

GREEN LUMINESCENCE SYSTEM CONTAINING THE Tb³⁺- β -DIKETONATE COMPLEX DOPED IN THE EPOXY RESIN AS EMITTER CENTER

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Optical properties of the terbium tri(acetylacetonate) tetrahydrated, [Tb(ACAC)₃(H₂O)₄], doped in the epoxy resin, in the solid state are reported. The polymeric-Tb³⁺-complex and the precursor compound were characterized by elemental analysis, thermogravimetry (TG), differential scanning calorimetry (DSC), infrared and electronic spectroscