃:Ce in 2001, a new material combining a high light yield of 70,000 photons per MeV of absorbed gamma ray energy, excellent energy resolution of 2.8% for 662 keV gamma ray detection and a short scintillation pulse decay time of 15 ns was obtained [1].

A major drawback of LaBr₃:Ce is that it contains ¹³⁸La isotopes that causes an intrinsic activity which spoils the capability of LaBr₃:Ce in low count rate experiments [2]. The drastically reduced intrinsic activity due to the absence of naturally occurring isotopes makes CeBr₃ a more interesting scintillator for the detection of low intensity gamma ray sources compared to LaBr₃:Ce. Below 3 MeV, the detection sensitivity of CeBr₃ is $5 \times$ higher than for LaBr₃:Ce and up to $16 \times$ higher for detection of ⁴⁰K [3, 4]. Because of the increased detection sensitivity, CeBr₃ offers superior performance for applications such as planetary remote sensing, homeland security and radionuclide identification [5–7]. Unfortunately CeBr₃ shows a poorer energy resolution of 4% at 662 keV compared to 3% for LaBr₃:5%Ce [3].

Because of the hexagonal crystal structure (space group P63/m) of CeBr3 and lanthanide halides in general, the crystals have anisotropic thermal expansion and plasticity resulting in build-up of stresses in the material during growth [8]. Together with the tendency to fracture along the prismatic slip planes, these materials are very brittle. Attempts were made to improve the crystal strength by aliovalent doping without negatively influencing the scintillation properties. Interestingly, the use of alkaline earth metals as dopant was shown to improve the energy resolution of LaBr₃:Ce and CeBr₃ considerably for small crystal samples. Sr²⁺ as a dopant improved the energy resolution to a value of 2% in LaBr₃:5%Ce [9], while Ca^{2+} as a dopant improved the energy resolution of CeBr₃ to 3.2% [10]. This improvement in energy resolution of the doped samples was ascribed to a more proportional response of the light yield compared to standard LaBr3:5%Ce or CeBr3. The exact origin of this improvement due to aliovalent doping remains controversial. A possible explanation by Åberg et al. is that the aliovalent dopants create Br vacancies acting as electron traps, which reduces Auger quenching of free carriers and that is expected to improve the linearity of the light yield as function of the energy of the incident radiation [11].

In a previous investigation, the energy resolution enhancement of Na^+ , Mg^{2+} , Ca^{2+} , Sr^{2+} and Ba^{2+} doped CeBr₃ samples was evaluated and an energy resolution of down to 3.0% for the Sr²⁺ doped sample was found [12]. In this chapter we report on the effects of these same dopants on the scintillation properties of CeBr₃.

2.2. Experimental methods

The undoped and Na⁺, Mg²⁺, Ca²⁺, Sr²⁺ and Ba²⁺ doped CeBr₃ crystals discussed in this chapter were grown at the University of Bern by the vertical Bridgman technique in sealed quartz ampoules and contained a nominal dopant concentration of 0.5%. It is expected that at maximum only a few hundred ppm of dopant is actually incorporated

in the crystal, since inductively coupled plasma analysis only showed 100-200 ppm of Sr^{2+} in Sr^{2+} co-doped LaBr₃:Ce [13]. The crystals were grown as ingots of 8 mm in diameter of which $30 - 50 \text{ mm}^3$ samples were cleaved. In a previous paper the effective enhancement of the energy resolution via aliovalent doping of the same CeBr₃ samples was discussed [12]. For all temperature dependent measurements a Janis VPF-800 cryostat operated with a LakeShore 331 temperature controller were used. The cryostat was baked to remove all water from the system prior to sample mounting because of the hygroscopic nature of the CeBr₃ crystals. The samples were mounted on the cold finger of the cryostat in a nitrogen filled glovebox. The PMT's were outside the cryostat and remained always at room temperature. X-ray excited emission and thermoluminescence spectra were recorded using an X-ray tube with Cu anode operating at 60 kV and 20 mA. The emission of the sample was focused via a quartz window and a lens on the entrance slit of an ARC VM504 monochromator (blazed at 300 nm, 1200 grooves/mm), dispersed, and recorded with a Hamamatsu R943–02 PMT. The spectra were corrected for the monochromator transmission and the quantum efficiency of the PMT. For thermoluminescence measurements the crystals were cooled down to 78 K and then irradiated with X-rays for approximately 5 min leading to a steady state X-ray excited luminescence (SSL). The thermoluminescence emission was monitored at the respective emission maximum of the sample while heating with a constant rate of 6 K/min using a Lakeshore 331 temperature controller. Photoluminescence excitation and emission spectra were recorded using an OSRAM XBO450W/4 Suprasil Xenon lamp in combination with a Horiba Gemini - 180 double grating monochromator. The emission light of the crystal was dispersed with a Princeton Instruments Acton SP 2300 monochromator and detected by a Perkin Elmer Photon Counting Module MP1993. The excitation spectra were corrected for the lamp spectrum while no further correction was applied for the emission spectra. Temperature dependent photoelectron yield measurements were derived from pulse height spectra which were recorded using a Hamamatsu R6231-100 PMT at a voltage of -500 V, a 137 Cs source (662 keV) and a 10 μ s shaping time. The time correlated single photon counting method using a pulsed picosecond X-ray source was used to record the scintillation time profiles. The X-ray tube with a S-20 type photocathode and tungsten anode was operated at a voltage of 40 kV (Emax = 40 keV). A HP 8116A Pulse/Function generator in combination with a Picoquant Sepia multi-channel laser diode were used to trigger the X-ray pulses. An IDQ id100 single photon detection module was used as detector. In short, photoelectrons are liberated by the emitted photons from the laser diode and are accelerated towards the anode, resulting in pulsed X-rays which are directed towards the sample. More detailed information on the pulsed X-ray setup can be found in [14].

2.3. Results and discussion

Fig. 2.1 shows the X-ray excited emission spectra of the studied CeBr₃ crystals at 298 K and 78 K. At room temperature, the undoped, Na⁺ and Mg²⁺ doped CeBr₃ samples show two overlapping peaks at 367 nm and 388 nm. In the case of Ba²⁺ doped CeBr₃ the two emission peaks are hardly distinguishable. The emission bands of Sr²⁺ and Ca²⁺ doped CeBr₃ samples are slightly red-shifted and show a maximum emission at 381 nm with a side band at longer wavelengths. At 78 K the undoped, Na⁺, Mg²⁺ and Ba²⁺ doped CeBr₃



Figure 2.1: X-ray excited emission spectra of undoped and Na, Mg, Ca, Sr and Ba doped CeBr₃ crystals recorded at (a) 298 K and (b) 78 K.

samples show the typical double-band Ce^{3+} 5d-4f emission according to the transitions into the ${}^{2}F_{5/2}$ and ${}^{2}F_{7/2}$ states centered at 361 nm and 390 nm, respectively. The 390 nm peak has approximately 60% of the intensity of the 361 nm peak except for Ba^{2+} doped $CeBr_{3}$ were the intensity of the 390 nm peak is approximately 80%. These double-band emissions are red-shifted by 5 nm when compared to $LaBr_{3}$:Ce with maxima at 355 nm and 385 nm [13]. The Sr^{2+} and Ca^{2+} doped $CeBr_{3}$ samples show the largest effect on the radio luminescence spectrum, i. e. a maximum emission at 386 nm with a shoulder at 367 nm and a second but 70% less intense peak at 412 nm. This is in contrast to Harrison *et al.* who reported the largest effect on the radio luminescence spectrum for Ba doped $CeBr_{3}$ [15].

Optically excited emission spectra of undoped and Ca^{2+} and Sr^{2+} doped $CeBr_3$ samples at 78 K are shown in Fig. 2.2. At 285 nm excitation, the undoped $CeBr_3$ sample shows a typical Ce^{3+} doublet at 362 nm and 392 nm. The Sr^{2+} doped $CeBr_3$ sample shows three poorly resolved bands at 366 nm, 388 nm and 412 nm, similar to the X-ray excited emission in Fig. 2.1. The Ca^{2+} doped $CeBr_3$ sample shows also three overlapping bands, but the 422 nm band is now almost as intense as the 392 nm band. The additional emission band around 420 nm in Sr^{2+} and Ca^{2+} doped $CeBr_3$ is believed to be from the doublet



Figure 2.2: Photoluminescence spectra of undoped and Ca and Sr doped CeBr₃ crystals recorded at 78 K and excited at 285 nm.

emission of perturbed Ce³⁺ sites. The presence of Sr²⁺ or Ca²⁺ dopant ions is believed to create Br vacancies as charge compensation, which change the local environment of the Ce³⁺ ion, leading to perturbed Ce³⁺ ions and causes a redshift in the emission compared to unperturbed Ce³⁺ emission. The three overlapping bands in the X-ray and optical excited emission spectra of Ca^{2+} and Sr^{2+} doped CeBr₃ therefore originate from two overlapping Ce³⁺ emission doublets: the unperturbed Ce³⁺ emission doublet centered at around 362 nm and 392 nm and the perturbed Ce³⁺ emission doublet centered at around 392 nm and 422 nm, labeled Ce_I and Ce_{II}, respectively. Looking at the excitation spectra of Ca^{2+} doped CeBr₃ in Fig. 2.3 at the three different emission wavelengths, it appears there are two excitation bands also indicating the presence of two different Ce^{3+} emission sites. The first excitation band is observed between 230 and 350 nm and is attributed to excitation of standard, unperturbed Ce³⁺ (Ce_I). Because of the high concentration of Ce^{3+} the five 4f $[{}^{2}F_{5/2}] \rightarrow 5d$ excitations are not resolved and therefore not visible in the excitation spectrum. The second excitation band is centered at 350 nm and is attributed to the excitation of perturbed Ce^{3+} ions (Ce_{II}). The excitation band of Ce_{II} overlaps with the emission of Ce_I, resulting in energy transfer from Ce_I to Ce_{II}, leading to an effective increase in the emission intensity of Ce_{II} at the cost of the emission of Ce_I. These perturbed Ce³⁺ sites are also reported for co-doped LaBr₃:Ce in [16], where Br⁻ vacancies and interstitial site occupation are suggested as possible charge compensation mechanisms for divalent co-doping.

Fig. 2.4 shows the relative photoelectron yield as function of temperature for the studied CeBr₃ crystals normalized to 100% at 300 K. The temperature dependencies of Na⁺ and Mg²⁺ doped CeBr₃ are similar to undoped CeBr₃ and are comparable with data reported for co-doped LaBr₃:Ce with the same dopants, i.e. a continuous decrease in light yield (LY) from 125% at 80 K to 70%-85% at 600 K depending on the type of dopant [13]. The temperature dependencies of Ca²⁺, Sr²⁺, and Ba²⁺ doped CeBr₃ crys-



Figure 2.3: Photoluminescence excitation spectra of a CeBr₃:Ca crystal recorded at 78 K while monitoring the (a) 370 nm, (b) 392 nm and (c) 422 nm emission.

tals were found to behave differently also with respect to LaBr₃:Ce with the same dopants. Ba²⁺ doped CeBr₃ shows a continuous decrease in LY from 115% at 80 K to 75% at 600 K, comparable to the linear decrease for undoped, Na⁺ and Mg²⁺ doped CeBr₃ samples. The light yield of the Ca²⁺ and Sr²⁺ doped CeBr₃ samples has a maximum at 250 K and decreases only by at most 5% as the temperature is further decreased. This is different from LaBr₃:Ce with the same dopants, where the LY is maximum at room temperature and decreases to 50-80% at 85 K. A strong decrease in LY to 55% and 75% for Ca²⁺ and Sr²⁺ doped CeBr₃ is observed as the crystals are heated to 600 K, which is comparable to the behavior observed for LaBr₃:Ce with the same dopants.

The thermoluminescence glow curves of the studied CeBr₃ crystals are shown in Fig. 2.5. Na⁺ and Mg²⁺ doped CeBr₃ show a maximum thermoluminescence emission of 0.2% of their steady-state X-ray excited luminescence intensity (SSL), which is similar to undoped CeBr₃. The maximum thermoluminescence emission for Ca²⁺, Sr²⁺ and Ba²⁺ doped CeBr₃ crystals are 60%, 70% and 40% of their SSL, respectively, two orders of magnitude higher than that of Na⁺, Mg²⁺ and undoped CeBr₃ and approximately 2× higher



Figure 2.4: Temperature dependence of the relative photoelectron yield of undoped and Na, Mg, Ca, Sr and Ba doped CeBr₃ crystals. The spectra are normalized to 100% at 300 K. The inset shows the thermoluminescence glow curve of Ca, Sr and Ba doped CeBr₃ samples on the same temperature scale.

compared to LaBr₃:Ce with the same dopants [13]. The Ca²⁺, Sr²⁺ and Ba²⁺ doped CeBr₃. crystals show multiple thermoluminescence glow peaks between 100 and 200 K as can seen from the inset of Fig. 2.4, which indicate the thermal release of charge carrier traps. As a charge compensation mechanism for the introduction of the divalent Ca^{2+} , Sr^{2+} and Ba²⁺ ions in the CeBr₃ lattice, it is believed that Br vacancies are formed which can act as electron traps. To explain the observed more or less constant behavior of the light yield between 80 K and 250 K for the Ca^{2+} and Sr^{2+} (and to a lesser extent Ba^{2+}) doped CeBr₃, a single trap model is employed (Fig. 2.6) which was shown to be successful in explaining electron traps in $Lu_3Al_5O_{12}$: Pr [17]. Prompt consecutive capture of charge carriers at Ce^{3+} , followed by radiative recombination (process 1 in Fig. 2.6) is the main route for emission in the CeBr₃ samples. A second route of emission (process 2) is via electron capture at the trap, followed by a delayed thermal release of the electron (process 3) which migrates and recombines radiatively with a hole at a Ce luminescent center. At temperatures below 200 K the trap lifetime is longer than the shaping time in the pulse height measurements and therefore emission via trapped electrons does not contribute to the luminescence, resulting in a decrease in the total light yield between 80 K



Figure 2.5: Thermoluminescence glow curves of undoped and Na, Mg, Ca, Sr and Ba doped CeBr₃ samples. The curves are normalized to the steady-state X-ray excited luminescence intensities (SSL)



Figure 2.6: Schematic representation of the single trap model in $CeBr_3$ with dopants. 1: migration of electron via conduction band to Ce^{3+} , 2: capture of electron by electron trap, 3: thermal release of electron followed by migration via conduction band to Ce^{3+} .

and 250 K as observed in Fig. 2.4. Above 200 K the trap lifetime becomes sufficiently short that process 3 now contributes to the measured signal, increasing the total light yield. Above 250 K the light yield starts to decrease again due to thermal quenching and increased self absorption. These effects are less pronounced in Ba^{2+} doped CeBr₃, probably because a smaller amount of electron traps are present. Unfortunately, with the used measurement setup it is not possible to perform reliable quantitative thermoluminescence analysis such as trap depth and lifetime determination.



Figure 2.7: Scintillation decay profiles of undoped and Na, Mg, Ca, Sr and Ba doped CeBr₃ crystals recorded at 300 K. The dashed lines show the background levels of (top line) undoped, Ca, Sr and Ba doped CeBr₃ and (bottom line) Na and Mg doped CeBr₃.

The scintillation decay profiles of the studied CeBr₃ samples are shown in Fig. 2.7. The scintillation decay profiles are fit with either one or two exponential functions depending on the dopant and the calculated decay constants are summarized in Table 2.1. Na⁺ doped CeBr₃ shows only a fast single exponential decay of around 17 ns like undoped CeBr₃. Mg²⁺ doped CeBr₃ shows a slightly increased decay time of 19ns. Ca²⁺, Sr²⁺ and Ba²⁺ doped CeBr₃ show besides an increased decay time of 19-24 ns, also a slow component of 86-223 ns which contributes only a few percent to the total scintillation output. The fast components of the decay time constants are plotted as function of temperature in Fig. 2.8. Two main features are apparent from the obtained data: 1) the decay time constant increases when dopants are added; and 2) the decay time constant increased decay time for Ca²⁺, Sr²⁺ and Ba²⁺ doped CeBr₃ samples can be partly ascribed to the red-shifted emission of Ce_{II}. Eq. 2.1 relates the decay rate of the excited state and the emission wavelength [19]:

$$\Gamma = \frac{1}{\tau} \sim \frac{n(n^2 + 2)^2}{\lambda^3} |\langle f | \mu | i \rangle|^2$$
(2.1)

where τ is the radiative lifetime of the Ce³⁺ excited state, λ is the wavelength of the Ce³⁺

emission, *n* is the refractive index of the host material, $|\langle f|\mu|i\rangle|^2$ is the matrix element connecting the initial and the final states via the electric dipole operator μ . Assuming $|\langle f|\mu|i\rangle|^2$ remains unchanged, the decay time of Ce_{II} can be calculated using $\tau_{II} = \tau_1 \cdot \lambda_{II}^3 / \lambda_I^3 = 20$ ns, which is accounting for only half of the increased decay time. Secondly, energy transfer from Ce_I to Ce_{II} emission also increases the decay time, which is also reported for Sr²⁺ co-doped LaBr₃:Ce [16]. The sudden increase in decay time for Sr²⁺ at 200 K can be linked to the thermal release of trapped electrons based on the glow curves of the inset of Fig. 2.4. However, a similar increase is then also expected for Ca²⁺ doped CeBr₃ which is not observed in Fig. 2.8. Although less pronounced, the use of co-dopants was also found to increase the decay time in LaBr₃:Ce [13]. Temperature dependent decay time data for the co-doped LaBr₃:Ce samples was not reported.

Table 2.1: Scintillation decay time components in nanoseconds of the studied $CeBr_3$ crystals and their relative contributions in percentage to the total scintillation output.

	Decay components			
Sample	Fast	Slow		
CeBr ₃	17 (100 %)	-		
CeBr ₃ :Na	17 (100 %)	-		
CeBr ₃ :Mg	19 (100 %)	-		
CeBr ₃ :Ca	24 (97 %)	223 (3 %)		
CeBr ₃ :Sr	23 (97 %)	211 (3 %)		
CeBr ₃ :Ba	19 (98 %)	86 (2 %)		



Figure 2.8: Temperature dependence of the scintillation decay time constant of undoped and Na, Mg, Ca, Sr and Ba doped CeBr₃ crystals. The scintillation decay time constants are obtained by fitting the decay profiles with a single exponential component.

2.4. Summary and Conclusion

The scintillation properties of Na⁺, Mg^{2+} , Ca^{2+} , Sr^{2+} and Ba^{2+} doped $CeBr_3$ crystals were evaluated. Three categories of dopants were classified: 1) Ca^{2+} and Sr^{2+} which showed the largest effect on the scintillation properties; 2) Ba^{2+} showed a moderate effect; and 3) Na⁺ and Mg²⁺ did not influence the scintillation properties. X-ray excited emission spectra and photoluminescence excitation and emission measurements of Ca²⁺ and Sr²⁺ doped CeBr₃ showed emission from two different Ce³⁺ emission centers. Additional to a standard, unperturbed Ce³⁺ site with the same optical properties as the Ce³⁺ in undoped CeBr₃, a perturbed Ce³⁺ site was observed. The 5d \rightarrow 4f emission bands of the perturbed Ce³⁺ sites are red-shifted by 30 nm as compared to the unperturbed Ce³⁺ sites. The light yield at temperatures of 300 K and below remained constant for the Ca²⁺ and Sr²⁺ doped CeBr₃ while the undoped, Na⁺, Mg²⁺ and Ba²⁺ doped CeBr₃ showed an 15-25% increased light yield at low temperatures. Thermoluminescence measurements showed multiple glow peaks between 120 and 150 K for Ca²⁺, Sr²⁺ and Ba²⁺ doped CeBr₃. Also an increase in decay time was observed for Ca²⁺ (24 ns) and Sr²⁺ (23 ns) doped CeBr₃ compared to undoped CeBr₃ (17 ns) at 300 K. The different behavior of the scintillation properties of Ca^{2+} and Sr^{2+} (and to a lesser extend Ba^{2+}) doped $CeBr_3$ was explained using a single trap model in which Br vacancies, formed as charge compensation for the aliovalent dopants, act as electron traps. The obtained results were found to be consistent with the results reported for LaBr₃:Ce with the same aliovalent dopants.

Interestingly, adding a small amount of aliovalent dopant ions has a pronounced effect on the scintillation properties of CeBr₃, which can be ascribed to the formation of electron traps. Especially interesting is the more proportional scintillation response resulting in a better energy resolution of 3% for Sr doping (reported in a previous paper), resulting in an outstanding scintillator for low intensity gamma detection. As mentioned in the introduction, the improved proportionality can be linked to the presence of these electron traps by reducing quenching of free charge carriers. A better understanding of dopant induced charge carrier traps may provide valuable insight in scintillation nonproportionality and ultimately lead to improvements in next generation scintillators and is therefore the scope for further research.

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X-ray induced valence change and vacuum referred binding energies of Bi³⁺ and Bi²⁺ in Li₂BaP₂O₇

Bismuth-doped Li₂BaP₂O₇ was prepared in air, showing no Bi³⁺ or Bi²⁺ related photoluminescence even at 10 K. Absorption measurements showed that only Bi³⁺ was present in the as-prepared samples of which the emission is completely quenched. During Xray excitation the characteristic deep-red radioluminescence of Bi²⁺ was observed. After X-ray irradiation, this red luminescence of Bi²⁺ could be excited optically indicating that upon X-ray irradiation the Bi³⁺ is reduced to Bi²⁺. Based on the spectroscopic results, the Bi³⁺ and Bi²⁺ energy levels were estimated in a vacuum referred binding energy (VRBE) scheme and were used to explain the observed luminescence behavior. The VRBE scheme provided an interpretation for the commonly observed Bi³⁺ pair emission in bismuth doped compounds. In the case of Li₂BaP₂O₇:Bi it was used to explain the self-quenching behavior of Bi³⁺. These findings show that is possible to initially dope compounds with Bi³⁺ ions while only radioluminescence is observed from Bi²⁺ when the sample is exposed to high-energy excitation. This phenomenon can be used to fabricate new types of luminescent materials.

This chapter is based on the publication: **R. H. P. Awater** and P. Dorenbos, "X-ray induced valence change and vacuum referred binding energies of Bi^{3+} and Bi^{2+} in $Li_2BaP_2O_7$ ", *Journal of Physical Chemistry C* **120** (2016), 15114.

3.1. Introduction

Bi³⁺ is a well-known activator and sensitizer in luminescent materials with emission usually located in the ultraviolet. On the contrary, data on Bi²⁺ as an activator ion are limited to only a few compounds due to the difficulties of stabilizing Bi²⁺ over Bi³⁺. In 1994, Blasse *et al.* discovered unusual orange luminescence in SrB₄O₇:Bi, which they attributed to divalent bismuth luminescence [1]. Bi²⁺ has the [Xe]4f¹⁴5d¹⁰6s²6p¹ electron configuration with the ²P ground state. The ²P level is split into the ²P_{1/2} ground state and ²P_{3/2} excited state by spin-orbit splitting (ϵ_{so}), which is illustrated in Fig. 3.1. The ²P_{3/2} level is further split into two sublevels, ²P_{3/2}(1) and ²P_{3/2}(2), by crystal field splitting (ϵ_{cfs}). At higher excitation energy, the allowed transition to the ²S_{1/2} (6p \rightarrow 7s) is observed. Transitions within the ²P states are parity forbidden but when incorporated in a crystal lattice, electron–phonon coupling will admix with the ²P and ²S wave functions, lifting the parity selection rule [2]. The characteristic orange-to-red emission of Bi²⁺ is due to the ²P_{3/2}(1) \rightarrow ²P_{1/2} transition and is attractive for LED and other photonic materials [3].



Figure 3.1: The energy levels of Bi²⁺ and the effects of spin–orbit splitting (ϵ_{so}) and crystal field splitting (ϵ_{cfs}).

Since 1994, Bi²⁺ luminescence has been reported in alkaline earth fluorides, sulfates, phosphates, borophosphates and borates [4–10]. Interestingly, these Bi²⁺-doped compounds were synthesized under oxidizing conditions (*i.e.* air atmosphere), whereas usually a reducing agent (*e.g.* H₂ or CO) is required to stabilize the divalent state. The common factor of these host lattices is that they contain Ba²⁺ or Sr²⁺ sites and are built up of a stiff network consisting of tetrahedral anion groups. According to Peng *et al.*, both appear to be crucial for stabilizing Bi²⁺ over Bi³⁺ by creating a driving force for the internal reduction and shielding of the divalent bismuth from oxygen attack, even in oxidizing atmospheres [11]. Furthermore, Peng *et al.* have shown that reduction of Bi³⁺ to Bi²⁺ is also possible in overstoichiometric barium phosphates, as an alternative to using reducing conditions during synthesis. As this approach does not require expensive and

sophisticated synthesis equipment, it is preferable to the standard approach of synthesizing in reducing atmospheres. Unfortunately, it is restricted to a limited number of specific host lattices.

Recently, also deep-red radioluminescence of a $Sr_2P_2O_7$:Bi²⁺ sample was reported which spectrally matches with silicon photomultipliers [12]. This red radioluminescence of Bi²⁺ doped compounds could potentially be used in next-generation scintillator materials. For this study, we selected bismuth-doped Li₂BaP₂O₇ to investigate the radioluminescence of Bi²⁺. Interestingly, we discovered that X-ray induced reduction of Bi³⁺ to Bi²⁺ in Li₂BaP₂O₇ is possible. Photoluminescence and thermoluminescence measurements were performed to investigate the valence change of the bismuth activator. The experimental data were used to estimate the energy level locations of Bi²⁺ and Bi³⁺ within a constructed vacuum referred binding energy scheme.

3.2. Experimental

Samples of $Li_2Ba_{1-x}P_2O_7$: Bi_x (x = 0.0, 0.004, 0.007 and 0.01) were prepared by means of conventional solid-state reactions. Stoichiometric amounts of the starting materials Li₂CO₃ (99.997%), BaCO₃ (99.999%), (NH₄)H₂PO₄ (99.999%) and Bi₂O₃ (99.999%) were thoroughly ground in acetone. Subsequently, the ground powders were sintered at 750 °C for 15 h in air. The phase purity of the samples was checked by X-ray diffraction using a PANalytical XPert PRO X-ray diffraction system with a Co K_{α} (λ = 0.71073 Å) X-ray tube. The measured patterns were compared with reference patterns from the Pearson's Crystal Database to identify the formed phase and impurities. X-ray excited emission spectra were recorded using an X-ray tube with Cu anode operating at 60 kV and 25 mA. Each sample as a pressed powder pill received an equal amount of X-ray dose during the measurement. The emission of the samples was focused via a quartz window and a lens on the entrance slit of an ARC VM504 monochromator, dispersed, and recorded with a Hamamatsu R943-02 PMT. The spectra were corrected for the monochromator transmission and the quantum efficiency of the PMT. Photoluminescence excitation and emission measurements were recorded using a Newport 66921 Xe lamp in combination with a Horiba Gemini 180 monochromator. The emission light of the samples was dispersed with a Princeton Instruments Acton SP 2300 monochromator and detected by a Perkin Elmer Photon Counting Module MP1993. Thermoluminescence measurements were recorded using a RISØ TL/OSL reader model DA-15 with a DA-20 controller. The samples were irradiated in the X-ray setup described above and were transported to the TL setup in complete darkness to prevent photobleaching. The diffuse reflectance measurements were recorded on a home-built setup using an Acton Research Corporation deuterium lamp and an Ocean Optics optical fiber detector. The reflectance measurements were corrected for the background and lamp intensity using a BaSO4 reference sample and were converted into the absorption spectra via the Kubelka-Munk method.

3.3. Results and discussion

3.3.1. Phase identification of $Li_2Ba_{1-x}P_2O_7$: Bi_x

Fig. 3.2 shows the XRD patterns of $Li_2BaP_2O_7$:0.7% Bi and a reference pattern from Pearson's Crystal Database No#1003786. The diffraction peaks of the sample match the ref-
erence data very well, indicating a phase pure sample. $\text{Li}_2\text{BaP}_2O_7$ crystallizes in the monoclinic crystal system with space group C2/c and has a calculated density of 3.51 g/cm³ [13]. The Li₂BaP₂O₇ lattice has one crystallographic Ba-site and three different crystallographic Li-sites. Considering the ionic radii, the large bismuth ion will most likely occupy the bigger Ba-site.



Figure 3.2: XRD patterns of Li₂BaP₂O₇:0.7% Bi and the reference No#1003786.

3.3.2. X-ray excited luminescence

Fig. 3.3 shows the X-ray excited emission spectra of the $Li_2BaP_2O_7$:Bi samples measured at room temperature. The $Li_2BaP_2O_7$ host lattice (x = 0.0%) only shows emission at 375 nm, which was assigned to an excitonic origin. This 375 nm emission band decreases with increasing bismuth concentration due to the competition with the recombination of the charge carriers on bismuth centers. The bismuth-doped samples show an additional emission band centered at 685 nm and is most intense at a concentration of 0.7 mol% Bi. The red radioluminescence cannot be ascribed to impurities as the undoped sample shows no emission in this region. Furthermore, it is well known that Bi³⁺ shows emission in the ultraviolet in other phosphate compounds [14]. On the other hand, Bi²⁺ luminescence is usually located in the red. The red radioluminescence of the studied $Li_2BaP_2O_7$:Bi samples is therefore assigned to Bi²⁺. Li *et al.* found similar red radioluminescence in Sr₂P₂O₇:Bi which they assigned to Bi²⁺ [12]. When the integrated emission of the Bi²⁺ radioluminescence is compared with a BaF₂ reference sample, estimated values of the light yield are found: 300, 1300 and 500 photons/MeV of absorbed X-ray energy for 0.4%, 0.7% and 1.0% of Bi, respectively.

The temperature dependence of the X-ray excited emission of $\text{Li}_2\text{BaP}_2\text{O}_7$:0.7% Bi is shown in Fig. 3.4. Starting from 100 K and increasing the temperature to 600 K, decreases the emission intensity and red-shifts the 680 nm main emission band by almost 10 nm. Furthermore, at temperatures above 200 K a second emission band at 644 nm appears, which increases in intensity when the temperature is increased. Since $\text{Li}_2\text{BaP}_2\text{O}_7$ has only one crystallographic Ba-site, emission from different occupied sites like in $\text{Sr}_2\text{P}_2\text{O}_7$:Bi is therefore excluded [15]. The energy difference between the main emission band and the higher energy sideband is approximately 1000 cm⁻¹ (0.12 eV). The high-energy sideband is therefore attributed to a vibronic transition involving the phosphate stretching vibration which is around 1100 cm⁻¹ in energy. As the temperature increases, the transition from the higher vibronic state becomes more probable resulting in the increased intensity of the sideband.



Figure 3.3: X-ray excited emission spectra of $Li_2Ba_{1-x}P_2O_7$: Bi_x (x = 0.0, 0.4, 0.7 and 1.0 mol%) measured at room temperature.



Figure 3.4: X-ray excited emission spectra of Li₂BaP₂O₇:0.7% Bi as function of temperature.

3.3.3. X-ray induced reduction of Bi³⁺ to Bi²⁺

No Bi³⁺ or Bi²⁺ luminescence was observed for the as-prepared Li₂BaP₂O₇:Bi samples under UV-vis photon excitation. The absorption spectrum of the Li₂BaP₂O₇:Bi in Fig. 3.5 shows the Bi³⁺ A-band absorption at 250 nm and no Bi²⁺ related absorption bands, indicating that initially only Bi³⁺ is present in the samples. Probably the Bi³⁺ emission is completely quenched as is the case in Ba₃(PO₄)₂ [11]. A possible explanation for the quenching of the Bi³⁺ luminescence via the formation of Bi³⁺-Bi³⁺ pairs is presented in the next section. The absorption band at 340 nm is observed in both the bismuth doped and undoped Li₂BaP₂O₇ sample and is related to absorption of defects in the host lattice.



Figure 3.5: UV – VIS absorption spectra of as-prepared $\text{Li}_2\text{BaP}_2\text{O}_7$ with 0.0 mol% Bi (undoped) and 1.0 mol % Bi.

Interestingly, after exposure to X-rays, the red Bi²⁺ luminescence as seen previously in the radioluminescence spectra can be excited optically. The photoluminescence emission spectra of Li₂BaP₂O₇:0.7% Bi using λ_{ex} = 270 nm after different X-ray exposure durations are presented in Fig. 3.6. Before X-ray irradiation (0 min), no Bi²⁺ related emission is observed. Already after 15 min of X-ray irradiation, an emission band at 667 nm appeared which is due to the ${}^{2}P_{3/2} \rightarrow {}^{2}P_{1/2}$ transition of Bi²⁺. With longer irradiation time, the luminescence intensity is enhanced and does not seem to saturate even after 5 h of irradiation time. This indicates that during X-ray exposure the bismuth activator ions are reduced, *i.e.* the X-ray induced reduction of Bi^{3+} to Bi^{2+} . This is similar to the production of the isoelectronic ions Tl⁰ and Pb⁺ using high-energy irradiation reported in the literature [16, 17]. The intensity of the Bi²⁺ luminescence increases as the irradiation time is increased, indicating that the concentration of Bi²⁺ ions has increased. However, measuring the optical absorption spectrum of the irradiated sample resulted in an identical absorption spectrum as the sample before irradiation, showing only Bi³⁺ related absorption bands. The amount of Bi²⁺ formed during X-ray irradiation is probably very low. The photoluminescence measurements were performed directly after the exposure to X-rays. Measuring the photoluminescence of the samples 12 h after the X-ray exposure showed that the Bi²⁺ photoluminescence had completely faded. This indicates that the captured electrons slowly disappear over time.



Figure 3.6: Bi^{2+} emission ($\lambda_{ex} = 270$ nm) as function of X-ray irradiation time of $Li_2BaP_2O_7$:0.7% Bi recorded at room temperature.

The photoluminescence excitation spectrum of the $Li_2BaP_2O_7$:0.7% Bi sample at 10 K after 5 h of X-ray irradiation is shown in Fig. 3.7a. Monitoring the red emission, three excitation bands at 272 nm (4.56 eV), 419 nm (2.96 eV) and 631 nm (1.96 eV) are observed. These three excitations are characteristic for Bi²⁺ and are assigned to transitions into the ${}^2S_{1/2}$, ${}^2P_{3/2}(2)$ and ${}^2P_{3/2}(1)$ excited states of Bi²⁺, respectively (see Fig. 3.7b).



Figure 3.7: Excitation spectrum at λ_{em} = 676 nm of the Li₂BaP₂O₇:0.7% Bi sample irradiated for 5 h recorded at 10 K.

In order to study the recombination mechanism in more detail, thermoluminescence measurements were performed. Fig. 3.8 shows the thermoluminescence (TL) emission spectrum $\text{Li}_2\text{BaP}_2\text{O}_7$:0.7% Bi after 1.5 h of X-ray irradiation. The TL glow peaks are located at 455 K and 496 K with a shoulder band on the higher temperature side. Dobrowolska *et al.* observed almost identical glow peaks in $\text{Li}_2\text{BaP}_2\text{O}_7$:Eu and estimated trap depths of 1.2 eV and 1.4 eV for electron trapping centers with glow peaks at 466 K and 514 K, respectively [18]. Only Bi²⁺ emission at 680 nm is observed, indicating the recombination occurs exclusively on Bi³⁺ ions. Upon thermal stimulation, electrons are released from the electron traps and are captured on Bi³⁺, forming Bi²⁺ in the excited state. De-excitation results in the observed Bi²⁺ thermoluminescence.



Figure 3.8: TL emission spectrum of $Li_2BaP_2O_7$:0.7% Bi recorded after X-ray irradiation for 1.5 h.

3.3.4. VRBEs of Bi^{3+} and Bi^{2+} in $Li_2BaP_2O_7$

The vacuum referred binding energy (VRBE) scheme of $\text{Li}_2\text{BaP}_2\text{O}_7$ including the estimated values for the Bi³⁺ and Bi²⁺ energy levels is shown in Fig. 3.9. The values for the conduction band bottom (–0.88 eV) and valence band top (–9.23 eV), and the method to construct the VRBE scheme of $\text{Li}_2\text{BaP}_2\text{O}_7$, are taken from Ref. [18] and references therein.

 ${\rm Bi}^{2+}$ emission originating from the ${}^{2}{\rm P}_{3/2}(1)$ excited state was observed in the X-ray excited emission spectra in Fig. 3.3 and 3.4. Therefore, the ${}^{2}{\rm P}_{3/2}(1)$ excited state should be located below the bottom of the conduction band. We estimate that the ${}^{2}{\rm P}_{3/2}(1)$ excited state is located at least 1 eV below the conduction band bottom to account for the emission at elevated temperatures. Based on the excitation energies from Fig. 3.7, the ${}^{2}{\rm P}_{1/2}$ ground state and the ${}^{2}{\rm P}_{3/2}(1)$, ${}^{2}{\rm P}_{3/2}(2)$ and ${}^{2}{\rm S}_{1/2}$ Bi²⁺ excited states are then located



Figure 3.9: Vacuum referred binding energy scheme showing the Bi^{3+} and Bi^{2+} energy levels in the $Li_2BaP_2O_7$ host lattice. The arrows and luminescence mechanism are explained in the text.

relative to the vacuum energy at -3.84 eV, -1.88 eV, -0.88 eV and 0.72 eV, respectively. This corresponds with the determined impurity level of Bi²⁺ in MgGeO₃:Mn,Bi of Katayama *et al.*, who placed the the Bi²⁺ ²P_{1/2} ground state at -3.7 eV relative to the vacuum level based on thermoluminescence measurements [19]. In order to estimate the Bi³⁺ ground state, the metal-to-metal charge transfer transition (MMCT) of Bi³⁺ to the conduction band can be used [20, 21]. Unfortunately, Li₂BaP₂O₇:Bi did not show any Bi³⁺ related emission nor was a MMCT transition observed. In order to position the Bi³⁺ energy level locations, the ³P₁ excited state is estimated at -2.38 eV, 1.5 eV below the conduction band and deeper than the Bi²⁺ excited state. Using the A-band absorption of 4.96 eV (250 nm) from Fig. 3.5 the ¹S₀ ground state is then located at -7.34 eV. The MMCT transition would then be around 6.46 eV, which is higher than the predicted value by Boutinaud *et al.* but corresponds to the Bi³⁺ MMCT energy in other phosphates [21]. These estimations are based on a literature study on the energy levels of Bi³⁺ in inorganic compounds which will be published elsewhere.

Upon X-ray irradiation, electrons from the valence band are excited over the bandgap to the conduction band of the host compound (arrow 1 in Fig. 3.9). These liberated electrons migrate via the conduction band (arrow 2) and are captured on Bi^{3+} ions forming Bi^{2+} . The holes migrate via the valence band and can be captured on another Bi^{3+} ion, forming Bi^{4+} (arrow 3). Interestingly, based on the energy level locations, Bi^{3+} can act as both electron and hole trap. Relaxation to the $^2P_{3/2}$ excited state (arrow 4) and deexcitation to the ground state (arrow 5) results in the red radioluminescence observed in Fig. 3.3.

By placing the bismuth levels in the VRBE scheme using these estimated energy values, the ${}^{3}P_{1}$ excited state of Bi³⁺ is located above the ${}^{2}P_{1/2}$ ground state of Bi²⁺. From literature it is known that ions with the *ns*² electronic configuration frequently enter the host lattice in the form of pairs [22, 23]. Especially at higher bismuth concentrations, it is speculated that the formation of Bi³⁺ dimers (when two Bi³⁺ ions occupy nearest cation sites) can quench the emission. Furthermore, energy transfer from an isolated Bi³⁺ ion to Bi³⁺ pairs is known to be possible [24]. Considering such a Bi³⁺ pair, the VRBE scheme shows that upon excitation in the A-band it is possible that the excited electrons jumps to a neighboring bismuth ion (arrow 6) forming a Bi²⁺ ion and Bi⁴⁺ ion:

$$(Bi^{3+})^* + Bi^{3+} \to Bi^{2+} + Bi^{4+} \to Bi^{3+} + Bi^{3+}$$
 (3.1)

This is not a stable situation and eventually the electron will jump back (arrow 7), returning to the initial Bi^{3+} pair in the ground state. Process 7 can be radiative resulting in a broad luminescence band in the visible region, which is the case in for example LaBO₃, La₂O₃ and LaOCI [25–27]. In the case of Li₂BaP₂O₇, process 7 occurs nonradiatively and as a result there is no bismuth pair emission. Actually, this nonradiative return to the ground state is probably the main reason for the completely quenched Bi^{3+} emission in Li₂BaP₂O₇:Bi.

3.4. Conclusions

Bismuth enters the Li₂BaP₂O₇ host lattice as Bi³⁺ of which the emission is completely quenched. The X-ray excited emission spectra of the Li₂BaP₂O₇:Bi samples showed luminescence originating from Bi2+ ions. After X-ray irradiation, this red luminescence of Bi^{2+} could be stimulated optically. Therefore, part of the Bi^{3+} ions are reduced to Bi^{2+} upon exposure to X-rays. The spectroscopic data in combination with data from literature were used to locate the Bi²⁺ and Bi³⁺ energy levels within a constructed vacuum referred binding energy (VRBE) scheme. The VRBE scheme showed that the excited state of Bi³⁺ is located above the ground state Bi²⁺. This revealed that for bismuth pairs, an electron from an excited Bi³⁺ ions can be transferred to a neighboring Bi³⁺ ion. This was used to interpret the additional luminescence associated with bismuth pair emission which is commonly observed in bismuth doped materials. In the case of Li₂BaP₂O₇:Bi, the formation of bismuth pairs leads to quenching of the Bi³⁺ luminescence by a nonradiative decay to the ground state. Interestingly, this work shows that even if bismuth is initially incorporated in its trivalent state, under high-energy irradiation, emission from the divalent state is possible. This could open up new ways of fabricating luminescent materials.

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4

Luminescence and charge carrier trapping in YPO₄:Bi

 YPO_4 doped with Bi³⁺ and/or Tb³⁺ samples were prepared in air. X-ray excited luminescence measurements showed emission from isolated Bi³⁺ and Bi-pairs, and also emission from Bi²⁺ was observed. Based on the obtained spectroscopic data, the electron binding energies in the ground and excited states of Bi³⁺ and Bi²⁺ were placed inside the vacuum referred binding energy (VRBE) scheme, and this was used to explain the luminescence of bismuth doped YPO₄. The VRBE scheme and additional thermoluminescence glow curves show that bismuth can act both as electron and as hole trap in YPO₄.

This chapter is based on the publication: **R. H. P. Awater**, L. C. Niemeijer-Berghuijs and P. Dorenbos, "Luminescence and charge carrier trapping in YPO₄:Bi", *Optical Materials* **66** (2017), 351.

4.1. Introduction

 ${\rm Bi}^{3+}$ is a well-studied activator and sensitizer for luminescent materials [1]. The ${\rm Bi}^{3+}$ ion has the 6s² outer electron configuration, and optical transitions to three 6s6p excited states, labeled as A-, B- and C-bands, can be observed. Also a metal-to-metal charge transfer (MMCT) from the ${\rm Bi}^{3+}$ ground state to the host conduction band (CB) is commonly observed. In addition to the characteristic A-band emission, a lower energy emission band is frequently reported (see column 8 in Table 5.1). There is still controversy about the origin of this lower energy emission band. In a recent evaluation, Boutinaud tentatively assigned it to CB \rightarrow Bi^{3+/4+} CT luminescence [2]. Alternatively, Srivastava assigned the additional emission band in Y₂Sn₂O₇ and GdAlO₃ to emission from Bi³⁺–Bi³⁺ pairs or clusters [3, 4]. Also Wolfert *et al.* proposed in the 1980s that bismuth pairs can be the origin of the additional emission observed in Bi³⁺-doped rare earth borates and oxychlorides [5, 6].

In 1994, Blasse *et al.* confirmed the luminescence of the unique divalent bismuth oxidation state in SrB_4O_7 and $BaSO_4$ [7, 8]. Later, Bi^{2+} luminescence was also reported in alkaline-earth fluorides, sulfates, phosphates and borates for white-light LED applications [9–13]. Divalent bismuth shows luminescence in the orange-red to infrared part of the spectrum. With excitation bands in the near ultraviolet (NUV) and blue, Bi^{2+} -doped phosphors are ideal to be used in combination with an InGaN LED in order to make white light LEDs [14]. Furthermore, the red radioluminescence reported for $Sr_2P_2O_7$: Bi^{2+} spectrally matches with the sensitivity of silicon-based semiconductor detectors, making Bi^{2+} a promising activator ion for the next-generation long wavelength emitting scintillator materials [15]. The difficulty is to stabilize Bi^{2+} in the host lattice. Usually, a reducing atmosphere is required. Peng *et al.* showed that Bi^{2+} can also be stabilized in over-stoichiometric barium phosphates [16]. In chapter 3 it was shown that X-ray induced reduction of Bi^{3+} to Bi^{2+} is possible in Li₂BaP₂O₇:Bi. In this chapter, radioluminescence of divalent bismuth originating from bismuth ions doped on trivalent host lattice sites is shown.

In order to gain more insight in the trapping and luminescence properties of bismuth, we selected YPO₄:Bi as our compound of interest. YPO_4 is a wide band gap compound with only one crystallographic Y-site where Bi³⁺ can be situated. Its electronic structure is well-established from lanthanide spectroscopy and also the spectroscopy of YPO₄:Bi has been discussed in literature [17–19]. In order to estimate the VRBE in the Bi²⁺ ground state in YPO₄, thermoluminescence measurements of co-doped YPO₄:Bi,Tb were performed. We discovered that bismuth acts as both electron and hole trap, and that in the case of YPO₄, the hole is trapped more shallower than the electron.

4.2. Experimental

Samples of YPO₄ doped with varying amounts of Bi³⁺ and/or Tb³⁺ were prepared using conventional solid-state sintering. Stoichiometric amounts of the starting materials Y_2O_3 (99.99%), (NH₄)H₂PO₄ (99.999%), Bi₂O₃ (99.999%) and Tb₂O₃ (99.999%) were thoroughly ground in acetone. The ground powders were first sintered at 400 °C for 10 h in air. After an intermediate grinding step in acetone, the samples were sintered at 1400 °C for 4 h in air. X-ray diffraction was used to check the phase purity of the samples us-

ing a PANalytical XPert PRO X-ray Diffraction system with a Co K_{α} (λ = 0.71073 Å) X-ray tube. The recorded patterns were compared with a reference pattern from the Pearson's Crystal Database to identify the formed phase and possible impurities. The X-ray excited luminescence (XEL) spectra were recorded using an X-ray tube with Cu anode operating at 60 kV and 25 mA. The emission of the samples was focused via a quartz window and a lens on the entrance slit of an ARC VM504 monochromator, dispersed, and recorded with a Hamamatsu R943-02 PMT. The recorded spectra were corrected for the monochromator transmission and the quantum efficiency of the PMT. Photoluminescence excitation and emission measurements were recorded using a Hamamatsu L1835 D₂ lamp in combination with an ARC VM502 monochromator. The emission light of the samples was dispersed with a Princeton Instruments Acton SP 2300 monochromator and detected by a PerkinElmer Photon Counting Module MP1993. Thermoluminescence spectra were recorded using a RISØ TL/OSL reader model DA-15 with a DA-20 controller using a heating rate of 5 K/s. The samples were irradiated for 2 h using a ⁶⁰Co Gamma Cell 220 source with a dose rate of 0.29 Gy/s and were transported to the TL setup in complete darkness to prevent photobleaching.

4.3. Results

The X-ray diffraction patterns of YPO_4 :1.0% Bi and a reference pattern from Pearson's Crystal Database No#1828017 are shown in Fig. 4.1. The diffraction peaks of the sample match well with that of the reference and no impurity phases are observed.

Figure 4.1: XRD patterns of YPO₄:1.0% Bi and the reference No#1828017.

The photoluminescence excitation and emission spectra of YPO_4 :1%Bi sample are shown in Fig. 4.2. The excitation spectrum was recorded while monitoring the 242 nm A-band emission and the emission spectrum was recorded while exciting at the 227 nm A-band absorption. Both spectra were recorded at room temperature. The 122 nm band in the excitation spectrum is attributed to the formation of free electrons and holes, *i.e.* the energy difference E_{VC} between the top of the valence band and the bottom of the conduction band. The wavelength of the host exciton creation in YPO₄ is well-established

at 145 nm from lanthanide spectroscopy [20]. At this wavelength a dip is observed in the excitation spectrum, indicating inefficient energy transfer from the exciton to Bi^{3+} . In the region between 150 and 200 nm excitation bands can be distinguished at 157 nm and 170 nm, which are attributed to the C-band excitation and $Bi^{3+} \rightarrow CB$ CT transitions, respectively. In the work by Cavalli *et al.* on YPO₄:Bi, only an unresolved broad band was reported in the region 150-200 nm [17]. The emission spectrum shows besides the 242 nm Bi^{3+} A-band emission a weak emission peaking at 325 nm. This emission has been attributed to Bi-pair emission by Srivastava *et al.*, and they observed that the emission intensity increases with increasing Bi^{3+} concentration [19]. We observed no significant increase in the intensity of the Bi-pair emission band with varying the Bi^{3+} concentration between 0.05 and 2.0%.

Figure 4.2: Photoluminescence excitation ($\lambda_{em} = 242 \text{ nm}$) and emission ($\lambda_{ex} = 227 \text{ nm}$) spectra of YPO₄:1%Bi³⁺ measured at room temperature.

The X-ray excited emission spectra of the $\mathrm{Bi}^{3+}\text{-}\mathrm{doped}\ \mathrm{YPO}_4$ at room temperature are shown in Fig. 4.3. Undoped YPO_4 shows no radioluminescence in the measured region of 200 nm to 800 nm. The bismuth-doped samples show the A-band emission at 242 nm, as well as additional emission bands at 300-550 nm and 670 nm. The A-band emission is most intense for a Bi³⁺ concentration of 0.05 mol% and decreases in intensity with increasing concentration of Bi³⁺. Srivastava et al. observed a similar concentration dependence of the A-band emission in photoluminescence measurements [19]. The emission band observed at 670 nm has not been reported before for YPO_4 . We attribute it to the ${}^2P_{3/2} \rightarrow {}^2P_{1/2}$ transition of Bi^{2+} caused by the trapping of an electron from the conduction band by Bi^{3+} to populate the excited Bi^{2+} state. Also the Bi^{2+} luminescence is most intense for the lowest concentration of 0.05% bismuth and decreases with increasing concentration. The broad 300-550 nm emission band centered at 405 nm is probably due to electron hole recombination near an unknown defect, i.e. a near defect exciton (NDE) emission. A similar NDE emission band was observed by Bos et al. in YPO_4 : Tb^{3+} [21]. The intensity of the NDE emission band increases with increasing Bi³⁺ concentration. For a concentration of 2% bismuth, the NDE emission band has disappeared and a new band at 340 nm appears. This new band is the same as the pair emission observed in the photoluminescence spectra in Fig. 4.2.

Figure 4.3: X-ray excited emission spectra of YPO_4 :Bi_x (x = 0.0, 0.05, 0.25 and 1.0 mol%) measured at room temperature. The asterisk at 480 nm marks the grating's second order reflection of the 242 nm A-band emission.

Fig. 4.4 shows the temperature dependence of the A-band, pair-, Bi^{2+} and NDE emission under X-ray excitation for the sample with the lowest Bi concentration (0.05% Bi). The A-band gradually increases about 10% in intensity from 85 K up to 250 K. Then, the intensity increases rapidly to approximately 80% at 350 K and reaching 100% at 450 K, after which the intensity decreases again to 50% at 600 K. At temperatures below 170 K, NDE emission dominated the pair emission band. It disappears above 170 K and the Bi-pair emission starts to dominate. At the same temperature the Bi^{2+} emission appears and grows until reaching maximum intensity at 300 K. Above 300 K, the Bi^{2+} emission decreases until it is absent at 450 K.

Figure 4.4: Temperature dependence of the A-, pair-, Bi^{2+} -, and NDE emission bands derived from X-ray excited luminescence measurements of YPO₄:0.05%Bi. The intensity at the emission maximum is shown normalized to unity for each band.

Fig. 4.5 compares the X-ray excited emission spectra for $YPO_4:1\%Tb^{3+}$ and $YPO_4:1\%Tb^{3+},1\%Bi^{3+}$ at room temperature. The bands labeled 1 and 2 originate from the Bi^{3+} A-band and NDE emission, respectively. The bands labeled 3 - 14 originate from Tb^{3+} 4f–4f transitions. Compared to the $YPO_4:1\%Tb^{3+}$ sample, the $YPO_4:1\%Tb^{3+},1\%Bi^{3+}$ shows an increased intensity of the ${}^5D_4 \rightarrow {}^7F_{6-0}$ transitions (peaks nr. 8–14), while the ${}^5D_3 \rightarrow {}^7F_{6-2}$ transitions (peaks nr. 3–7) have slightly lower or the same intensity.

Figure 4.5: X-ray excited emission spectra of YPO_4 : $1\%Tb^{3+}$ and YPO_4 : $1\%Bi^{3+}$, $1\%Tb^{3+}$ measured at room temperature.

The thermoluminescence (TL) emission spectra of YPO₄:1%Tb³⁺ and YPO₄:1%Tb³⁺, 1%Bi³⁺ are shown in Fig. 4.6. The TL glow peak of the YPO₄:1%Bi³⁺ sample is located at 425 K and shows Bi³⁺ A-band emission and a weak and broad emission at around 400 nm originating from mainly NDE emission. The TL glow peaks of the YPO₄:1%Tb³⁺, 1%Bi³⁺ sample are observed at 425 K and 485 K with a shoulder glow on the higher temperature side, and it shows Bi³⁺ A-band emission, Tb³⁺ 4f-4f emission and weak broad band emission between 280 and 380 nm which is due to Bi-pair emission. No Bi²⁺ emission is observed during the TL measurements for both samples. The TL emission spectra of undoped YPO₄ and YPO₄:1%Tb³⁺ showed no thermoluminescence emission.

Figure 4.6: TL emission spectrum of a) YPO_4 :1.0% Bi and b) YPO_4 :1.0% Bi, 1.0% Tb recorded after γ -ray irradiation for 2 h using a ^{60}Co source.

4.4. Discussion

Fig 4.7 shows the vacuum referred binding energy (VRBE) scheme for YPO₄:Bi,Tb. The VRBE of the electron at the conduction band bottom ($E_C = -0.53 \text{ eV}$), the valence band top ($E_V = -9.77 \text{ eV}$), in the exciton state ($E^X = -1.22 \text{ eV}$), and in the Tb³⁺ ground and excited states are taken from Ref. [22]. In order to estimate the VRBE in the 6s² ground state of Bi³⁺, the Bi³⁺ \rightarrow CB CT energy has been used. Similar to the interpretation of the Pr³⁺ and Tb³⁺ \rightarrow CB CT-energies in Ref. [23], we regard this energy (see arrow 1) as the energy difference between the Bi³⁺ ground state and $\frac{E_C+E_X}{2}$. One obtains –8.2 eV for the VRBE in the Bi³⁺ ground state, and by adding the 5.5 eV A-band energy (see arrow 2), –2.7 eV for the VRBE in the ³P₁ excited state.

The X-ray excited emission spectra of YPO₄:Bi in Fig. 4.3 shows Bi²⁺ emission which is only possible when the ²P_{3/2}(1) excited state is located well below the conduction band. Fig. 4.4 shows that the Bi²⁺ emission starts to quench at 350 K. With a typical radiative life time of 20 μ s for Bi²⁺ emission [9, 13] one may derive, with methods similar as in Ref. [24], the quenching energy barrier $\Delta E_q \approx T_{0.9}/620$ eV. Where $T_{0.9}$ is the temperature where the intensity has dropped by 10%. It corresponds more or less with the onset of thermal quenching. For quenching, the electron does not need to go fully to the conduction band and we will assume that $\frac{E_C + E_X}{2} - \Delta E_q$ provides a fair estimate for the VRBE in the Bi²⁺ excited state. The VRBE in the ²P_{3/2}(1) excited state is then at -1.43 eV in Fig. 4.7 and using the emission energy of 1.85 eV (670 nm) (see arrow 3), the VRBE in the ²P_{1/2} ground state is at -3.3 eV.

Figure 4.7: Vacuum referred binding energy scheme of YPO_4 including the Tb^{3+} , Bi^{3+} and Bi^{2+} energy levels.

Upon X-ray irradiation, conduction band electrons can be captured by Bi^{3+} to first populate the excited state of Bi^{2+} , which is then followed by red luminescence of the ${}^{2}P_{3/2}(1) \rightarrow {}^{2}P_{1/2}$ transition (see arrow 3). Bi^{2+} is not stable in YPO_4 and will eventually recombine with a hole from the valence band or disappear by electron transfer to neighboring defects. Therefore, no Bi^{2+} related luminescence is observed under photon excitation.

The XEL spectrum of YPO₄:Bi,Tb in Fig. 4.5 shows enhanced emission intensity from the Tb³⁺ ⁵D₄ \rightarrow ⁷F₇₋₀ transitions as compared to that of single-doped YPO₄:Tb. In single doped YPO₄:Tb³⁺, the Tb³⁺ ions can be excited by first trapping a hole and later an electron, or by means of e-h pair capture, or energy transfer from the exciton or NDE state. In the presence of Bi³⁺ even more excitation paths are envisaged. Angiuli *et al.* already showed that Bi³⁺ \rightarrow Tb³⁺ energy transfer occurs in YPO₄:Bi,Tb [25]. Probably also Tb⁴⁺-Bi²⁺ pairs can be formed during X-ray exposure, and electron back transfer from Bi to Tb is more likely to populate the ⁵D₄ level.

In the TL emission spectrum of YPO₄:1%Bi in Fig. 4.6, a single glow peak is observed at 425 K which shows Bi^{3+} A-band emission at 243 nm. This indicates that the luminescence occurs from either electron capture on Bi^{4+} or from hole capture on Bi^{2+} . According to Fig. 4.7, Bi^{3+} can act as a 1.57 eV deep hole trap to form Bi^{4+} but also as a 2.97 eV deep electron trap to form Bi^{2+} . The hole will be released earlier from its trap than the electron. Besides the $Bi^{2+/3+}$ electron trap, other unknown host related electron traps may be present. Electron release from these unknown defects would generate besides Bi^{3+} emission (recombination on Bi^{4+}) also Bi^{2+} emission (recombination on Bi^{3+}). Since no Bi^{2+} emission is observed, the Bi^{3+} emission of the 425 K glow is attributed to Bi^{4+} hole release that recombines with Bi^{2+} . Supporting evidence is the absence of any thermoluminescence glow for $YPO_4:Tb^{3+}$. The formation of Bi^{4+} may seem strange since it is regarded as chemically unstable. Indeed during synthesis Bi^{4+} is never formed, however, since the Bi^{3+} ground state is above the valence band it can trap a hole to form Bi^{4+} as a meta-stable defect.

The TL emission spectrum of YPO_4 :Bi,Tb in Fig. 4.6 shows both Bi^{3+} A-band and Tb^{3+} 4f-4f glow at 425 K and 485 K. We attribute the observed Tb^{3+} glow at 425 K to hole release from Bi^{4+} and recombination on Bi^{2+} followed by $Bi^{3+} \rightarrow Tb^{3+}$ energy transfer. In the VRBE scheme Tb^{3+} is predicted to act as a 2.1 eV deep hole trap which is 0.5 eV deeper than that by Bi^{3+} . We therefore attribute the 485 K glow peak to hole release from Tb^{4+} , which recombines with the Bi^{2+} electron trap, producing Bi^{3+} A-band glow and Tb^{3+} glow via energy transfer. Also weak Bi–Bi pair emission near 320 nm is visible at 485 K. Apparently, once the hole on Tb^{4+} is released it can migrate towards an electron trapped in a Bi–Bi pair resulting in the 3.65 eV pair emission. The rather low probability of this recombination mechanism accounts for its weak intensity. In the case of YPO_4 :Bi, the pair emission is probably dominated by the NDE emission observed at around 400 nm.

4.5. Conclusions

X-ray excited emission spectra of YPO_4 :Bi samples showed luminescence originating from Bi³⁺, Bi²⁺ and Bi-pair emission. This chapter showed that characteristic red radioluminescence of Bi²⁺ is observed in YPO_4 doped with Bi³⁺. Such type of radioluminecence may be utilized as red emitting scintillator material. The spectroscopic data on Bi³⁺ and Bi²⁺ was used to determine the VRBE of the electron in the ground and excited states of the Bi²⁺ and Bi³⁺. The VRBE scheme, together with data from thermoluminescence glow curves showed that Bi³⁺ can act as both electron and hole trap in YPO_4 .

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5

The Bi³⁺ 6s and 6p electron binding energies in relation to the chemical environment of inorganic compounds

This chapter provides an overview and interpretation of the spectroscopic data of the Bi^{3+} activator ion in 117 different inorganic compounds. The energies of the metal-to-metal charge transfer and the interconfigurational transitions of Bi^{3+} were collected from the archival literature. Using these energies, in combination with the electron binding energies in the host conduction and valence band, the binding energies in the 6s ground state and 6p excited state were determined relative to the vacuum level. The locations of the Bi^{3+} energy levels within the forbidden gap of the host compound provides valuable insight in the physical properties of the Bi^{3+} activator ion in different compounds.

This chapter is based on the publication: **R. H. P. Awater** and P. Dorenbos, "The Bi³⁺ 6s and 6p electron binding energies in relation to the chemical environment of inorganic compounds", *Journal of Luminescence* **184** (2017), 221.

5.1. Introduction

The luminescence of the Bi^{3+} activator in compounds has been extensively studied over the past 50 years [1]. Depending on the type of host compound, the Bi^{3+} emission spans from the ultraviolet to the green, which makes Bi^{3+} attractive for scintillator or phosphor applications. Furthermore, Bi^{3+} is commonly used as a sensitizer for lanthanide luminescence in various luminescent materials [2–6]. Although extensively investigated, the luminescence of Bi^{3+} in relation to the type of compound is still poorly understood [7].

Like the Tl⁺ and Pb²⁺ ions, the Bi³⁺ ion has a 6s² outer electron configuration with ¹S₀ ground state. Optical transitions to the 6s¹6p¹ configuration result in ³P_{0,1,2} triplet and ¹P₁ singlet excited states, which are shown on the left side of Fig. 5.1. The optical transitions from the ¹S₀ ground state to the ³P₁, ³P₂ and ¹P₁ excited states are commonly labeled as A-, B- and C-band, respectively. The ¹S₀ \rightarrow ³P₀ and ¹S₀ \rightarrow ³P₂ are spin-forbidden, although the transition to the ³P₂ can be induced by coupling with lattice vibrational modes [8]. As a result of spin-orbit coupling and mixing with the ¹P₁ state, the A-band transition becomes allowed. The A-band is also the emitting state (³P₁) \rightarrow ¹S₀) in Bi³⁺ doped materials. A more detailed discussion on the optical transitions of 6s² ions can be found in the literature [9–11].

Figure 5.1: The interconfigurational electronic transitions, intervalence charge transfer (IVCT) transition and the metal-to-metal charge transfer (MMCT) transition of the Bi³⁺ ion.

Wang *et al.* showed a quantitative relationship between the A- and C- band excitation energies and the chemical environment [12]. Using the environment factor h_e , they showed that the larger the four parameters determining the chemical environment (coordination number, fractional covalence, bond volume polarizability and effective charge), the lower the energy of the Bi³⁺ absorption bands. In order to understand the performance of Bi³⁺ activated materials, knowledge on the location of the Bi³⁺ levels within the band gap of host compounds is required. Previous work on lanthanide spectroscopy showed systematics in lanthanide luminescence with the chemistry of compounds [13]. Based on the established models by Dorenbos, it is now possible to determine the lanthanide 4f and 5d electron binding energies relative to the vacuum en-

ergy [14]. Combined with the binding energy at the conduction band bottom and valence band top, it is not only possible to understand but also to predict the properties of lanthanide activated materials. In 2014 these models were applied to determine the vacuum referred binding energies of the 3d electron in transition metal ions [15]. In this chapter, based on the same models, the binding energies of the 6s and 6p electrons of Bi^{3+} are determined in 44 different compounds.

An important parameter required to determine the binding energy of an electron is a charge transfer transition. In the case of the lanthanides, the charge transfer of a valence band electron to a trivalent lanthanide can be used to locate the ground state of the divalent lanthanide relative to the top of the valence band [16]. Additionally, in for example Pr^{3+} or Tb^{3+} doped phosphors, an electron transfer from the 4f ground state to the conduction band is possible, which can be used to locate the ground state of the trivalent lanthanide relative to the bottom of the conduction band. These charge transfer transitions are called metal-to-metal charge transfer (MMCT) transitions [17]. Also in the case of Bi^{3+} a MMCT transition of an electron from Bi^{3+} to a host cation (M^{n+}) exists, *i.e.* $Bi^{3+}/M^{n+} \rightarrow Bi^{4+}/M^{(n-1)+}$, which is shown on the right side of Fig. 5.1 [18, 19]. Recently, Boutinaud et al. reported on the systematics of the Bi3+ MMCT transition in d^0 and d^{10} transition metal oxides [20, 21]. Another possibility proposed by Srivastava et *al.* is that Bi^{3+} ions enter the host lattice in the form of pairs or clusters [22, 23]. Considering a pair of neighboring Bi³⁺ ions, a charge transfer of an electron from one Bi³⁺ ion to its neighbor is possible, forming Bi²⁺ and Bi⁴⁺ ions. To distinguish between the charge transfer of an electron from a Bi³⁺ ion to either the conduction band or to a neighboring Bi³⁺ we introduce the term intervalence charge transfer (IVCT). Here we refer to IVCT as a charge transfer between two neighboring Bi³⁺ ions, as indicated by the dashed arrows in Fig. 5.1. The IVCT is only possible when the ground state of Bi^{2+} is located below the first excited state of Bi³⁺. In chapter 3 we suggested that the ground state of Bi^{2+} is indeed located below the ${}^{3}P_{1}$ excited state of Bi^{3+} in $Li_{2}BaP_{2}O_{7}$. Furthermore, we suggested that the intervalence charge transfer transition between neighboring bismuth ions can be responsible for the complete quenching of the Bi³⁺ A-band luminescence.

In most of the Bi^{3+} -doped compounds an additional emission band is observed at energies lower than the A-band. In the literature this emission band is commonly labeled as VIS-band, since the emission energy is usually in the visible part of the spectrum, whereas the A-band is usually located in the ultraviolet. The origin of the lower energy emission band remains controversial, however, emission from Bi^{3+} -pairs or from the charge transfer state have been suggested [20, 22]. Dedicated experiments are required to determine the binding energy of an electron in the ground state of the Bi^{2+} ion in compounds in order to make a thorough evaluation and distinction of the Bi^{3+} MMCT and IVCT transition energies.

5.2. Methodology

Spectroscopic data on the electronic transitions of Bi³⁺ in 117 different compounds have been collected from the archival literature and have been compiled in Table 5.1. The first column lists the 7-digit compound identification number that characterizes the type of compound and reflects an increasing value for the redshift of the 4f-5d transition of lan-

Figure 5.2: The U value as function of the compound identification number. O-X indicates the U value of the oxyhalides.

thanide luminescence centers for increasing values of the identification number [24]. The second column lists the compound name. Column three lists the U value of the corresponding host compounds, which are taken from previous reports [15, 25]. When the U value was not available from previous literature, it was estimated based on the U value of compounds with similar chemical composition and are shown in italics. The U value, or Coulomb repulsion energy, is defined as the binding energy difference of the electron in the 4f ground state of Eu²⁺ with that in Eu³⁺. The U value of many different compounds were derived empirically from lanthanide spectroscopy and were found to be a good representation of the chemical environment of the activator ion [26]. Within the chemical shift model, the knowledge of the U value for a specific compound provides the vacuum referred binding energies (VRBE) for an electron in the ground state for all divalent and trivalent lanthanides in that compound. The value ranges from 7.65 - 7.3 eV in fluorides, 7.2 - 6.4 eV in oxides, 6.8 - 6.6 eV in chlorides and bromides, around 6.3 eV in iodides and around or lower than 6.2 eV in sulfides and selenides. The U value is strongly connected to the nephelauxetic sequences that formed the bases for encoding compounds with the identification number, which is illustrated in Fig. 5.2. An increasing value of U corresponds to a decreasing identification number of the host compound. Here, the U value of the host compound is used to show luminescence trends when the chemical environment of the Bi³⁺ ion is changed. The excitation energies corresponding to the A-, C-, and MMCT-bands are shown in columns 4, 5 and 6, respectively. The predictability of the chemical shift model in combination with a differentiation in the spectral band widths of the MMCT (0.8 eV) and A- or C-band (0.3 eV), allowed for a reassignment of the spectroscopic bands. In Appendix A a detailed account is provided on the obtained MMCT energies that are listed in column 6. Here, the MMCT transition energies are used to locate the Bi³⁺ 6s ground state relative to the conduction band bottom in the VRBE diagrams. The A-band emission energies are shown in column 7. As mentioned in the introduction, an additional lower energy emission band is observed originating from either the IVCT (Bi³⁺ pairs) or MMCT state. Since the distinction between the emission of these charge transfer transitions is not possible with the data currently available, we labeled the lower energy emission as the CT emission band. The CT emission energies are listed in column 8. The last column shows the reference to the original literature, which needs to be consulted for additional information on the spectroscopic data.

ID				Excitat	tion	Em	ission	
number	Compound	U	А	С	MMCT	А	СТ	Ref.
0000000	free ion	_	9.42	14.21	_	-	_	[27]
1190010	BaF ₂	7.38	5.75	7.92	9.36	4.75	-	[28]
1190020	SrF ₂	7.32	5.78	8.16	9.11	5.47	-	[28]
1190030	CaF ₂	7.31	5.86	8.2	9.17	5.51	-	[28]
1190404	NaYF ₄	7.51	6.2	-	-	-	2.82	[29]
2290002	RbCl	6.8	3.44	5.44	5.9	_	_	[30]
2290003	KCl	6.8	3.76	5.85	6.39	-	-	[30, 31]
2290004	NaCl	6.7	3.81	5.85	6.26	-	-	[30]
2290041	CsMgCl ₃	6.8	4.37	-	-	-	2.96	[32]
2290101	Cs ₂ NaLaCl ₆	6.67	3.9	-	-	3.63	-	[33]
2290401	Cs ₂ NaYCl ₆	6.67	3.84	-	-	3.72	-	[34]
3390003	KBr	6.6	3.35	5.32	5.74	_	_	[30]
3390071	CsCdBr ₃	6.6	4.33	_	-	_	2.12	[32]
3390401	Cs ₂ NaYBr ₆	6.53	3.36	_	-	3.16	_	[33]
4490003	KI	6.25	3.26	5.17	5.56	_	_	[30]
5174020	Sr ₃ AlO ₄ F	6.8	3.97	_	-	2.9	_	[35]
5190400	YOF	6.7	4.62	-	6.2	3.76	-	[1]
5290100	LaOCl	6.44	3.72	-	4.61	3.6	2.79	[1]
5290300	GdOCl	6.46	3.67	_	4.82	2.79	2.5	[36]
5290400	YOCl	6.47	3.73	-	4.86	3.1	-	[1]
5390100	LaOBr	6.4	_	_	4.26	3.37	2.5	[37]
5532100	LaP ₃ O ₉	7.26	5.27	_	-	_	2.72	[38]
5532105	LiLaP ₄ O ₁₂	7.29	5.4	_	-	_	2.89	[39]
5532300	GdP ₃ O ₉	7.24	5.15	_	-	_	_	[38]
5532400	YP ₃ O ₉	7.19	5.15	_	-	4.23	_	[38]
5532500	LuP ₃ O ₉	6.99	5.15	_	-	4.29	_	[38]
5532600	ScP ₃ O ₉	6.9	4.9	_	-	4.34	-	[38]
5534100	LaPO ₄	7.18	5.06	-	7.1	-	2.75	[40]
5534400	YPO ₄	7.09	5.51	_	7.3	5.08	3.7	[41]
5534500	LuPO ₄	7.08	5.51	_	-	5.25	3.72	[8]
5550013	KBaBP ₂ O ₈	7.0	5.17	_	-	3.2	_	[42]
5552100	LaB_3O_6	7.15	4.77	-	-	3.26	-	[34]
5552140	LaMgB ₅ O ₁₀	7.15	4.19	_	-	3.69	-	[43]
5552160	LaZnB ₅ O ₁₀	7.0	4.16	-	-	3.69	-	[43]
5552170	$LaCdB_5O_{10}$	7.0	4.2	-	-	3.76	_	[43]
5552440	YMgB ₅ O ₁₀	7.17	4.63	-	_	3.8	-	[43]

Table 5.1: The spectroscopic data of Bi³⁺ in 117 different compounds. All energies are in electronvolts.

ID			Excitation		Emission			
number	Compound	U	А	С	MMCT	А	СТ	Ref.
5552460	YZnB ₅ O ₁₀	7.0	4.59	-	-	3.73	_	[43]
5552470	YCdB ₅ O ₁₀	7.0	4.63	-	-	3.77	_	[43]
5554000	GaBO ₃	6.9	4.49	-	-	4.28	2.93	[44]
5554000	InBO ₃	6.86	4.37	-	-	4.08	3.1	[45]
5554035	LiCaBO ₃	6.9	4.08	-	-	3.28	_	[46]
5554100	LaBO ₃	6.93	4.62	-	-	3.46	2.69	[47]
5554400	$YAl_3B_4O_{12}$	7.04	4.77	-	-	4.28	_	[1]
5554400	YBO ₃	6.85	5.02	-	6.2	4.22	3.87	[48]
5554500	LuBÕ ₃	6.95	4.32	-	-	3.95	_	[45]
5554600	ScBO ₃	6.86	4.26	-	-	4.14	_	[47]
5555120	SrLaBO ₄	6.8	4.77	-	-	_	2.34	[49]
5555330	$Ca_4GdO(BO_3)_3$	6.8	4	-	-	3.18	_	[50]
5555430	$Ca_4 YO(BO_3)_3$	6.82	4	5.39	-	3.26	_	[51]
5555430	CaYBO ₄	6.8	4.44	-	-	_	_	[52]
5563400	$Y_2 Sn_2 O_7$	6.75	4.43	-	-	3.76	2.43	[22]
5564025	$Li_4SrCa(SiO_4)_2$	6.8	4	5.51	-	3.26	_	[53]
5564030	CaSnO ₃	6.7	4.07	-	-	3.44	2.74	[19]
5564040	MgGeO ₃	6.75	4.27	-	4.9	3.44	_	[54]
5564060	Zn_2GeO_4	6.8	4.13	-	-	-	2.41	[55]
5564405	LiYSiO ₄	6.91	4.43	-	-	_	_	[52]
5565300	Gd_2GeO_5	6.8	4	5.28	-	2.75	_	[56]
5565400	$Y_2 SiO_5$	6.8	4.5	-	-	3.6	2.08	[57]
5565500	Lu ₂ SiO ₅	6.83	4.2	5.95	-	3.47	2.25	[58]
5570000	ZnGa ₂ O ₄	6.8	3.44	-	4.43	3.02	2.3	[59]
5570100	LaAlO ₃	6.67	4.35	-	-	3.31	_	[60]
5570100	LaGaO ₃	6.7	4.04	-	5.17	3.26	_	[61]
5570100	LaInO ₃	6.73	3.65	-	-	2.95	_	[60]
5570130	CaLaAl ₃ O ₇	6.7	4.2	-	-	_	3.1	[62]
5570300	$Gd_3Al_5O_{12}$	6.77	4.48	-	-	3.25	-	[63]
5570300	$Gd_3Ga_5O_{12}$	6.77	4.0	-	4.59	_	2.64	[64, 65]
5570300	GdÅlO ₃	6.75	4.27	-	-	3.72	2.58	[66]
5570300	Gd ₂ GaSbO ₇	6.7	4.27	-	-	3.35	-	[67]
5570400	$Y_3 \tilde{Al}_5 O_{12}$	6.77	4.7	5.68	5.93	4.09	2.64	[68]
5570400	$Y_3Ga_5O_{12}$	6.77	4.07	-	4.3	3.87	2.95	[23, 64]
5570400	YÅlO ₃	6.81	4.42	-	6.5	3.67	_	[69]
5570400	$Y_4Al_2O_9$	6.7	4.42	5.5	-	3.4	_	[70]
5570500	$Lu_3Al_5O_{12}$	6.77	4.63	-	5.95	4.15	2.56	[71]
5581030	CaMoO ₄	7	-	-	3.83	-	2.17	[20]
5582030	CaWO ₄	7.1	-	_	4.32	-	2.65	[72]
5582060	ZnWO ₄	7.1	-	_	3.65	-	2.21	[73]
5582070	CdWO ₄	7.1	-	_	3.54	-	2.25	[73]
5582400	Y ₂ WO ₆	7.1	-	-	3.63	-	2.41	[1]

Table 5.1: Continued

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Table 5.1. Continued

ID				Excita	tion	Em	ission	
number	Compound	U	А	С	MMCT	А	СТ	Ref.
5583100	LaVO ₄	6.8	-	_	3.83	-	2.26	[20]
5583300	GdVO ₄	6.8	_	_	3.78	-	2.23	[74]
5583400	YVO ₄	6.8	_	-	3.72	_	2.18	[74]
5583500	LuVO ₄	6.8	_	-	3.7	_	2.15	[75]
5583600	ScVO ₄	6.8	_	-	3.49	2.67	1.95	[76]
5584030	$CaNb_2O_6$	6.8	_	-	3.88	_	2.43	[20]
5584100	LaNbO ₄	6.8	-	-	4.07	3.02	-	[77]
5584300	GdNbO ₄	6.8	-	-	4.04	2.79	-	[78]
5584400	YNbO ₄	6.8	-	-	3.95	2.79	2.3	[79]
5585300	GdTa ₇ O ₁₉	6.7	4	-	-	2.58	-	[80]
5585400	YTaO ₄	6.7	4.27	-	4.96	2.95	-	[81]
5586030	CaTiO ₃	6.7	-	-	3.35	-	2.14	[82]
5586400	$Y_2 Ti_2 O_7$	6.7	-	-	3.87	-	2.25	[83]
5590020	SrZnO ₂	6.45	3.29	-	-	-	2.41	[84]
5590020	SrO	6.4	3.38	-	4.7	2.85	-	[85]
5590030	CaO	6.31	3.59	-	5.1	3.15	-	[27, 85]
5590030	CaZrO ₃	6.55	3.87	-	-	3.18	-	[60]
5590030	CaHfO ₃	6.55	4.03	-	-	3.25	-	[23]
5590100	La ₂ O ₃ ^a	6.45	4.03	-	4.98	2.73	2.58	[74]
5590104	NaLaO ₂	6.45	3.53	-	-	2.23	-	[86]
5590110	BaLa ₂ ZnO ₅	6.55	-	-	3.87	3.02	-	[87]
5590200	$La_2Zr_2O_7$	6.55	4.27	-	5.06	3.22	2.41	[88]
5590300	Gd ₂ O ₃ ^a	6.52	3.57	-	-	2.92	2.29	[89]
5590304	NaGdO ₂	6.5	3.61	4.9	-	3.22	-	[86]
5590305	LiGdO ₂	6.5	3.74	4.82	-	2.69	-	[86]
5590400	Y ₂ O ₃ ^a	6.46	3.73	-	4.71	3.03	2.58	[1]
5590404	NaYO ₂	6.45	3.51	4.88	-	3.22	-	[86]
5590405	LiYO ₂	6.45	3.82	-	-	2.23	-	[86]
5590500	Lu ₂ O ₃ ^a	6.5	3.33	-	-	3.07	2.42	[74]
5590504	NaLuO ₂	6.5	3.52	4.85	-	3.24	-	[86]
5590505	LiLuO ₂	6.5	3.76	-	-	2.42	-	[86]
5590600	Sc ₂ O ₃ ^a	6.45	3.7	-	4.6	3.05	2.46	[90]
5590604	NaScO ₂	6.4	3.43	4.8	-	3.25	-	[86]
5590605	LiScO ₂	6.4	3.92	4.9	-	3.06	-	[86]
6690010	BaS	6.25	2.77	3.35	-	2.21	-	[91]
6690020	SrS	6.25	2.95	-	3.75	2.6	1.55	[92]
6690030	CaS	6.17	3.01	3.54	4.2	2.75	2.05	[27]
6690040	MgS	6.22	2.98	3.83	4.4	2.84	-	[93]
7790030	CaSe	6.22	2.74	3.2	3.6	2.5	1.94	[27]

^a: In the case of the sesquioxides M_2O_3 (M = Sc, Y, La, Gd, Lu) a 75% occupation of the C_2 site was reported for the Bi³⁺ ion and therefore only the C_2 site was considered when discussing the sesquioxides [21].

5.3. Results

5.3.1. Overview of the spectroscopic data on Bi³⁺

Fig. 5.3a shows the A-band excitation energy of the Bi³⁺ ion in 117 different inorganic compounds. The compounds are categorized using the 7-digit compound identification number. A strong decrease in the A-band energy from 9.4 eV in the free bismuth ion to 2.7 eV in selenide compounds is observed. The decrease follows the nephelauxetic sequence of decreasing Pauling electronegativity of the coordinating anion [26]:

$$F^- < O^{2-} < Cl^- < Br^- < l^- < S^{2-} < Se^{2-}$$
 (5.1)

Fig. 5.3b is an expanded view of the A-band excitation energy in the 93 different oxide compounds. Now the A-band energy follows the sequence of decreasing electronegativity of the cation that binds the anion ligands:

$$P^{5+} < B^{3+} < Si^{4+} < Al^{3+} < RE^{3+} < AE^{2+}$$
(5.2)

where RE and AE are abbreviations for rare earth and alkaline earth, respectively. The decrease in the A-band excitation energy is similar to the reported redshift of the Ce^{3+} 4f-5d₁ transition, which is defined as the energy by which the 5d₁ level of Ce^{3+} is lowered relative to the 4f ground state when doped in a compound compared to the 5d₁ energy of the free Ce^{3+} ion [13, 25]. However, the variation in the A-band excitation energy of Bi^{3+} appears to be stronger when compared to the 4f-5d excitation energies of Ce^{3+} .

Figure 5.3: The A-band excitation energies of the Bi^{3+} ion in a) 117 different inorganic compounds and b) 93 different oxide compounds. TM = transition metal, RE = rare earth, AE = alkaline earth.

Fig. 5.4a shows the A- and C-band excitation energies of the Bi^{3+} ion as a function of the U value of the host compound. The A- and C-band excitation energies increase linearly with U. Two different emission bands can be observed in Bi^{3+} activated materials, namely an intense and sharp (~0.3 eV wide) A-band emission and a weaker and considerably broader (~0.8 eV wide) CT emission band. Fig. 5.4b shows the A-band and CT emission energy as function of the U value. Similar to the A-band excitation energy, the A-band emission energy varies systematically with changing chemical environment. The A-band emission energy decreases from 6.5 eV in fluorides to 2.5 eV in selenides and appears to follow the nephelauxetic sequence. In contrast, the CT emission band does not show such a clear trend. The average CT emission energy is located at 2.5 eV and varies only by 1 eV when going from selenides to fluorides. Exceptions are YBO₃, YPO₄ and LuPO₄, where the CT emission band is located at around 4 eV. Fig. 5.5 shows the distribution of the Stokes shift between A-band absorption and emission of Bi³⁺. The average Stokes shift is about 0.7 eV. The Stokes shift varies between 0.1 and 2 eV and shows no trend with the U value.

Figure 5.4: The a) A- and C-band excitation energies and b) A-band and CT emission energies of the Bi³⁺ ion as function of the U value of the host compound.

Figure 5.5: Distribution of the Stokes shift in the A-band of Bi³⁺.

5.3.2. The vacuum referred binding energies in the 6s and 6p electronic states of ${\rm Bi}^{3+}$

Based on the available lanthanide spectroscopy combined with the chemical shift model, it is possible to determine the vacuum referred electron binding energy at the top of the valence band and the bottom of the conduction band in compounds. The energy of an electron at rest in the vacuum is here defined as energy zero. For 44 of the compounds in Table 5.1 this information was already gathered in Refs. [15, 26, 94–96]. Using the MMCT energy, one may determine the electron binding energy in the 6s² ground state with respect to that at the bottom of the conduction band as shown in Fig. 5.1. The A-band excitation energy was then used to determine the VRBE of the electron in the ³P₁ excited state. The vacuum referred binding energies of the electron in the 6s and 6p levels of the Bi³⁺ ion in 44 different compounds are shown in Fig. 5.7. E_X is the VRBE of the electron in the host exciton state. The binding energies in the 6s ground state tend to decrease with increasing U value and are always located above the valence band top. The binding energy in the 6p level also indicates a general decrease with increasing U value but appears to follow the location of the host conduction band.

Figure 5.6: The vacuum referred electron binding energy in the ${}^{1}S_{0}$ ground state and ${}^{3}P_{1}$ excited state of the Bi³⁺ ion as function of the U value.

Fig. 5.6 shows the VRBE in the 1S_0 ground and 3P_1 excited state of the Bi $^{3+}$ ion as function of the U value of the host compound. The 1S_0 ground state VRBE decreases from -5 eV to -10 eV with increasing U value, while the 3P_1 excited state VRBE decreases less strongly from -2 to -4 eV. The difference between both sets of data are then the A-band energies shown in Fig. 5.3. Both the 6s and 6p data points in Fig. 5.6 spread by ± 1 eV around the drawn lines.

Figure 5.7: Stacked VRBE diagrams for Bi^{3+} doped compounds. E_C is the binding energy at the conduction band bottom, E_X is the binding energy in the host exciton, E_V is the binding energy at the valence band top and 6s and 6p are the binding energies in the Bi^{3+} ground and excited state, respectively.

5.4. Discussion

To further verify the method of the chemical shift model as applied to Bi^{3+} in inorganic compounds, we collected information on the VRBE in the Bi^{3+} ground state from independent methods. From photo-electron spectroscopy studies on BiF_3 , Poole *et al.* [97] found the Bi^{3+} 6s² band at -9.6 eV and the fluorine valence band at -12.2 eV relative to the vacuum level. When we compare those numbers with those for Bi^{3+} as impurity in CaF_2 , SrF_2 , and BaF_2 in Fig. 5.7 we see almost the same number for $CaF_2:Bi^{3+}$. Note that Ca^{2+} and Bi^{3+} have a similar value for the ionic radius. Many studies have been performed on the electronic properties of $BiVO_4$ because of its photo-catalytic activity in water splitting. Bi^{3+} and La^{3+} also have similar ionic radius, and it is of interest to compare binding energy information from electro-chemistry on $BiVO_4$ with that from the chemical shift model on $LaVO_4:Bi^{3+}$ in Fig. 5.7. Kim *et al.* [98] and Hong *et al.* [99] located the top of the valence band at about 2.5 eV below the electrochemical potential of the normal hydrogen electrode. Relative to the vacuum level this translates to a VRBE of -6.94 eV. Note that for $LaVO_4:Bi^{3+}$ in Fig. 5.7, a value of -7.16 eV was found from the chemical shift method and the spectroscopy of Bi^{3+} which all agrees reasonably well.

Figure 5.8: The effect of the chemical environment on the vacuum referred binding energy of electrons in the 6s and 6p levels of Bi^{3+} and in the 4f and 5d levels of Ce^{3+} . E_{CS} shows the chemical shift of the electron due to the chemical environment in YPO₄.

When discussing the Bi^{3+} 6s and 6p VRBEs it is useful to compare those binding energies to the Ce^{3+} 4f and 5d VRBEs. When doped in a compound, the Bi^{3+} or Ce^{3+} cations will be screened by effectively an equal amount of negative charge from the surrounding anions in a way that minimizes the total energy. The Coulomb repulsion between that screening charge and the electrons in the Bi^{3+} or Ce^{3+} orbitals results in a decreased binding energy (less negative VRBE). The amount by which the binding energies shifts

upward is termed the chemical shift E_{CS} [14]. Bi^{3+} and Ce^{3+} have the same charge and almost identical effective ionic radii of 103 pm and 101 pm for a six-fold coordination, respectively [100]. From this, one may expect that the size of the chemical shift for both ions should be comparable. This is illustrated in Fig. 5.8 where the VRBEs in the ground and excited state for the free Ce^{3+} ion and for Ce^{3+} in YPO₄ are compared to those for Bi^{3+} . The -36.76 eV and -45.29 eV for the VRBE in the ground state of the free ions Ce^{3+} and Bi^{3+} are the same as the 4th ionization potentials of Ce and Bi. For Ce^{3+} the VRBE in the 5d state is 6.22 eV weaker and for Bi^{3+} the 6s6p state is 9.4 eV weaker bonded. The corresponding VRBE values in YPO₄ are also shown in Fig. 5.8, and for Bi^{3+} are also marked in Fig. 5.6. In YPO₄ the 6s electron binding energy for Bi^{3+} is located at -7.8 eV, corresponding with a chemical shift of 37.5 eV. For the Ce^{3+} ground state the chemical shift amounts 30.79 eV, which is significantly smaller than for Bi^{3+} with similar ionic radius.

Fig. 5.9 compares the VRBE in the 6s ground state of Bi^{3+} , the 4f ground state of Ce^{3+} and the excited state of both ions as function of U. The fitted line through the Bi^{3+} ground state and excited state data was taken from Fig. 5.6, whereas the Ce^{3+} and ground and excited state data are taken from Refs [14, 15]. By definition within the chemical shift model, the VRBE for Ce^{3+} changes linearly with U. Interestingly, the data of the Bi^{3+} ground state in Fig. 5.6 also appears to change linearly with U. In the free Bi^{3+} ion the electron in the ground state is 8.53 eV stronger bonded than in the free Ce ion (see Fig. 5.8). In fluorides (U = 7.4 eV) the difference has reduced to about 3 eV and it further reduces to about 1 eV for sulfides with (U = 6.2 eV). Yet, as found in this chapter, in compounds the Bi^{3+} ground state will always appear below that of Ce^{3+} .

Figure 5.9: The vacuum referred binding energy of electrons in the 6s ground state and 6p excited state of Bi^{3+} and the 4f ground state state and 5d excited state of Ce^{3+} .

Comparing the size of the chemical shift for compounds with U value typical for fluorides, typical for oxides, and typical for sulfides, the chemical shift for Bi^{3+} appears about 18 ±3% larger than that for Ce^{3+} , which leads to the steeper slope of the ground state VRBE in Fig. 5.9 as compared to that for Ce^{3+} . As such it is an interesting finding that the chemical shift of different ions are strongly correlated. One may speculate on why the size of the chemical shift for Bi^{3+} in compounds appears about 18% larger than for Ce^{3+} . The electron in the 4f-orbital of Ce^{3+} is relatively close (on average about 20-25 pm) to the nucleus and it is screened from the chemical environment by the 5s and 5p electrons. The electron in the 6s orbital of Bi^{3+} on the other hand extends to the periphery of the cation resulting in an overlap and hybridization with the anion orbitals. Possibly part of the explanation lies in those hybridization effects in the ground state.

Fig. 5.9 also compares the VRBE in the 5d excited state of Ce^{3+} with that in the 6s6p excited state of Bi^{3+} . The 5d electron VRBE does not depend on the U value and is on average at -1.8 eV with up to ± 1 eV compound to compound variation caused by changing crystal field splitting of the 5d-states [15]. For Bi^{3+} we do observe that the VRBE changes with the U value, yet less strongly as for the ground state. The Bi^{3+} excited state tends to be at lower energy than the Ce^{3+} 5d-state although, depending on the size of the crystal field splitting, in some compounds the situation can be reversed.

Y₂O₃

Figure 5.10: The vacuum referred binding energies of the divalent (red) and trivalent (blue) lanthanide 4felectrons and the 6s and 6p electrons of Bi^{3+} in Y_2O_3 .

In most of the compounds shown in Fig. 5.7, the VRBE of the 6s and 6p electrons of Bi^{3+} are located inside the band gap of the host compound. This type of electronic structure is optimal for luminescent applications. Both A-band excitation and emission can be observed in the luminescence spectra of these Bi^{3+} -doped compounds. The A-band excitation and emission energies increase linearly with increasing value of U, which allows engineering of the luminescence properties of materials with Bi^{3+} doping. The av-

erage Stokes shift in Fig. 5.5 is at 0.7 eV, while an average Stokes shift of 0.45 eV was reported for Ce^{3+} . The Stokes shift originates from the relaxation of the host lattice due to coupling of the excited electron orbital with lattice phonons and can therefore be regarded as a property of both the lattice and dopant. Apparently, the 6p electron orbital of Bi³⁺ has somewhat stronger interaction with the lattice than that of the 5d electron orbital of Ce³⁺.

Bi³⁺ is also reported to be a good sensitizer for lanthanide intra-4f luminescence [2– 6]. In Fig. 5.10 the VRBE in the 6s and 6p state of Bi^{3+} and the VRBEs in the 4f and 5d of the di- and trivalent lanthanides in Y_2O_3 are shown as an example. The Bi³⁺ Aband excitation and emission energy spectrally matches with the 4f transitions of the lanthanides. Furthermore, the VRBE of the Bi³⁺ 6p level is located above the VRBE of the Eu^{2+} 4f level, indicating that an electron transfer from the excited state of Bi^{3+} to Eu³⁺ is possible. In such case, both radiative and electron transfer from Bi³⁺ to Eu³⁺ can be used as sensitizing mechanism for Eu³⁺ luminescence. Only for transition metal based compounds in which the conduction band is located at or below -3 eV, the 6p excited state of Bi³⁺ is located inside the conduction band. This is for example the case in Bi^{3+} activated YVO₄ shown in Fig. 5.7. Since all Bi^{3+} interconfigurational transitions are located inside the conduction band, no A-band luminescence occurs. This is similar to the situation when the lowest 5d level of Ce^{3+} is located inside the conduction band. Upon excitation in the 5d state, the electron auto-ionizes and emission is not possible. Instead of A-band emission, in the case of Bi³⁺, excitation and emission of the MMCT transition can be observed.

The VRBE diagrams in Fig. 5.7 show that the Bi^{3+} 6s ground state is always located above the valence band. There appears to be a correlation of the electron binding energies in the 6s state and the valence band top. The chemical shift increases when the electronegativity of the anion decreases. As a result of this weaker bonding, the VRBE of both the anion electrons and Bi^{3+} become less negative. Since the Bi^{3+} ground state is always located above the valence band, Bi³⁺ can act as a trap for holes created in the valence band when exposed to ionizing radiation. Depending on the type of the host compound, Bi^{3+} can act as a deep hole trapping level, like in CaF_2 or NaCl (2.82 and 2.33 eV, respectively), or a shallow trap, like in the vanadates (0.6 \pm 0.15 eV). Ce³⁺ can only trap an electron to become Ce^{2+} when the CB bottom is above -1 eV, which is usually not the case. Whether Bi³⁺ can trap an electron to become Bi²⁺ depends on the VRBE in the lowest $6s^26p$ level of Bi^{2+} compared to that at the CB bottom. In the past 20 years, Bi²⁺ luminescence has been reported in wide band gap compounds such as sulfates, borates and phosphates [101–103], which is only possible when both the excited and ground state of Bi²⁺ are located below the bottom of the conduction band, and for those compounds Bi³⁺ would also act as an electron trap. Unfortunately, no information on a charge transfer transition between the host VB and Bi³⁺ was reported that could be used as a method to locate the Bi²⁺ ground state within the band gap. The fact that Ce³⁺ 5d-4f emission has never been reported in Bi³⁺ based compounds suggests that, upon excitation to the 5d-level, emission is fully quenched by electron transfer to the Bi. In other words it suggests that the VRBE in the Bi^{2+} ground state is below the VRBE in the Ce³⁺ 5d excited state. In chapter 6 the VRBE in the Bi²⁺ levels and its implications for the luminescence properties will be presented.

5.5. Summary and conclusions

Spectroscopic data on the metal-to-metal charge transfer and interconfigurational Aand C-band transitions of the Bi³⁺ ion in 117 different compounds were collected from the archival literature. Using the MMCT and A-band transition energies it was possible to determine the vacuum referred binding energy of the 6s and 6p electron in 44 different compounds. The obtained VRBE in the Bi³⁺ ground state are consistent with the VRBE obtained independently from photo-electron spectroscopy and electrochemical studies on BiF₃ and BiVO₄. The VRBE of the electron in the 6s and 6p levels were found to linearly decrease with the U value and follow the nephelauxetic sequence. The VRBE in the Bi³⁺ 6s ground state is always located above the valence band and below the VRBE of the Ce³⁺ 4f ground state. The chemical shift for Bi³⁺ was found 18 ±3% larger than that for Ce³⁺. A larger hybridization of the Bi³⁺ 6s orbital with the anion orbitals compared to that of the shielded Ce³⁺ 4f electrons was suggested as a possible origin.

5.6. Appendix A. The metal-to-metal charge transfer energies of Bi³⁺

Below an account is given on how the metal-to-metal charge transfer energies in Table 5.1 were obtained. The data is extracted from excitation or absorption spectra reported in the original literature. Since the assignment of the Bi³⁺ interconfigurational electronic transitions (A-, B-, and C-band) are rather straightforward, we limit ourselves to discussing only the MMCT transitions and refer to the original literature for additional information on the other Bi³⁺-related transitions.

Alkaline earth fluorides. Oboth *et al.* reported the optical absorption and emission of Bi^{3+} -doped CaF_2 , SrF_2 and BaF_2 [28]. An approximately 0.5 eV broad absorption band at 9.17 eV in the absorption spectrum of CaF_2 was observed, which they interpreted as the MMCT band. Similar assignments for the MMCT-band in Bi^{3+} -doped SrF_2 and BaF_2 were given.

Alkali halides. Nistor *et al.* observed an additional band on the high-energy side of the C-band in the optical absorption spectrum of KCl:Bi³⁺ [31]. Due to the very pronounced temperature dependence this band was attributed to a charge transition. We interpreted this charge transfer transition as the MMCT transition. Radhakrishna reported on the optical absorption of the Bi³⁺-doped alkali halides NaCl, KCl, RbCl, KBr and KI [30]. They also observed an additional band on the high-energy side of the C-band, which was labeled as C_3 . However, based on the report of Nistor, we propose that this C_3 band in the absorption spectra corresponds to the MMCT transition.

Oxyhalides. Blasse and Bril reported two absorption bands for the oxyhalides YOF, LaOCl and YOCl [1]. However, only excitation in the higher energy excitation band resulted in emission. Probably the emission under excitation in the lower energy A-band is completely quenched. Similar observations were made in the oxychlorides by Jacquier and by Wolfert and Blasse [36, 104]. We propose that the high-energy excitation is the MMCT transition. Wolfert and Blasse observed an excitation band at 4.3 eV in Bi³⁺-doped LaOBr, which they assigned to a charge transfer from the valence band to the Bi³⁺ ion [37]. However, we propose that this charge transfer transition is in fact the MMCT transition. **Phosphates**. Cavalli *et al.* observed a broad excitation band in the range of 6.53 - 8.27 eV

in Bi³⁺-doped YPO₄, which they assigned to the C-band [41]. However, this excitation appears to consist of two bands, namely the C- and MMCT-bands. We estimated the MMCT transition at the low energy side of the broad excitation band at approximately 7.3 eV. In the excitation spectrum of Bi³⁺-doped LaPO₄ reported by Moncorgé *et al.* an excitation band is visible at 7.1 eV, overlapping on the low energy side of the host exciton band at 7.75 eV [40]. They did not assign this excitation band to an optical transition, however, we propose that this excitation is the MMCT transition, in-line with the 7.3 eV MMCT transition of YPO₄:Bi³⁺. Wang *et al.* observed a broad excitation band between 6.2 and 6.9 eV in LaPO₄:Bi³⁺, which they assigned to the C-band. A sharp excitation peak at 10.5 eV was assigned to a CT transition, which would mean that the Bi³⁺ related charge transfer transition is larger than the 8.64 eV band gap of LaPO₄. Considering the broad shape of the 6.2-6.9 eV band, we propose that this excitation is due to the MMCT.

Borates. Chen *et al.* reported two emission bands at 4.22 and 3.87 eV in Bi^{3+} -doped YBO₃, originating from two different sites [48]. However, we propose that these are from isolated Bi^{3+} centers and emission from the charge transfer state, in agreement with the assignment of the emission bands in LaBO₃: Bi^{3+} by Wolfert *et al.* [47]. Furthermore, A-band excitation and emission was observed, originating from only one site in YBO₃: Bi^{3+} reported by Zhang *et al.* [105]. The broad shoulder on the the low energy side of the C-band, centered at 6.2 eV in the excitation spectrum reported by Chen *et al.*, could then be assigned to the MMCT transition.

Germanates. Katayama *et al.* reported a broad MMCT transition at 4.9 eV in Bi³⁺-doped MgGeO₃ [54].

Aluminates. Krasnikov *et al.* reported a broad excitation band located at E > 6 eV in YAlO₃, which they suggested to be from the C-band transition [69]. However, we propose that this band is from the MMCT transition and estimated the center of this band at approximately 6.5 eV. Zorenko *et al.* reported charge transfer transitions in Bi³⁺-doped Y₃Al₅O₁₂ at 5.25, 5.93 and 6.85 eV, of which they propose that the 5.93 eV transition originates from the Bi³⁺ \rightarrow Bi⁴⁺ + e⁻ charge transfer transition [68]. The band excitation band at 5.95 eV observed in Bi³⁺-doped Lu₃Al₅O₁₂, originally assigned to the C-band by Babin *et al.*, we also propose to be the MMCT transition [71].

Gallates. Jacquier *et al.* observed an approximately 0.8 eV wide excitation band at 5.17 eV in the excitation spectrum of $LaGaO_3:Bi^{3+}$ [61]. Srivastava also observed a rising edge of an excitation band at approximately 4.9 eV, but due to the limitations of their setup they were not able to determine the center of this band [106]. According to Boutin-aud, this energy would match with the energy expected for the MMCT transition [21]. Zhuang observed an excitation band at 4.43 eV in $ZnGa_2O_4:Bi^{3+}$, which they attributed to a transition of exciton-like origin ($Bi^{4+} + e^-$) [59]. However, we propose that this is the MMCT transition. Setlur and Srivastava showed an excitation band at 4.36 eV for Bi^{3+} -doped $Y_3Ga_5O_{12}$ [23]. Ilmer *et al.* reported two bands at 4.43 and 4.07 eV for Bi^{3+} -doped $Y_3Ga_5O_{12}$ [64]. They mentioned that the ³P₁ level is probably very close to the conduction band and that the highest excitation band can be considered as an impurity bound exciton in which the hole resides on the Bi^{3+} ion and the electron is inside the conduction band. We therefore assigned the 4.43 eV excitation to the MMCT transition. Ilmer also observed two excitation bands at 4.59 and 4.0 eV in $Gd_3Ga_5O_{12}$, with identical assignments as in $Y_3Ga_5O_{12}$.
Transition metal compounds. Compounds containing transition metal cations generally have a conduction band located at or below -3 eV relative to the vacuum level. In those compounds it is to be expected that the $Bi^{3+} P_1$ excited state is located inside the conduction band of the host compound. The only Bi^{3+} related excitation band in these type of compounds that is expected to be observed is the MMCT transition. The exception are the tantalates, which have a conduction band at energies slightly higher than -3 eV, in which the A-band can also be observed. Boutinaud and Cavalli systematically investigated the MMCT transition of Bi^{3+} in a variety of closed-shell transition metal oxides [20]. In most cases the MMCT transition is observed as a shoulder on the low energy side of the host exciton band. With this knowledge, we have extracted additional MMCT transitions from the literature on Bi^{3+} -doped transition metal oxides (see references in Table 5.1).

Zirconates. Srivastava and Beers reported two excitation bands in Bi^{3+} -doped $La_2Zr_2O_7$ [88]. The low-energy excitation band at 4.27 eV was assigned to the A-band, whereas the high-energy excitation band at 5.06 eV was tentatively assigned to the absorption into the excitonic state. However, we know the host lattice exciton energy is located at around 6 eV. Since the MMCT is also exciton-like, we propose that the 5.06 eV excitation band is due to the MMCT transition.

Simple oxides. Ellervee reported a 0.5 eV wide excitation band centered at 5.1 eV and 4.7 eV in the excitation spectra of Bi³⁺-doped CaO and SrO, respectively [85]. Yamashita and Asano reported this excitation band at 5.28 eV in CaO and van der Steen and Dijcks at approximately 4.84 and 4.71 eV for CaO and SrO, respectively [27, 107]. Similar to the assignment of the MMCT transition in the sulfides and selenides, we propose that these excitation bands correspond to the MMCT transitions. We have taken the values reported by Ellervee for the MMCT transition listed in Table 5.1 of this chapter. Boulon reported on the luminescence of Bi^{3+} in Ln_2O_3 with Ln = Sc, Lu, Y, Gd and La [74]. In the excitation spectra of these rare-earth oxides doped with Bi³⁺, a high-energy band at 4.6-5.0 eV was observed depending on the type of rare-earth. Blasse also reported a high energy excitation band at 4.71 eV for Y_2O_3 :Bi³⁺ and Bordun observed these excitation bands in $Bi^{3+} Y_2O_3$ and Sc_2O_3 ceramics [1, 90]. Although the original literature assigns these bands to the C-band, we agree with the assignment of Boutinaud, whom suggested that these excitation bands should be assigned to the MMCT transition [21]. A charge transfer transition was also suggested by van de Craats and Blasse to account for the luminescence quenching in Y_2O_3 :Bi³⁺ [108].

Other oxides. Chang reported a 0.5 eV wide excitation band centered at 3.17 eV, which was concentration dependent, becoming broader and intenser with increasing Bi^{3+} concentration in $BaLa_2ZnO_5$ [87]. This excitation band was assigned to the A-band, however, we propose that this should be assigned to the MMCT transition especially since the observed decay time was very slow (milliseconds), whereas typically the Bi^{3+} A-band is in the order of nanoseconds to microseconds.

Sulfides and selenides. Yamashita and Asano investigated the luminescence of Bi^{3+} in Ca(S : Se) [27]. For CaS: Bi^{3+} they observed excitation bands at 4.5, 4.2, 3.54 and 3.02 eV which were assigned to the D-band, C-band, B-band and A-band, respectively. The D excitation band was attributed to the paired ion center, however, this should be the host exciton band. The assigned B-band is as intense as the observed the A-band. Due

to the forbidden nature of the B-band transition, this is more likely the allowed C-band transition. The 4.2 eV excitation band which was assigned to the C-band is then more likely the MMCT transition, especially regarding the temperature dependence and the fact this excitation band is twice as broad as the A-band. For CaSe:Bi³⁺ excitation bands were observed at 3.85, 3.6, 3.2 and 2.74 eV, which were assigned to the D-band, C-band, B-band and A-band, respectively. Again we propose that the D-band to be the exciton band and the broad 3.6 eV excitation band to be the MMCT transition, considering the high temperature dependency of the transition. The 3.2 eV excitation assigned to the forbidden B-band transition is more likely the C-band. Yamashita *et al.* showed three bands in the excitation spectrum of SrS:Bi³⁺ at 4.65, 3.75 and 2.95 eV, which were assigned to exciton, C-band and A-band, respectively [92]. However, due the strong temperature dependence of the 3.75 eV excitation band and the fact that this excitation band is rather broad (0.5 eV) we suggest that this band should be assigned to the MMCT transition. For MgS, Asano and Yamashita reported excitation bands at 4.4, 3.83 and 2.98 eV, which they assigned to the charge transifer transition, the C-band and the A-band, respectively [93].

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6

Towards a general concentration quenching model of Bi³⁺ luminescence

The vacuum referred binding energy of the electron in the Bi^{2+} ground state has been determined in 15 different compounds. This shows that the electron binding energy in the ground state of Bi^{2+} is at lower (more negative) energy compared to the electron binding energy in the excited state of Bi^{3+} . This means that electron transfer from the excited state of one Bi^{3+} to a neighboring Bi^{3+} , forming a $Bi^{2+}-Bi^{4+}$ pair, acts as a quenching route for the Bi^{3+} emission. Electron back transfer in the $Bi^{2+}-Bi^{4+}$ pair is then suggested to be the origin for the frequently observed pair emission. This paper shows that vacuum referred electron binding energy diagrams can provide a unique physical insight in the properties of inorganic compounds.

This chapter is based on the publication: **R. H. P. Awater** and P. Dorenbos, "Towards a general concentration quenching model of Bi³⁺ luminescence", *Journal of Luminescence* **188** (2017), 487.

6.1. Introduction

Bi³⁺ shows a complex temperature and concentration dependent luminescence quenching [1–3]. The luminescence quenching cannot be explained by multi-phonon relaxation because of a too large energy gap between the ${}^{3}P_{1}$ excited state and the ${}^{1}S_{0}$ ground state. A transfer of an electron from the excited state to the conduction band is often used to describe the quenching, like for example in (In,Lu)BO₃ and Ca(Zr,Sn)O₃ compounds [4, 5]. Another possibility for luminescence quenching is via the charge transfer (CT) of an electron in the valence band to a bismuth ion as has been proposed for example in LaOBr [6]. These two quenching routes involving CT apply to isolated bismuth centers. A third type of quenching occurs in Bi-pairs, which becomes important at high concentrations of bismuth or in the case of preferential formation of bismuth pairs or clusters [7–9]. The mechanism of such type of quenching has always remained open.



Figure 6.1: Stacked VRBE diagrams for 10 different Bi^{3+} doped compounds with E_C the binding energy at the conduction band bottom, E_V the binding energy at the valence band top and E_X the binding energy in the exciton state. The blue bars labeled ${}^{1}S_0$ and ${}^{3}P_1$ indicate the electron binding energy in the Bi^{3+} ground and excited state, respectively.

In chapter 5 we determined the vacuum referred electron binding energies (VRBE) in the Bi³⁺ [¹S₀] and [³P₁] levels in 44 different compounds. The VRBE of Bi³⁺ in a selection of 10 of those compounds is shown in Fig. 6.1. The VRBE in the ¹S₀ ground state varies between -5 and -10 eV and in the ³P₁ excited state between -2 and -4 eV in a systematic fashion with the chemical and structural properties of the compound. When Bi³⁺ is doped in compounds with a low lying conduction band, e.g. compounds containing transition metals with d⁰ configuration such as YNbO₄ and YVO₄ in Fig. 6.1, the ³P₁ excited state appears above the conduction band (CB) bottom. In these cases photoionization to the CB occurs without generating any Bi³⁺ emission. In stead a MMCT emission is often observed in these compounds [10, 11]. Besides Bi³⁺, also luminescence from Bi²⁺ has been reported in fluorides, phosphates, sulphates and borates [12–15]. For

 $Li_2BaP_2O_7$ in chapter 3 and YPO₄ in chapter 4, we concluded that the VRBE in the Bi²⁺ ${}^2P_{1/2}$ ground state is at approximately –4 eV, which is below the 3P_1 excited state of Bi³⁺ in those compounds. This means that electron transfer from the excited state of one Bi³⁺ ion to a neighboring bismuth ion is energetically favorable. Such electron transfer provides then a mechanism for the quenching of the Bi³⁺ emission.

This mechanism is illustrated in Fig. 6.2. Two Bi^{3+} ions labeled I and II that form a Bi–Bi pair are shown, and after exciting an electron to the ${}^{3}P_{1}$ excited state of bismuth ion I (arrow 1), energy will be released in the electron transfer to the neighboring bismuth ion II (arrow 2), effectively forming a Bi^{2+} – Bi^{4+} pair. Recombination of the Bi^{2+} – Bi^{4+} electron-hole pair (arrow 3) can be radiately or non-radiatively. If the recombination occurs radiatively, a broad emission band with a large Stokes shift typical for CT transitions can be observed. This also provides a quenching route for the $Bi^{3+} {}^{3}P_{1} \rightarrow {}^{1}S_{0}$ emission. In this paper we will demonstrate that electron transfer between neighboring bismuth ions is a general phenomenon for Bi^{3+} -doped compounds when either Bi enters in pairs or cluster or when the Bi concentration is sufficiently high to generate pairs based on statistical grounds. For that we have collected information on Bi^{2+} in the 15 compounds listed in Table 6.1.



Figure 6.2: The proposed quenching model of the Bi^{3+} luminescence and generation of pair emission via electron transfer between neighboring bismuth ions.

6.2. Results and Discussion

For almost all compounds, the listed VRBE at the valence band (VB) top and conduction band (CB) bottom were presented earlier, and together with the VRBE in the Bi³⁺ ground and excited states are taken from chapter 5. There are several methods to estimate the VRBE in the Bi²⁺ ground and excited state. One may seek for information on the VB \rightarrow Bi³⁺ charge transfer. However, we only found a report on MgGeO₃ by Katayama *et al.* from which a Bi²⁺ location at –3.8 eV is derived [16]. Another method is to determine the energy barrier ΔE_q for thermal quenching of Bi²⁺ emission. However, detailed studies on the quenching of Bi²⁺ emission to determine such energy barrier are not available. At best one may find the temperature T_{0.9}, the temperature where the intensity has dropped by 10% and the quenching starts to become significant. Using methods similar as described in Ref. [17], the quenching energy barrier ΔE_q is approximately T_{0.9}/620 eV. The electron does not need to be transferred as high in energy as the conduction band bottom for quenching to occur. We will assume that $\frac{E_C + E_X}{2} - \Delta E_q - E_{em}^{Bi^{2+}}$ provides a fair estimate for the VRBE in the Bi²⁺ ground state ($E_{g.s.}^{Bi^{2+}}$), with E_C the VRBE of the electron at the conduction band bottom and E_X the VRBE of the electron in the exciton state. The relevant values, including the references to the literature we used for $E_{em}^{Bi^{2+}}$, the Bi²⁺ emission energy, can be found in Table 6.1.

Table 6.1: Spectroscopic and VRBE data on Bi²⁺ in compounds and on pure Bi compounds. All energies are in eV.

Host compound	E_V	E_C	$E_{em}^{\mathrm{Bi}^{2+}}$	T _{0.9}	$E_{\rm g.s.}^{{\rm Bi}^{2+}}$	Ref.
BiF3	-9.60	-4.42	-	-	-4.42	-
SrF ₂	-12.13	-0.69	2.07	> 300 K	-3.66	[12]
CaF ₂	-12.33	-0.34	2.25	> 300 K	-3.52	[12]
BaSO ₄	-8.93	-0.12	1.98	> 300 K	-2.91	[13]
SrB ₄ O ₇	-9.59	-0.79	2.1	> 300 K	-3.70	[15]
SrSO ₄	-8.93	0.12	2.03	> 300 K	-2.73	[14]
BaBPO ₅	-8.98	-0.29	1.94	> 300 K	-3.03	[18]
SrBPO ₅	-9.08	-0.21	1.94	> 300 K	-2.96	[18]
CaBPO ₅	-9.24	-0.01	1.97	> 300 K	-2.80	[18]
CaSO ₄	-8.94	0.19	2.1	> 300 K	-2.73	[13, 14]
Li ₂ BaP ₂ O ₇	-9.23	-0.86	1.81	> 600 K	-3.95	[19]
BiPO4	-8.73	-3.83	-	_	-3.83	_
YPO ₄	-9.77	-0.53	1.85	350 K	-3.28	[20]
MgGeO ₃	-8.70	-2.22	n.a.	n.a.	-3.81	[16]
Bi ₂ O ₃	-7.53	-4.18	-	_	-4.18	-

A final method to obtain indications on the VRBE in the Bi-levels is from pure Bicompounds. Usable information was found for BiF₃, BiPO₄, and Bi₂O₃. The top of the VB has here a strong Bi³⁺ ground state component and the CB-bottom a strong Bi²⁺ contribution. For BiF₃ the VRBE at the VB-top is found at –9.6 eV and is derived from the X-ray photoelectron spectroscopy studies by Poole *et al.* [21], and by adding information on the band gap the CB-bottom is found at –4.4 eV. BiPO₄ has been studied for its photocatalytic activity and from those studies the CB-bottom was found about 0.4 eV above that of TiO₂ [22] which has the CB-bottom at –4.2 eV [23]. Bi₂O₃ has also been studied for its photocatalytic activity and Lin *et al.* [24] places the CB-bottom at 0.1 eV below the H⁺/H₂ redox potential which translates to a VRBE of –4.5 eV. Combined with band gap information, the data as compiled in Table 6.1 is obtained. The data from Table 6.1 is presented as a stacked VRBE diagram in Fig. 6.3. The Bi^{3+} ground and excited states are indicated by the blue horizontal bars within the forbidden gap. The VRBE in the Bi^{2+} ground state (g.s.) and excited states are illustrated by the horizontal red bars. In the case when only an upper limit VRBE estimation is available, a small down pointing arrow has been added. The true position, depending on the actual emission quenching temperature, can well be 0.5 eV lower. For the pure Bi-based compounds the top of the valence band is regarded as representative for the VRBE in the Bi^{3+} g.s. and the bottom of the CB for the VRBE in the Bi^{2+} g.s. For example, the electron binding energy in the ground state of Bi^{2+} in YPO₄ is located at -3.28 eV, which is only 0.5 eV different from the binding energy in the conduction band bottom of $BiPO_4$. Also the electron binding energy in Bi^{3+} ground state located at -8.2 eV in YPO₄ differs only by 0.5 eV from the binding energy in the valence band top of $BiPO_4$. The different crystal structures, but also orbital mixing of Bi-states with that of other host atoms may account for such 0.5 eV difference.



Figure 6.3: Stacked diagram with the VRBE at the VB-top and the CB-bottom of various compounds with the Bi^{3+} (blue horizontal bars) and Bi^{2+} (red horizontal bars) levels. Also the VRBE at the CB-bottom and VB-top in BiF_3 , $BiPO_4$, and Bi_2O_3 is shown.

The newly added information in this chapter is the VRBE in the Bi²⁺ ground state, which appears in the range between –3.5 and –4 eV, and is indeed below the Bi³⁺ excited state usually found near –3 eV in fluoride and oxide compounds. The energy difference between the VRBE in the Bi³⁺ excited state and Bi²⁺ ground state appears to be about 0.7 \pm 0.3 eV. If approximately 1 eV lattice relaxation loss is taken into account, the remaining energy for possible pair emission is then expected 1.7 \pm 0.5 eV lower than the Bi³⁺ emission energy which is indeed the typical Stokes shift observed. For example, in YPO₄:Bi³⁺ the pair emission band at 325 nm is 1.6 eV Stokes shifted from the Bi³⁺ excitation.

Depending on the transition rates, complicated dynamics as in the Bi³⁺-doped garnet systems may now occur [25–27]. It is not the purpose of this work to enter in too much detail but some general ideas can be launched. The typical luminescence decay rate at room temperature of the Bi³⁺ emission is found near 1 μ s and the luminescence decay of the pair emission is typically 100 μ s [27, 28]. The radiative decay rate of the Bi³⁺ emission (arrow 1 in Fig. 6.2) has to compete with the electron transfer rate to the neighboring Bi ion (arrow 2 in Fig. 6.2). In such electron transfer, the above found 0.7 ± 0.3 eV energy difference needs to be emitted as phonons and a temperature dependent rate is to be expected. Once in the Bi⁴⁺–Bi²⁺ pair state, again different rates apply. With only 0.7 eV difference, thermally activated electron back transfer to the higher energy Bi³⁺[³P₁]–Bi³⁺[¹S₀] pair state is possible (reverse of arrow 2 in Fig. 6.2). In addition, there are the luminescence decay rates of Bi-pair emission, and its thermal quenching rate.

Srivastava *et al.* [2] studied the luminescence intensity as function of temperature of the Bi³⁺ emission at 385 nm (3.22 eV) and observed an additional emission band labeled as VIS-band at 514 nm (2.41 eV) when exciting Bi³⁺ at 290 nm (4.27 eV) in 0.5% Bi³⁺ doped La₂Zr₂O₇. From 0 K to 60 K the Bi³⁺ emission quenches whereas the VIS-band grows in intensity, and from 70 K to 120 K the process is reversed and the Bi³⁺ emission grows again with decreasing VIS-band intensity. Above 120 K both bands quench until at RT no emission remains. Although the origin of the VIS-band was not clear, one may also interpret it as Bi-pair emission. Intensity first grows at the expense of Bi³⁺ emission, above 70 K electron back transfer to the Bi³⁺ excited state reverses the process, and above 120 K the radiation-less electron transfer to the Bi³⁺ ground state starts to quench both emissions. Energetically such processes are well-possible. More detailed studies on the luminescence dynamics as function of temperature and Bi concentration would be required to verify or falsify such dynamics.

6.3. Conclusion

We have shown that in general the VRBE in the Bi^{2+} ground state is located at lower energy as compared to the VRBE in the $Bi^{3+3}P_1$ excited state. In the presence of Bi–Bi pairs, it is then upon excitation of one Bi^{3+} energetically favorable for electron transfer to occur, and to form a $Bi^{2+}-Bi^{4+}$ pair. Based on the estimated VRBEs in the Bi^{2+} ground state in 15 different compounds, the electron transfer between neighboring Bi ions appears always energetically possible. The recombination of the electron and hole on the $Bi^{2+}-Bi^{4+}$ pair can occur non-radiatively or radiatively, in the later case the resulting visible emission band is often observed and ascribed to pair emission. The proposed mechanism of charge transfer between Bi-pairs can also explain the quenching behavior of Bi^{3+} luminescence when the Bi^{3+} concentration increases.

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7

The vacuum referred electron binding energies in the 1S_0 and 3P_1 states of Pb²⁺ and Tl⁺ in inorganic compounds

An overview of the spectroscopic data on Tl^+ and Pb^{2+} in 37 and 126 different inorganic compounds, respectively, is presented. Using the metal-to-metal charge transfer and A-band transition energies, the electron binding energies in the ${}^{1}S_{0}$ ground state and ${}^{3}P_{1}$ excited state of Tl^+ and Pb^{2+} were determined relative to the vacuum level. By constructing vacuum referred binding energy (VRBE) diagrams that display the energy levels of the Tl^+ and Pb^{2+} activator ion together with that of the host compound, insight in the luminescent properties of these activated compounds are gained. The obtained VRBEs of the electron in the ${}^{1}S_{0}$ and ${}^{3}P_{1}$ states of Tl^+ and Pb^{2+} are compared to those of Bi^{3+} .

This chapter is based on the publication that has been submitted to Journal of Luminescence: **R. H. P. Awater** and P. Dorenbos, "The vacuum referred electron binding energies in the ${}^{1}S_{0}$ and ${}^{3}P_{1}$ states of Pb²⁺ and Tl⁺ in inorganic compounds".

7.1. Introduction

The continuous research on Ce^{3+} -doped halides for scintillation materials led to the discovery of $LaBr_3$:Ce,Sr. It has an energy resolution of 2% for the detection of 662 keV γ -photons and a light yield of 70.000 photons per MeV of ionization energy [1]. Despite these excellent scintillation properties, a major drawback for γ -ray detection is their low density (~5 g/cm³). Oxides containing high Z elements such as Hf, Ta, W, Tl, Pb or Bi are attractive candidates for high density (~8 g/cm³) scintillator materials. In order to evaluate if Ce³⁺ can luminescence in these type of host compounds, knowledge on the activator levels relative to the host conduction and valence bands is required. By means of the chemical shift model, it is possible to determine the 4f and 5d electron binding energies are graphically presented in vacuum referred binding energy (VRBE) diagrams, which provide valuable insight on the performance of luminescent materials.

In 2014 the VRBEs of the 5d electrons in Hf⁴⁺, Ta⁵⁺ and W⁶⁺ ions were determined [3, 4]. In chapter 5 we have determined the VRBEs in the Bi³⁺ 6s [¹S₀] ground state and 6p [³P₁] excited state in 44 different inorganic compounds. This showed that the VRBE in the ¹S₀ ground state of Bi³⁺ decreases from –5 eV to –10 eV when changing the chemical environment (from sulfides to fluorides), while the ³P₁ excited state decreases from –2 eV to –4 eV. In this chapter, we determined the electron binding energies in the ¹S₀ and ³P₁ states of Tl⁺ and Pb²⁺. Like the Bi³⁺ ion, Tl⁺ and Pb²⁺ ions have a 6s² outer electron configuration.

In order to determine the absolute electron binding energy in an energy level of an activator ion, knowledge on the energy of a charge carrier (electron or hole) transition either from the valence band top to the activator ion or from the activator ion to the conduction band bottom is necessary. In the absorption spectra of $6s^2$ activated compounds there is, besides the interconfigurational A-, B- and C-bands, often an additional band visible which is traditionally labeled as D-band [5, 6]. This D-band can be interpreted as a metal-to-metal charge transfer (MMCT) from the 1S_0 ground state to the conduction band bottom [7]. Boutinaud reported on the systematics of the MMCT of Pb²⁺ and Bi³⁺ in closed shell transition metal oxides and provided a model to predict the energy of the MMCT transition [8, 9].

Together with the D-band absorption, also an additional emission is often observed and labeled as D-band emission [10–12]. The D-band emission can be interpreted as the radiative recombination of the electron and hole formed in the MMCT. A charge carrier transfer between dimers or pairs of $6s^2$ ions is also possible [13, 14]. In chapter 6 we have shown by means of VRBE diagrams that $(Bi^{3+})^* \rightarrow Bi^{3+}$ electron transfer to form a Bi^{2+} – Bi^{4+} pairs is energetically favorable and can quench the Bi^{3+} luminescence. Considering the large variety of optical and CT transitions possible in the spectra of $6s^2$ ion activated compounds, it can be difficult to assign and interpret the spectral bands and luminescence properties of these compounds. By locating the electronic levels of Tl^+ and Pb^{2+} , we attempt to provide the general trends of the luminescent properties of Tl^+ and Pb^{2+}

7.2. Methodology

Spectroscopic data on the interconfigurational and charge transfer transitions of Pb²⁺ in 126 and of Tl⁺ in 37 different inorganic compounds have been collected from the literature and are presented in Tables 7.1 and 7.2. We have used the same methodology as in chapter 5 on the electron binding energies in Bi^{3+} , which can be consulted for additional information. The compound identification number and compound name are listed in the first and second column, respectively. The third column lists the U value of the host compound, which was taken from previous reports [3, 15] and otherwise it was estimated based on the U value of compounds with similar chemical composition. These estimated values are shown in italics. The excitation energies of the A-, Cand MMCT-bands are shown in columns 4, 5 and 6, respectively. The emission energies corresponding to the A-band and CT luminescence are shown in columns 7 and 8. In most of the compounds listed in Tables 7.1 and 7.2 we used the original spectroscopic assignments. However, in some cases we used an alternative interpretation of the spectroscopic data which is motivated per compound in Appendices A and B. The reference to the original literature, which can be consulted for additional information on the spectroscopic bands, is shown in the last column.

ID		U	E	xcitation	[eV]	Emiss	Emission [eV]	
number	Compound	[eV]	А	С	MMCT	А	СТ	Ref.
0000000	free	_	7.98	11.82	-	7.49	-	[16]
1190004	NaF	7.50	5.75	-	-	_	4.51	[17]
1190010	BaF ₂	7.38	6.12	7.95	9.15	6.57	4.85	[18]
1190015	LiBaF ₃	7.42	6.34	-	-	5.01	4.02	[19]
1190020	SrF ₂	7.32	6.10	8.04	9.40	7.00	5.65	[18]
1190030	CaF_2	7.31	6.08	8.05	9.70	7.05	5.70	[18]
1190035	LiCaAlF ₆	7.65	6.08	-	-	5.93	-	[20]
1190040	MgF ₂	7.32	6.80	8.29	10.80	6.57	5.16	[18]
1190043	KMgF ₃	7.66	6.63	8.98	9.76	5.71	-	[21]
1190100	LaF ₃	7.51	6.20	7.80	-	4.43	-	[21]
1190300	GdF ₃	7.56	6.02	7.61	8.75	-	-	[21]
1190400	YF ₃	7.51	5.90	7.65	8.85	4.31		[21]
1190410	BaY ₂ F ₈	7.60	6.63	8.38	8.86	5.44	-	[21]
2190010	BaFCl	7.15	5.83	-	-	4.96	2.79	[12]
2190020	SrFCl	7.10	5.65	-	-	5.28	3.10	[12]
2190030	CaFCl	7.10	5.52	-	-	5.28	3.44	[12]
2290000	NH ₄ Cl	7.05	4.73	-	-	3.62	2.85	[22]
2290001	CsCl	6.80	4.46	6.20	-	-	-	[23]
2290002	RbCl	6.80	4.56	6.26	6.45	3.45	-	[24, 25]
2290003	KCl	6.80	4.57	6.23	6.53	3.50	3.00	[26]
2290004	NaCl	6.70	4.55	6.25	6.65	_	-	[26]
2290030	CdCl ₂	6.70	4.35	-	5.16	3.87	2.48	[27]
3190010	BaFBr	7.00	5.26	_	-	4.28	2.38	[12]

Table 7.1: The spectroscopic data of Pb²⁺ in 126 different compounds.

ID		U	E	Excitation [eV]		Emis	sion [eV]	
number	Compound	[eV]	Α	С	MMCT	А	СТ	Ref.
3190020	SrFBr	7.00	5.12	-	-	4.13	2.82	[12]
3390001	CsBr	6.60	4.00	5.64	-	_	-	[23]
3390002	RbBr	6.60	4.16	5.50	6.10	3.20	-	[28, 29]
3390003	KBr	6.60	4.16	5.54	6.20	3.44	-	[26, 29]
3390004	NaBr	6.55	4.08	5.64	-	-	-	[24]
3390031	CsCdBr ₃	6.90	3.72	-	-	2.45	-	[30]
4490001	CsI	6.25	3.32	4.79	5.77	_	-	[23]
4490002	RbI	6.25	3.54	4.50	5.42	_	-	[28]
4490003	KI	6.25	3.54	4.53	5.45	_	-	[28]
4490004	NaI	6.25	3.47	3.90	4.68	3.04	2.06	[31]
4490020	SrI ₂	6.25	2.98	3.25	4.38	2.53	1.82	[32]
5234030	$Ca_{10}(PO_4)_6Cl_2$	7.10	5.14	-	-	_	2.88	[33]
5251020	SrB ₅ O ₉ Cl	7.20	5.00	-	5.58	4.32	3.03	[34]
5251030	CaB ₅ O ₉ Cl	7.20	5.00	-	-	4.31	-	[35]
5290020	Sr ₄ OCl ₆	6.70	4.07	-	-	_	2.88	[36]
5351020	SrB ₅ O ₉ Br	7.00	4.88	-	5.68	4.46	3.35	[34]
5390020	Sr ₄ OBr ₆	6.50	3.81	-	-	_	2.70	[36]
5514010	BaSO ₄	7.26	5.64	-	-	_	3.65	[37]
5514020	SrSO ₄	7.26	5.64	-	-	4.35	3.26	[37]
5514030	CaSO ₄	7.19	5.64	7.75	7.29	5.28	-	[37]
5524010	BaCO ₃	7.20	4.77	-	5.77	_	3.54	[38]
5524020	SrCO ₃	7.20	5.39	-	-	4.35	3.44	[38]
5524030	CaCO ₃	7.16	5.12	-	5.95	4.00	-	[39]
5534010	$Ba_4(PO_4)_2SO_4$	7.00	4.96	-	-	_	3.35	[40]
5534020	$Sr_4(PO_4)_2SO_4$	7.00	4.86	-	-	_	3.59	[40]
5534025	LiSrPO ₄	7.10	5.02	-	-	_	-	[41]
5534030	$Ca_{10}(PO_4)_6(OH)_2$	7.10	5.08	-	-	_	2.88	[42]
5551020	SrB ₆ O ₁₀	7.31	4.51	-	-	4.00	-	[43]
5551020	SrB_4O_7	7.26	4.59	-	-	4.04	-	[44]
5552010	BaB_2O_4	6.80	5.17	-	-	3.35	-	[45]
5552015	$LiBa_2B_5O_{10}$	7.10	5.06	-	-	_	2.88	[46]
5552020	SrB ₂ O ₄	6.76	4.49	-	-	3.40	-	[44]
5552030	CaB_2O_4	6.75	4.88	-	-	3.82	-	[35]
5553030	$Ca_2 \tilde{B}_2 O_5$	6.75	4.86	5.77	-	3.65	-	[47]
5554010	$Ba_2Mg(BO_3)_2$	6.98	4.23	-	-	3.25	-	[48]
5554020	$Sr_2Mg(BO_3)_2$	6.97	4.77	-	-	3.76	-	[48]
5554024	$NaSr_4(BO_3)_3$	6.90	4.26	5.28	-	3.37	-	[49]
5554025	$LiSr_4(BO_3)_3$	6.90	4.37	5.39	-	3.78	-	[50]
5554030	$CaZr(BO_3)_2$	7.00	4.96	_	-	_	2.99	[51]
5554030	$Ca_3(BO_3)_2$	7.00	4.88	5.90	-	3.66	2.88	[47]
5554035	Li ₄ CaB ₂ O ₆	6.75	4.71	_	-	4.16	-	[52]
5554035	LiĊaBO₃	6.90	4.68	_	-	4.19	-	[53]

Table 7.1: Continued

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Table 7.1: 0	Continued
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ID		U	Excitation [eV]		Emission [eV]			
number	Compound	[eV]	А	С	MMCT	А	СТ	Ref.
5554120	La ₂ Sr ₅ Mg(BO ₃) ₆	7.00	4.88	-	-	-	3.40	[54]
5554420	Sr ₆ YAl(BO ₃) ₆	6.90	4.48	-	-	-	3.34	[54]
5555010	Ba ₂ Be ₂ B ₂ O ₇	6.80	4.37	-	-	3.13	-	[55]
5555010	$BaAl_2B_2O_7$	6.85	4.66	-	-	_	2.93	[56]
5555020	SrAl ₂ B ₂ O ₇	6.85	4.48	-	-	-	2.95	[57]
5555030	$CaAl_2B_2O_7$	6.85	4.73	-	-	3.52	-	[57]
5555120	SrLaBO ₄	6.85	4.68	-	-	_	2.70	[58]
5555130	$Ca_4 LaO(BO_3)_3$	6.85	4.77	-	-	3.35	3.02	[47]
5560010	BaSi ₂ O ₅	6.80	5.06	-	-	_	3.54	[59]
5562020	SrSiO ₃	6.80	5.30	-	-	3.44	2.79	[60]
5562030	CaSiO ₃	6.70	4.82	-	-	3.58	-	[60]
5564020	BaSrSiO ₄	6.60	4.96	-	-	_	3.10	[59]
5564020	Sr ₂ SiO ₄	6.59	5.06	-	-	_	3.10	[59]
5564025	Li_{4} SrCa(SiO ₄) ₂	6.80	4.98	-	-	4.26	-	[61]
5564025	Li ₂ SrSiO ₄	6.91	4.58	-	-	3.25	-	[62]
5564040	CaMgSiO ₄	6.80	4.96	-	-	_	3.44	[59]
5565400	Y ₂ SiO ₅	6.80	4.95	-	5.25	3.67	2.70	[63]
5565420	$\tilde{Sr}_{2}Y_{8}(SiO_{4})_{6}O_{2}$	6.80	4.73	-	-	3.33	-	[64]
5565420	$Sr_4Y_6(SiO_4)_6O$	6.80	4.77	-	-	3.35	-	[64]
5565430	$Ca_2Y_8(SiO_4)_6O_2$	6.80	4.73	-	-	3.35	-	[64]
5565430	$Ca_4Y_6(SiO_4)_6O$	6.80	4.73	-	-	3.35	-	[65]
5565440	$Mg_4Y_6(SiO_4)_6O$	6.80	4.66	-	-	3.26	-	[64]
5565440	$Mg_2Y_8(SiO_4)_6O_2$	6.80	4.64	-	-	3.26	-	[64]
5565500	Lu ₂ SiO ₅	6.83	4.75	-	5.45	3.65	2.72	[63]
5570010	BaAl ₂ O ₄	6.80	5.17	5.77	-	4.28	3.44	[11]
5570010	BaAl ₁₂ O ₁₉	7.10	4.77	-	-	-	3.14	[66]
5570010	BaGa ₂ O ₄	6.80	4.51	-	-	3.70	3.06	[11]
5570020	SrAl ₂ O ₄	6.80	4.96	-	6.05	3.94	-	[11]
5570020	SrAl ₁₂ O ₁₉	7.06	4.77	-	-	4.00	-	[66]
5570020	SrGa ₂ O ₄	6.75	4.68	-	-	4.07	2.95	[11]
5570020	$SrGa_{12}O_{19}$	7.00	4.43	-	-	3.59	3.02	[11]
5570040	MgAl ₂ O ₄	6.80	4.31	-	-	-	3.00	[66]
5570130	CaLaAl ₃ O ₇	6.70	4.63	-	-	-	3.33	[67]
5570300	$Gd_3Ga_5O_{12}$	6.77	4.40	-	-	-	-	[68]
5570400	$Y_3Al_5O_{12}$	6.77	4.55	6.05	5.62	3.30	1.97	[69]
5570500	$Lu_3Al_5O_{12}$	6.77	4.75	-	5.35	3.61	-	[70]
5581030	CaMoO ₄	7.00	-	-	3.87	-	2.38	[71]
5582010	Ba ₂ MgWO ₆	7.00	-	_	3.65	-	2.30	[72]
5582030	CaWO ₄	7.10	-	-	4.59	_	2.70	[73]
5582125	SrLaLiWO ₆	7.00	-	_	3.25	-	2.35	[72]
5583100	LaVO ₄	6.80	-	_	3.68	-	2.00	[8]
5583300	GdVO ₄	6.80	-	-	3.65	_	2.41	[8]

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ID		U	E	xcitation	n [eV]	Emission [eV]		
number	Compound	[eV]	А	С	MMCT	А	СТ	Ref.
5583400	YVO ₄	6.80	-	_	3.66	-	2.41	[8]
5584400	YNbO ₄	6.80	_	-	3.81	_	2.30	[8]
5585400	YTaO ₄	6.70	-	-	4.13	-	2.25	[8]
5586000	ZnTiO ₃	6.70	-	-	2.56	-	2.00	[74]
5586020	SrTiO ₃	6.70	-	-	-	-	2.36	[75]
5590010	BaZrO ₃	6.55	4.77	-	-	-	2.50	[76]
5590010	Ba_2ZrO_4	6.50	4.28	-	-	-	2.34	[76]
5590020	SrHfO ₃	6.55	4.59	5.77	-	3.65	-	[77]
5590020	SrZrO ₃	6.55	4.51	-	-	3.44	-	[76]
5590020	SrO	6.40	3.50	-	4.40	3.16	-	[78]
5590020	SrZnO ₂	6.45	3.94	-	-	-	2.76	[79]
5590030	CaO	6.31	3.65	-	4.59	3.40	-	[78]
5590030	CaZrO ₃	6.55	4.35	-	-	3.40	-	[80]
5590120	SrLa ₂ BeO ₅	6.50	4.13	-	-	3.26	2.30	[81]
6690020	SrS	6.25	3.53	4.68	4.51	-	-	[82]
6690030	CaS	6.17	3.60	4.80	5.10	3.49	2.91	[16]
6690040	ZnS	6.40	2.53	3.11	3.35	2.43	1.94	[83]
7790020	SrSe	6.26	3.37	4.35	4.11	_	-	[82]
7790030	CaSe	6.22	3.44	4.30	-	3.35	2.80	[16]

Table 7.1: Continued

7.3. Results

7.3.1. Overview of the spectroscopic data on Pb^{2+} and Tl^+

The A-band excitation energies of Pb²⁺ and Tl⁺ are plotted against the compound identification number in Fig. 7.1a and Fig. 7.2, respectively. The A-band excitation energy decreases from 8 eV in the free Pb²⁺ ion to on average 3 eV in the sulfides and selenides. For Tl⁺ the A-band excitation energy decreases from 6.5 eV in the free ion to approximately 4.5 eV in the iodides. The Tl^+ A-band excitation energy in oxides is located on average at around 5.5 eV except in the simple oxides CaO and SrO, in which the excitation energy is 2 eV lower. Similar to the A-band excitation energy of the Bi³⁺ ion, the decrease in the A-band excitation energy of Pb²⁺ and Tl⁺ follows the nephelauxetic sequence of decreasing electronegativity of the coordinating anion [84]. A comparison with chapter 5 shows that Bi3+ depends stronger on the electronegativity of the coordinating anion than that of Pb²⁺, which depends again stronger than that of Tl⁺. Fig. 7.1b shows an expanded view of the Pb²⁺ A-band excitation energies for the oxide compounds. As was shown for Bi^{3+} , the A-band excitation energy of Pb^{2+} decreases with decreasing electronegativity of the cation that binds the anion. For Tl⁺ only in 9 different oxide compounds the Aband excitation energy was found and a clear trend with decreasing electronegativity of the host cation was not visible, although a similar trend is to be expected.

Table 7.2: The spectroscopic data of Tl^+ in 37 different compounds.

ID		U		Excitati	on [eV]	Em	ission [eV]	
number	Compound	[eV]	А	С	MMCT	A	СТ	Ref.
0000000	free	_	6.47	9.38	_	_	_	[6]
1190002	RbF	7.50	5.93	7.67	-	-	-	[85]
1190003	KF	7.50	5.42	7.13	8.56	-	-	[86]
1190004	NaF	7.50	5.77	7.54	-	-	-	[85]
1190043	KMgF ₃	7.66	6.12	8.27	10.87	5.85	3.69	[87]
1190401	CsY_2F_7	7.60	6.23	-	-	4.81	-	[88]
1190402	RbY ₂ F ₇	7.50	6.23	-	-	4.86	-	[88]
1190402	Rb_2YF_5	7.60	5.40	-	-	4.80	-	[88]
1190403	K ₂ YF ₅	7.50	5.98	-	-	4.80	-	[88]
1190403	KYF ₄	7.50	6.00	-	-	4.26	-	[88]
1198003	KZnF ₃	7.65	6.00	-	-	5.45	-	[89]
2290000	NH ₄ Cl	6.80	5.17	5.77	-	-	-	[90]
2290001	CsCl	6.80	5.00	6.32	-	-	-	[85]
2290002	RbCl	6.80	5.06	6.36	7.10	-	-	[25, 85]
2290003	KCl	6.80	5.03	6.36	6.72	3.94	-	[91, 92]
2290004	NaCl	6.70	4.87	6.19	7.60	-	-	[6]
3390000	NH ₄ Br	6.60	4.84	5.51	-	-	-	[90]
3390001	CsBr	6.60	4.71	5.79	-	-	-	[85]
3390002	RbBr	6.60	4.80	5.85	6.23	-	-	[25, 85]
3390003	KBr	6.60	4.79	5.93	6.50	-	-	[6]
3390004	NaBr	6.55	4.64	5.74	6.60	-	-	[25, 85]
4490000	NH_4I	6.25	4.27	4.77	-	-	-	[90]
4490001	CsI	6.25	4.15	5.14	5.21	-	-	[85, 93]
4490002	RbI	6.25	4.33	5.17	5.45	-	-	[25]
4490003	KI	6.25	4.38	5.30	5.53	-	-	[94]
4490004	NaI	6.25	4.25	5.30	5.50	2.95	-	[25, 95]
4490005	LiI	6.25	4.07	-	-	2.64	-	[96]
4490011	CsBa ₂ I ₅	6.40	4.68	-	-	2.48	-	[97]
5532003	KZn(PO ₃) ₃	7.25	5.39	-	-	4.59	3.65	[98]
5534002	RbH ₂ PO ₄	7.30	5.58	7.35	-	-	-	[99]
5534003	KH_2PO_4	7.30	5.60	7.30	_	4.50	-	[100, 101]
5534030	$Ca_3(PO_4)_2$	7.00	5.08	-	-	3.99	-	[78]
5552100	LaB ₃ O ₆	7.15	5.80	6.50	7.85	3.35	-	[102]
5552300	$GdB_{3}O_{6}$	7.14	5.50	6.80	7.90	-	-	[102]
5555120	SrLaBO ₄	6.80	4.77	_	-	-	2.79	[58]
5590020	SrO	6.40	3.01	_	3.96	2.73	2.11	[78]
5590030	CaO	6.31	3.42	-	3.56	2.64	2.31	[78]

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Figure 7.1: The A-band excitation energies of the Pb^{2+} ion in a) 126 different inorganic compounds and b) 80 different oxide compounds. RE = rare earth, AE = alkaline earth.



Figure 7.2: The A-band excitation energies of the Tl⁺ ion in 37 different inorganic compounds.

Figs. 7.3a and b show the A- and C-band excitation energy as function of the U value of the host compound for the Pb^{2+} and Tl^+ ions, respectively. Both A- and C-band excitation energy increases linearly with increasing U value, although the scatter in data can be quite substantial. The two data points of the Tl^+ A-band excitation energy in Fig. 7.3b that are located approximately 1.5 eV below the average are again for CaO and SrO. The increase in the excitation energies for the Pb^{2+} ion appears more steep as compared to the excitation energies for the Tl^+ ion. The sensitivity of the A-band energy on the chemical environment is exploited by for example Duffy *et al.*, who uses the $6s^2$ ions as a probe ion in molten salts. [103, 104] The $6s^2$ ions are used to probe basicity and ionic–covalent interactions in glasses, since a small variation in the chemical environment has a strong effect on the emission wavelength.



Figure 7.3: The A- and C-band excitation energies of a) Pb^{2+} and b) Tl^+ as function of the U value of the host compound. The solid lines are a guide to the eye.

Fig. 7.4 compares the A-band energies of Tl^+ , Pb^{2+} and Bi^{3+} when doped in the same compound. The A-band excitation energies for Bi^{3+} are taken from chapter 5. The A-band excitation energy is on average 0.5 eV larger in Pb^{2+} than in Bi^{3+} and the A-band excitation energy of Tl^+ in turn is on average 0.5 eV larger than that of Pb^{2+} when doped in the same host compound. The difference in A-band excitation energy between Tl^+ , Pb^{2+} and Bi^{3+} when doped in the same compound is at maximum ± 1 eV.



Figure 7.4: Comparison of the the A-band excitation energy when doped in the same inorganic compound.

Fig. 7.5 shows the Stokes shift distribution between the A-band absorption / excitation and emission of Pb^{2+} . The Stokes shift has a mean value around 0.9 eV and varies between 0.1 and 2 eV. Since there are only 17 compounds in which both the A-band excitation and emission energy of Tl^+ were found, there is insufficient statistics to show a meaningful distribution of the Stokes shift in the A-band emission of Tl^+ .



Figure 7.5: Distribution of the Stokes shift of the A-band emission of Pb²⁺.

7.3.2. The vacuum referred electron binding energies in the ${}^{1}S_{0}$ ground state and ${}^{3}P_{1}$ excited state of Pb²⁺ and Tl⁺

Figs. 7.6 and 7.7 show the stacked VRBE diagrams of different inorganic compounds with the ${}^{1}S_{0}$ ground state and ${}^{3}P_{1}$ excited state of Pb²⁺ and Tl⁺. Information on how the VRBEs at the valence band top and conduction band bottom of the host compounds were derived can be found in Ref. [105]. The electron binding energies in the ${}^{1}S_{0}$ ground state and ${}^{3}P_{1}$ excited state of Pb²⁺ and Tl⁺ show a decreasing (more negative) trend with increasing U value. It appears that the ${}^{3}P_{1}$ excited state of both Pb²⁺ and Tl⁺ tend to follow the location of the conduction band bottom. In compounds with a very low lying conduction band, for example transition metal based compounds such as YNbO₄ or YVO₄, the ${}^{3}P_{1}$ level is located inside the conduction band. When this is the case, the A-band emission is completely quenched and the emission originates exclusively from CT luminescence. For the Tl⁺ ion no CT data was available when doped in compounds containing transition metal ions.



Figure 7.6: Stacked VRBE diagrams for Pb^{2+} doped compounds. E_C is the binding energy at the conduction band bottom, E_X is the binding energy in the host exciton, E_V is the binding energy at the valence band top and ${}^{1}S_0$ and ${}^{3}P_1$ are the binding energies in the Pb^{2+} ground and excited state, respectively.



Figure 7.7: Stacked VRBE diagrams for TI^+ doped compounds. E_C is the binding energy at the conduction band bottom, E_X is the binding energy in the host exciton, E_V is the binding energy at the valence band top and ${}^{1}S_0$ and ${}^{3}P_1$ are the binding energies in the TI^+ ground and excited state, respectively.

Figs. 7.8a and b show the electron binding energies in the ${}^{1}S_{0}$ ground state and ${}^{3}P_{1}$ excited state of Pb^{2+} and Tl^{+} as function of the U value of the host compound. A decreasing trend of the electron binding energy with U value is observed. The ${}^{3}P_{1}$ excited state of Pb^{2+} decreases from –2 eV to –3 eV, while the ${}^{1}S_{0}$ ground state varies from –5 eV to –10 eV with increasing value of U. The ${}^{3}P_{1}$ excited state of Tl^{+} decreases from –1 eV to –4 eV, while the ${}^{1}S_{0}$ ground state varies from –5 eV to –10 eV with increasing value of U. The data points in Figs. 7.8a and b spread by approximately ± 1 eV around the linear fitted lines. The linear decrease of the electron binding energies with U value of Pb^{2+} and Tl^{+} is comparable to that of Bi^{3+} , in which the ${}^{1}S_{0}$ ground state varies from –5 to –10 eV and the ${}^{3}P_{1}$ excited state varies from –2 to –4 eV [84].

7.4. Discussion

In chapter 5, we verified the results from the chemical shift model by comparing the obtained VRBE in the Bi³⁺ ground and excited states with values obtained from independent methods, namely from photo-electron spectroscopy and electrochemical measurements. From photo-electron spectroscopy measurements on PbF₂, Poole *et al.* found the Pb²⁺ top of the 6s² valence band at –9 eV and the top of the fluorine valence band at –11.8 eV relative to the vacuum level [106]. This is comparable to the approximately –10 eV VRBE in the Pb²⁺ ground state we found in CaF₂, SrF₂ and BaF₂ in Fig. 7.6. No reports on electrochemical measurements of Pb²⁺ or Tl⁺ were found that could be used to verify the VRBE in the ground and excited states of Tl⁺ and Pb²⁺.



Figure 7.8: The vacuum referred electron binding energy in the ${}^{1}S_{0}$ ground state and ${}^{3}P_{1}$ excited state of a) Pb^{2+} and b) Tl^{+} plotted as function of the U value.

When doped in a compound, the ionic charge of Tl^+ , Pb^{2+} and Bi^{3+} will be screened by an equal amount of negative charge from the surrounding anions in a way that minimizes the total energy. The Coulomb repulsion between that screening charge and the electrons in the Tl^+ , Pb^{2+} or Bi^{3+} orbitals results in a decreased electron binding energy. The amount by which the binding energy shifts upward is termed the chemical shift E_{CS} [2]. This is illustrated in Fig. 7.9, which shows the chemical shift of the VRBE of the electron in the 1S_0 ground state and 3P_1 excited state of Tl^+ , Pb^{2+} and Bi^{3+} when



Figure 7.9: The chemical shift E_{CS} of the electron in the ${}^{1}S_{0}$ and ${}^{3}P_{1}$ levels of Tl^{+} , Pb^{2+} and Bi^{3+} and in the lowest 4f and 5d levels of Eu^{2+} and Gd^{3+} due to the chemical environment in CaO.

doped in CaO. The VRBE of –20.4 eV for Tl⁺, –31.9 for Pb²⁺ and –45.3 eV for Bi³⁺ in the ground state of the free ions are the same as the 2nd, 3rd and 4th ionization potentials of the atoms, respectively. The ¹S₀ ground state of Tl⁺, Pb²⁺ and Bi³⁺ in CaO undergoes a chemical shift of 15.1 eV, 25.6 eV and 38.4 eV. The chemical shift of the VRBE of the electron in the ³P₁ level of Tl⁺, Pb²⁺ and Bi³⁺ appears smaller and is 12.0 eV, 21.2 eV and 32.6 eV, respectively. In both cases the chemical shift increases approximately in the ratio 1:2:3, similar to the ratio in the amount of screening charge. The increase in the chemical shift of the VRBE of the electron in the ¹S₀ and ³P₁ states when going from the free Tl⁺ to the free Pb²⁺ ion and when going from the free Pb²⁺ to the free Bi³⁺ shift to almost the same energy. Also the VRBE of the excited state of Tl⁺, Pb²⁺ and Bi³⁺ shift to approximately the same energy.

 Tl^+ , Pb^{2+} and Bi^{3+} are isoelectronic and the only difference relates to an increasing nuclear charge. Eu^{2+} and Gd^{3+} are also isoelectronic and for comparison we have added the relevant VRBEs for these two lanthanides in Fig. 7.9. The chemical shifts of the lowest 4f and 5d levels are of the same order of magnitude as in Pb^{2+} and Bi^{3+} , although also important differences are present. While the VRBE of the electron in the ground state of Tl^+ , Pb^{2+} and Bi^{3+} shift to almost the same energy when doped in a compound, the VRBE of the lowest 4f level of Eu^{2+} shifts to an energy that is 8 eV higher compared to the VRBE of the lowest 4f level of Gd^{3+} .

For a lanthanide ion Ln embedded in a chemical environment A, the chemical shift of the 4f electron binding energy is simply represented by the Coulomb interaction of the 4f electron and the total screening charge Q that is located at a screening distance $R_Q(A)$ from the lanthanide nucleus as defined by Eq. (7.1). The interactions between the chemical environment and the 4f electron of the lanthanide are highly complex. However, the simple chemical shift model can satisfactorily model the 4f electron binding energies of the lanthanide ions [2]. It is interesting to verify if Eq. (7.1) also holds for modeling the chemical shift of the 6s electron binding energies of Tl⁺, Pb²⁺ and Bi³⁺.

$$E_{CS}(Ln^Q, A) = \frac{Q}{4\pi\epsilon_0} \frac{e^2}{R_Q(A)}$$
(7.1)

For the lanthanides it was found that the screening distance is almost equal to the Shannon [107] in-crystal ionic radii. Using the shannon crystal radii of 173 pm, 143 pm and 131 pm for Tl⁺, Pb²⁺ and Bi³⁺ ions as the screening distance in Eq. (7.1), we derive chemical shift energies of 8.3 eV, 20.1 eV and 33 eV, respectively. Applying these values of chemical shift to the ground state of the free 6s² ions, the ground states of Tl⁺, Pb²⁺ and Bi³⁺ should shift to -12.1 eV, -11.8 eV and -12.3 eV. Eq. (7.1) predicts a much stronger bonding than the observed VRBE of -5.4 eV, -6.4 eV and -6.9 eV for Tl⁺, Pb²⁺ and Bi³⁺, respectively. The observed chemical shift is much larger than Eq. (7.1) would suggest. Or one may also state that the effective screening distance between the 6s electron and the screening charge from the anion is much less than suggested from the Shannon radius. Using the observed values of the chemical shift of 15.1 eV, 25.6 eV and 38.4 eV for Tl⁺, Pb²⁺ and Bi³⁺, respectively. Apparently the interaction between the electrons in the spatially extended 6s orbital and the surrounding anion orbitals is larger than that of 4f the atomic-like electrons, and a treatment in terms of Eq. (7.1) fails.

Fig. 7.10 shows a comparison of the VRBE in the ${}^{1}S_{0}$ ground and ${}^{3}P_{1}$ excited states of Tl^+ , Pb²⁺ and Bi³⁺ when doped in the same compound. This shows that when doped in the same compound, the VRBE of the electron in the ground state in Tl⁺, Pb²⁺ and Bi³⁺ are found to differ only by 1 to 2 eV. The VRBE of the electron in the ground state of Bi^{3+} is generally located at more negative energy (stronger bonding) than that of Pb^{2+} , which in turn is located at more negative energy than that of Tl⁺. However, occasionally the ordering is reversed or disrupted. Whether this is real or whether this has to do with charge compensation defects, broad excitation bands, presence of pairs [108], different site occupations or wrong spectral assignments remains open. All of these aspects can result in errors in the VRBE, which can alter the observed order of VRBE. Therefore the data should not be considered conclusive as future investigations might yield more dedicated spectroscopic data. This chapter does not provide an interpretation of the luminescent properties of individual compounds, but shows the general trends of the VRBE of the electron in the ${}^{1}S_{0}$ and ${}^{3}P_{1}$ levels of Tl⁺ and Pb²⁺ when doped in different compounds. The VRBE diagrams illustrate that Tl⁺, Pb²⁺ and Bi³⁺ dopant ions show luminescence (from UV to red) in a broad variety of inorganic compounds and these diagrams can aid in the design of new Tl⁺, Pb²⁺ or Bi³⁺ activated luminescent materials.



Figure 7.10: Comparison of the vacuum referred electron binding energy in the a) ${}^{1}S_{0}$ and b) ${}^{3}P_{1}$ levels of TI^{+} , Pb^{2+} and Bi^{3+} when doped in the same inorganic compounds.

erage at -1.8 eV with ± 0.5 eV compound to compound variation [109], which is marked by the shaded region in Fig. 7.10b. In order for Ce^{3+} to show efficient luminescence, the VRBE at the CB bottom should be at higher energy than the VRBE of the 5d electron of Ce^{3+} , *i.e.* higher than -1.8 eV. The VRBE of the electron in the ${}^{3}P_{1}$ excited state generally decreases in going from Tl⁺ to Pb²⁺ to Bi³⁺ and varies between -1 eV and -4 eV depending on the U value of the host. The conduction band bottom in Tl-, Pb- and Bi-based compounds will be determined by the VRBE of the electron in the ground state of Tl⁰, Pb^+ and Bi^{2+} . In chapter 6 we showed that the VRBE in the ground state of Bi^{2+} is located close to or slightly deeper than the VRBE in the ${}^{3}P_{1}$ excited state of Bi $^{3+}$. Assuming that the ground state in Tl⁰ and Pb⁺ ions are also located close to or slightly above the ³P₁ level of Tl⁺ and Pb²⁺, the results suggest that the highest probability to achieve efficient Ce^{3+} luminescence is in Tl- or Pb-based iodide compounds. Since the ground and excited state of TI^+ are found to be at the highest VRBE (least negative VRBE) of the $6s^2$ ions and efficient Ce³⁺ luminescence is already reported in Tl-based halide crystals [110-112], Tl-based halide compounds show the most potential as Ce^{3+} activated scintillator materials. Unfortunately, it is not possible to achieve high densities (>8 g/cm³) for halide compounds.

7.5. Summary and conclusions

Spectroscopic data on the metal-to-metal charge transfer and interconfigurational transitions of the Tl^+ and Pb^{2+} activator ions in 37 and 126 different inorganic compounds, respectively, were gathered from the available literature. The MMCT energy was used to determine the vacuum referred electron binding energy in the ¹S₀ ground state of Tl⁺ and Pb^{2+} in 10 and 32 different host compounds. By adding the A-band energy, the electron binding energy in the ³P₁ excited state of Tl⁺ and Pb²⁺ was obtained. The VRBE of the electron in the ${}^{1}S_{0}$ ground state and ${}^{3}P_{1}$ excited state of Tl⁺ and Pb²⁺ were found to linearly decrease with U value and follow the nephelauxetic sequence. When doped in CaO, the ${}^{1}S_{0}$ ground state of Tl⁺, Pb²⁺ and Bi³⁺ was found to be subjected to a chemical shift of 15.1 eV, 25.6 eV and 38.4 eV for Tl⁺, Pb²⁺ and Bi³⁺, while the chemical shift of the VRBE of the electron in the ³P₁ level of Tl⁺, Pb²⁺ and Bi³⁺ appeared smaller and was found to be 12.0 eV, 21.2 eV and 32.6 eV, respectively. In both cases the chemical shift was found to increase approximately in the ratio 1:2:3, similar to the ratio in the amount of screening charge. When doped in a compound, the ground state of Tl^+ , Pb^{2+} and Bi^{3+} were found to shift to almost the same energy. It was found that the simple chemical shift model that can satisfactorily model the 4f electron binding energies of the lanthanide ions, does not apply to the 6s electron binding energies. Apparently the interaction between the electrons in the spatially extended 6s orbital and the surrounding anion orbitals is larger than that of 4f the atomic-like electrons. Based on the determined VRBEs of the electron in ¹S₀ and ³P₁ states, Tl-based compounds (mainly iodides) showed the most potential for Ce^{3+} activated scintillator materials.

Appendices

In most of the literature we reviewed, the MMCT transition is labeled as D-band. In some of the compounds however, our interpretation deviated from the originally assigned transitions. In Appendices A and B we justify our interpretation and tentative assignments of the spectroscopic data of the Pb^{2+} and Tl^+ ions in the reported literature data. The assignment of the interconfigurational A- and C-band transition are rather straightforward and therefore we limit ourselves to discussing only the MMCT transitions and refer to the original literature for more information.

7.6. Appendix A. The metal-to-metal charge transfer energies of Pb²⁺

SrI₂ Chen *et al.* studied single crystals of SrI₂:Pb²⁺ [32]. They assigned the 3.25, 3.72 and 4.38 eV excitation bands to the A-, B- and C-bands, respectively. However, we suggest that the 2.98 eV, 3.25 eV and 4.38 eV transitions should be attributed to the A-, C- and MMCT-bands, respectively. Due to the forbidden nature of the B-band it should be very weak in photoluminescence spectra. Furthermore, the lowest energy excitation band at 2.98 eV should be assigned to the A-band. The most intense excitation band at 3.25 eV we believe is from the allowed C-band transition. The high-energy of the 4.38 eV transition, indicates this is the MMCT transition band.

 SrB_5O_9Cl Meijerink *et al.* reported a dominant excitation band at 5.58 eV for Pb²⁺-doped $Sr_2B_5O_9Cl$, which they assigned to the C-band. [34] However, they mentioned that it could not be excluded that the C-band (partly) overlapped with a charge transfer transition. Therefore, we assigned the 5.58 eV to the MMCT transition, taken into account that there is probably an error in the exact energy of the MMCT transition due to the overlap with the C-band.

 SrB_5O_9Br Similar to SrB_5O_9Cl , Meijerink *et al.* reported a dominant excitation band at 5.68 eV for $Sr_{1.9}B_5O_9Br$; Pb_{0.1}, which they assigned to the C-band [34]. However, they mentioned that it could not be excluded that the C-band (partly) overlapped with a charge transfer transition. Therefore, we assigned the 5.68 eV to the MMCT transition, taken into account that there is probably an error in the exact energy of the MMCT transition due to the overlap with the C-band.

CaCO₃ Kraienhemke *et al.* recognized a weak band close to the fundamental absorption edge but did not assign this band to a transition [39]. Due to the high energy of the 5.95 eV band, we attribute this band to the MMCT transition.

Y₂SiO₅ and Lu₂SiO₅ Babin *et al.* observed excitation bands at 4.95 eV and 5.25 eV for Y₂SiO₅, which they assigned to A-band emission of Pb²⁺ on a different crystallographic sites [63]. However, the 5.25 eV excitation and corresponding emission at 2.7 eV, showed a concentration dependence typical for the MMCT transition. This lead us to suggest that this transition is more likely from the MMCT transition. In the same report, Babin *et al.* reported excitation at 5.45 eV Lu₂SiO₅, which we tentatively assigned to the MMCT transition instead of excitation of Pb²⁺ on a different crystallographic site.

 $Lu_3Al_5O_{12}$ Babin *et al.* reported the A-band excitation and emission at 4.75 eV and 3.61 eV, respectively [70]. Another excitation at 5.35 eV was observed which they related to a different but undefined origin, which we assigned to the MMCT transition.

CaMoO₄ Yang *et al.* attributed the photoluminescence of CaMoO₄:Pb²⁺ to a charge transition involving Pb²⁺–Ca²⁺ charge transfer [71]. Compounds containing transition metal ions generally have a low-lying conduction band. Therefore, we expect the ³P₁ excited state to be located inside the conduction band. The excitation spectrum of the 2.37 eV emission showed a maximum at 4.59 eV with a shoulder band at 3.87 eV, which were not assigned to specific transitions. However, the 4.59 eV is due to excitation into the host exciton band. The 3.89 eV we assigned to the Pb²⁺–Ca²⁺ charge transfer transition.

Ba₂**MgWO**₆ and **SrLaLiWO**₆ Bleijenberg *et al.* reported the excitation and emission spectra of Ba₂MgWO₆ and SrLaLiWO₆ doped with 5% Pb²⁺ [72]. For Ba₂MgWO₆, emission was observed at 2.3 eV, while host emission was observed at 3.0 eV with a side band at 2.5 eV [72]. The excitation spectrum closely resembled the pure host excitation spectrum, although in the text the authors mentioned an additional excitation band for the Pb²⁺ doped sample at 3.65 eV. Since the low conduction band of tungstates the A-band will be located inside the CB and the reported 3.65 eV most probably originates from the MMCT transition. For SrLaLiWO₆ the authors mentioned an additional excitation band for the Pb²⁺ doped sample at 3.65 eV, which we again assigned to the MMCT transition. **ZnTiO**₃ Wang *et al.* reported on the photoluminescence of ZnTiO₃:0.5% Pb²⁺ nanocrystals [74]. They reported excitation and emission bands at 2.56 eV and 2.0 eV, respectively. These transitions were assigned to excitation and emission from the VB top to Pb²⁺. However, general for elements with the 6s² electron configuration this charge transfer is from the Pb²⁺ to the CB bottom.

SrO and CaO Van der Steen *et al.* reported C-band excitation at 4.4 eV and 4.59 eV for SrO and CaO, respectively [78]. Unfortunately they did not show the excitation spectrum. However, due to the high energy and by comparing with the MMCT in SrO:Bi³⁺, we assign the 4.4 eV and 4.59 excitation bands to the Pb²⁺ \rightarrow CB CT.

SrS and SrSe Yamashita *et al.* reported on the luminescence of SrS:Pb and SrSe:Pb [82]. For SrSe:Pb they found excitation bands at 3.37, 4.11 and 4.35 eV and for SrS:Pb they found excitation bands at 3.53, 4.51 and 4.68 eV. The first were assigned to A-band excitations which we believe to be correct. The second were not assigned, however, considering the 0.5 eV broad band and high-energy this is probably the MMCT transition. The third excitation band was also unassigned, however based on the narrow width of approximately 0.2 eV we suggest this is the C-band.

7.7. Appendix B. The metal-to-metal charge transfer energies of Tl⁺

KMgF₃ Scacco *et al.* assigned excitation at 8.98 eV to the D-band [87]. However, in a later review by Sun *et al.*, they assigned the 10.87 eV excitation band in the report of Scacco *et al.* to the D-band [113].

CsI Kawai *et al.* reported a VB \rightarrow Tl⁺ CT-band at 5.21 eV [93]. As motivated in the introduction we assigned this band to the Tl⁺ \rightarrow CB transition.

 LaB_3O_6 and GdB_3O_6 Tanimizu *et al.* observed an excitation band overlapping with the host exciton band for LaB_3O_6 and GdB_3O_6 [102]. They showed that this broad band consisted of three overlapping bands which they attributed to three C-bands. However, we suggest it is more probable that at least one of these bands originates from the MMCT.
SrO and CaO Van der Steen *et al.* reported excitation bands at 3.01 eV and at 3.96 eV for SrO and 3.42 eV and 3.56 eV for CaO which were not assigned to any optical transitions [78]. The low excitation has a high-energy emission, matching with the A-band transition. The high-energy excitation is broader and has lower energy emission band, matching with a MMCT transition.

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Summary

Scintillation materials convert high-energy radiation into many visible photons and, in combination with a photodetector, are used as ionizing radiation detectors. Since the discovery of ionizing radiation, there have been intensive research efforts in finding new, better performing scintillators, resulting in the development of a large variety of scintillation materials. Each scintillation material is tailored to a specific application to have the best performance. With ever-increasing material demands set by the various applications of scintillators, the search for even better performing scintillation materials remains an active field. This thesis explores new avenues of scintillation materials research in order to find the next-generation of scintillation materials.

A valuable technique to improve the scintillation properties of existing scintillation materials is by means of co-doping. Chapter 2 discusses the effects of Na⁺, Mg²⁺, Ca²⁺, Sr²⁺ and Ba²⁺ doping on the scintillation properties of CeBr₃. Adding a small amount of aliovalent dopant ions has a pronounced effect on the scintillation properties of CeBr₃. Ca²⁺ and Sr²⁺ doped CeBr₃ show an additional perturbed Ce³⁺ site in the photoluminescence and X-ray excited luminescence spectra. Furthermore, these crystals show a nearly constant light yield below 300 K, an increased decay time and the appearance of multiple glow peaks in the thermoluminescence spectra. We proposed that bromine vacancies, formed as charge compensation for the aliovalent dopants, act as electron traps which can then explain the different scintillation behavior of Ca²⁺ and Sr²⁺ doped CeBr₃.

In chapters 3 and 4 we investigate the potential of Bi^{2+} , a relatively new dopant ion in the field of scintillation materials with red-to-infrared emission wavelengths. Chapter 3 discusses the X-ray induced valence change of Bi^{3+} to Bi^{2+} and the luminescence and scintillation properties of bismuth doped $Li_2BaP_2O_7$. Although absorption measurements show that Bi^{3+} is present in the as-prepared $Li_2BaP_2O_7$:Bi, no bismuth-related photoluminescence is observed even at temperatures as low as 10 K. However, during Xray excitation the characteristic deep-red radioluminescence of Bi^{2+} is observed. After X-ray irradiation, the red luminescence of Bi^{2+} can be excited optically, indicating that upon X-ray irradiation the Bi^{3+} is reduced to Bi^{2+} . Chapter 4 deals with the luminescence and charge carrier trapping in YPO_4 :Bi. X-ray excited luminescence measurements show emission from isolated Bi^{3+} , Bi-pairs and Bi^{2+} . Thermoluminescence measurements of YPO_4 doped with Bi^{3+} and/or Tb^{3+} show that bismuth can act both as electron and as hole trap. The spectroscopic data obtained in chapters 3 and 4 are used to locate the electron binding energies in the ground and excited states of Bi^{3+} and Bi^{2+} inside the vacuum referred binding energy diagrams of $Li_2BaP_2O_7$ and YPO_4 , respectively. The constructed VRBE diagrams provide an interpretation for the commonly observed Bi-pair emission.

Currently most research is focused on halide crystals with lanthanide dopants and Ce^{3+} and Eu^{2+} in particular. These crystals have a rather low density of at maximum around 6 g/cm³, while scintillation materials with the highest density and stopping power are needed for radiation detectors in the next-generation of particle colliders and other high-energy physics experiments. To find new and denser scintillation materials, oxide materials with densities that can go beyond 8 g/cm^3 are attractive candidates. To ensure a high stopping power, oxides containing elements with a high atomic number such as Tl, Pb and Bi are among the most promising materials. In order to evaluate if Ce^{3+} can luminescence efficiently in these type of host compounds, knowledge on the impurity levels of Tl^+ , Pb^{2+} and Bi^{3+} relative to the host conduction and valence bands is required. The electron binding energies in the impurity levels of Bi³⁺ and how these change in relation to the chemical environment are discussed in chapter 5, while those of Tl^+ and Pb^{2+} are discussed in chapter 7. To determine these vacuum referred electron binding energies, spectroscopic data on the metal-to-metal charge transfer and interconfigurational A- and C-band transitions of Tl⁺, Pb²⁺ and Bi³⁺ were collected form the archival literature. The VRBE in the ³P₁ excited state and ¹S₀ ground state of Tl⁺, Pb²⁺ and Bi³⁺ decreases linearly with the U value of the host compound and follow the nephelauxetic sequence. The chemical shift of Tl⁺, Pb²⁺ and Bi³⁺ increases approximately in the ratio 1:2:3, which is similar to the ratio in the amount of screening charge. Although the difference in the amount of screening charge around the dopant ion when incorporated in a compound, the ground state of Tl^+ , Pb^{2+} and Bi^{3+} shift to almost the same energy. It was found that the simple chemical shift model that can satisfactorily model the 4f electron binding energies of the lanthanide ions, does not apply to the 6s electron binding energies of Tl⁺, Pb²⁺ and Bi³⁺. Apparently the interaction between the electrons in the spatially extended 6s orbital and the surrounding anion orbitals is larger than that of 4f the atomic-like electrons. Based on the determined VRBEs of the electron in the ${}^{3}P_{1}$ and ¹S₀ states, it is unlikely that Ce³⁺ will luminescence in Tl-, Pb- or Bi- based oxide compounds. However, this research provides new directions for further exploration of high density, high stopping power scintillation materials, namely to evaluate if Pb²⁺ will show luminescence in a Bi-compound or Bi³⁺ in a Pb-compound. Also the possibility of Tl⁺, Pb²⁺ and Bi³⁺ as activator in suitable, dense host compounds could be explored.

In chapter 6 we show that in general the VRBE in the Bi^{2+} ground state is located at lower energy as compared to the VRBE in the ${}^{3}P_{1}$ excited state of Bi^{3+} . In the presence of Bi-pairs, it is then upon excitation of one Bi^{3+} energetically favorable for electron transfer to occur, and to form a $Bi^{2+}-Bi^{4+}$ pair. The recombination of the electron and hole on the $Bi^{2+}-Bi^{4+}$ pair can occur non-radiatively or radiatively, in the later case the resulting visible emission band is often observed and ascribed to pair emission. The proposed mechanism of charge transfer between Bi-pairs can also explain the quenching behavior of Bi^{3+} luminescence when the Bi^{3+} concentration increases.

Samenvatting

Scintillatie materialen zetten hoogenergetische straling om in zichtbaar licht en worden in combinatie met een fotodetector gebruikt als stralingsdetector. Sinds de ontdekking van ioniserende straling is er intensief onderzoek naar nieuwe en betere scintillatie materialen. In de loop der jaren zijn er diverse scintillatie materialen ontwikkeld die zijn afgestemd op de specifieke eisen van de verschillende toepassingen. De eisen die aan scintillatie materialen worden gesteld, worden steeds strikter waardoor het onderzoeksveld naar scintillatie materialen nog steeds zeer actief is. In dit proefschrift verkennen we nog onbekend terrein in het onderzoek naar scintillatie materialen met als doel om de volgende generatie aan scintillatie materialen te vinden.

Een waardevolle techniek om de scintillatie eigenschappen van een materiaal te verbeteren is door middel van co-dotering. In hoofdstuk 2 worden de effecten van Na⁺, Mg²⁺, Ca²⁺, Sr²⁺ en Ba²⁺ dotering op de scintillatie eigenschappen van CeBr₃ besproken. Het toevoegen van een kleine hoeveelheid aan doterings ion heeft een significante invloed op de scintillatie eigenschappen van CeBr₃. Voor Ca²⁺ en Sr²⁺ gedoteerd CeBr₃ wordt een verstoorde Ce³⁺ roosterlocatie waargenomen in de foto- en Röntgengeëxciteerde emissie spectra. Verder hebben deze gedoteerde kristallen een bijna constante lichtopbrengst boven 300 K, een langere vervaltijd en meerdere gloeipieken in het thermoluminescentie spectrum. Ons voorstel is dat broom vacatures die gevormd zijn om de lading van het doterings ion te compenseren, werken als elektron vangstcentra wat het afwijkende scintillatie gedrag van Ca²⁺ en Sr²⁺ gedoteerd CeBr₃ kan verklaren. In hoofdstukken 3 en 4 onderzoeken we Bi²⁺, een relatief nieuw doterings ion in het

onderzoek naar scintillatie materialen met emissie in het rood tot infrarood. In hoofdstuk 3 worden de Röntgen geïnduceerde valentie verandering van Bi³⁺ naar Bi²⁺ en de luminescentie en scintillatie eigenschappen van bismut gedoteerd Li2BaP2O7 besproken. Ondanks dat absorptie metingen laten zien dat er Bi³⁺ aanwezig is in het gesynthetiseerde Li₂BaP₂O₇:Bi, is er geen bismut-gerelateerde fotoluminescentie waargenomen, zelfs niet wanneer het materiaal wordt afgekoeld tot 10 K. Tijdens Röntgen geëxciteerde luminescentie metingen wordt echter wel de karakteristieke dieprode radioluminescentie van Bi²⁺ waargenomen. Na Röntgen bestraling is het mogelijk om deze rode luminescentie ook optisch te exciteren, wat er op duidt dat ten gevolge van de Röntgen straling het Bi³⁺ wordt gereduceerd naar Bi²⁺. In hoofdstuk 4 worden de luminescentie en de eigenschap om ladingsdragers in te vangen van bismut in YPO₄ besproken. In de Röntgen geëxciteerde luminescentie spectra zijn emissie banden van geïsoleerd Bi³⁺, Bi-paren en Bi^{2+} waargenomen. Thermoluminescentie metingen van YPO_4 gedoteerd met Bi^{3+} en/of Tb³⁺ tonen aan dat bismut zowel als elektronen als gaten vangstcentrum kan dienen. De spectroscopie data uit hoofdstukken 3 en 4 worden gebruikt om de bindingsenergieën van elektronen in de grond en aangeslagen toestanden van Bi³⁺ en Bi²⁺ te bepalen, welke worden weergegeven in een vacuüm gerefereerd bindingsenergie schema

van $Li_2BaP_2O_7$ en YPO₄. Deze VRBE schema's geven daarnaast een interpretatie voor de vaak geobserveerde Bi-paren emissie.

Op dit moment wordt er veel onderzoek gedaan naar halide kristallen met lanthanide doterings ionen en voornamelijk Ce^{3+} en Eu^{2+} . Deze kristallen hebben een nogal lage dichtheid van maximaal rond de 6 g/cm³, terwijl juist scintillatie materialen met een zo hoog mogelijke dichtheid en stoppend vermogen nodig zijn voor de stralingsdetectoren in de volgende generatie deeltjesversnellers en andere hoogenergetische fysica experimenten. Om nieuwe materialen te vinden met een nog hogere dichtheid zijn juist oxides met een dichtheid hoger dan 8 g/cm³ het meest interessant. Oxides gebaseerd op Tl, Pb of Bi, elementen met een hoog stoppend vermogen, zijn het meest belovend. Om er achter te komen of Ce^{3+} luminescentie geeft in dit type materialen, is het noodzakelijk om te weten hoe de energieniveaus van Tl⁺, Pb²⁺ en Bi³⁺ als doterings ion zich verhouden tot de geleidings en valentie band van het gastrooster. De bindingsenergieën van de elektronen in de energieniveaus van Bi³⁺ en hoe deze veranderen in relatie tot de chemische omgeving zijn het onderwerp van hoofdstuk 5 en die van Tl^+ en Pb^{2+} van hoofdstuk 7. Om de VRBEs te bepalen, is de spectroscopische data met betrekking tot de metaal-naar-metaal ladingsoverdracht en interconfigurationele A- en C-band overgangen verzameld uit de beschikbare literatuur. De VRBE in de ³P₁ aangeslagen toestand en de ¹S₀ grond toestand van Tl⁺, Pb²⁺ en Bi³⁺ laten een lineaire afname met de U waarde van het gastrooster zien en volgen de nephelauxetic serie. De chemische verschuiving van Tl⁺, Pb²⁺ en Bi³⁺ neemt ongeveer toe in de verhouding 1:2:3, wat gelijk is aan de verhouding tussen de hoeveelheid aan afschermingslading. Ondanks het verschil in de hoeveelheid aan afschermingslading die de ionen omringt wanneer deze zijn ingebouwd in een gastrooster, verschuift de grond toestand van Tl⁺, Pb²⁺ en Bi³⁺ naar ongeveer dezelfde energie. Het eenvoudige chemische verschuivings model dat de 4f elektron bindingsenergieën van de lanthanide ionen kan voorspellen, blijkt niet te gelden voor de 6s elektron bindingsenergieën van Tl⁺, Pb²⁺ en Bi³⁺. Blijkbaar is de interactie tussen de elektronen in de ruimtelijk verspreide 6s orbitalen en de chemische omgeving dusdanig groter dan die van de gelokaliseerde, atoom-achtige 4f elektronen. Op basis van de VRBEs van de elektronen in de ${}^{3}P_{1}$ en ${}^{1}S_{0}$ toestanden, is het onwaarschijnlijk dat Ce³⁺ luminescentie zal geven in Tl-, Pb- of Bi-gebaseerde oxide verbindingen. Dit onderzoek geeft echter wel nieuwe richtingen voor verdere verkenning van scintillatie materialen met een hoge dichtheid en een groot stoppend vermogen. Namelijk, het bekijken of Pb^{2+} in een Bi-verbinding luminescentie geeft of Bi³⁺ in een Pb-verbinding. Ook de mogelijkheid om Tl^+ , Pb^{2+} en Bi^{3+} te gebruiken als activator in geschikte gastroosters met een hoge dichtheid zou onderzocht kunnen worden.

In hoofdstuk 6 laten we zien dat de VRBE in de grond toestand van Bi^{2+} over het algemeen bij een lagere energie ligt dan de VRBE in de $Bi^{3+} {}^{3}P_{1}$ aangeslagen toestand. Wanneer Bi-paren aanwezig zijn, is het energetisch gunstig dat er na excitatie een elektron van het ene Bi^{3+} ion wordt overgedragen op het andere ion waarbij zich een $Bi^{2+}-Bi^{4+}$ paar vormt. De recombinatie van het elektron en gat op het $Bi^{2+}-Bi^{4+}$ paar kan zowel stralingsloos als via de uitzending van een foton, waarbij in het laatste geval de resulterende zichtbare emissie wordt toegeschreven aan Bi-paar emissie. Dit voorgestelde mechanisme van ladingsoverdracht tussen Bi-paren kan ook het uitdovingsgedrag van Bi^{3+} luminescentie bij hogere Bi^{3+} concentraties verklaren. Vier jaar onderzoek aan de TU Delft is voorbij gevlogen! Mijn beslissing om naar Delft te verhuizen en promotieonderzoek te gaan doen is een van mijn beste beslissingen geweest. Het heeft mij ontzettend veel opgeleverd, niet alleen aan ervaring op het gebied van wetenschappelijk onderzoek maar ook op persoonlijk vlak. Het scintillatie onderzoek was voor mij een compleet nieuw onderwerp waar ik mij volledig in kon verdiepen en veel nieuwe technieken en methoden kon leren. Daarnaast kwam mijn interesse voor de lucht- en ruimtevaart sterk naar boven en daar kan ik mij de komende jaren bij Fokker Aerostructures verder in kan verdiepen. Echter het voltooien van een PhD is niet iets wat je alleen doet. En daarom ben ik een hoop mensen dankbaar voor hun bijdrage aan dit proefschrift.

Allereerst wil ik mijn ouders bedanken voor hun onvoorwaardelijke vertrouwen en steun. Door jullie ben ik in staat mijn dromen waar te maken en ben ik geworden wie ik nu ben. Natuurlijk wil ik ook mijn geweldige familie bedanken. Jullie begrijpen niet altijd waar ik mee bezig ben, maar daardoor waardeer ik jullie interesse en steun des te meer. Maikel, jouw wil ik in het bijzonder bedanken voor het coverdesign van dit proefschrift.

Elise, jou ben ik nog het meest dankbaar. Ik kan altijd rekenen op jouw onvoorwaardelijke liefde. Jij laat mij mezelf zijn en samen zijn we een perfect team. Ook wil ik mijn tweede familie die ik erbij heb gekregen bedanken voor hun interesse en enthousiasme voor mijn onderzoek.

Of course this thesis was not possible without the help of my promotor and colleagues. Pieter, thank you for your supervision the past four years. I am grateful for everything you thought me, from quantum mechanics to writing papers and from the Dorenbos models to presenting science. A big thanks to the technicians for keeping all the setups up and running. I would like to thank all of my colleagues for their valuable scientific discussions as well as our group activities at 't Koepeltje and more. Also I would like to thank Vladimir Ouspenski and colleagues at Saint Gobain for their contribution and industrial view on this project. I am also grateful to Karl Krämer and Daniel Biner for growing the crystals I have investigated. Thanks to all the other people I met which I forgot to mention here in making my time as a PhD candidate an incredible experience.

Wie ik als laatste wil bedanken maar zeker niet wil vergeten te vermelden zijn mijn goede vrienden uit Wijchen. Ook al woont het merendeel van ons nu verspreid over Nederland en is het contact minder dan voorheen, waardeer ik de momenten dat we samen een biertje drinken des te meer!

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Curriculum Vitæ

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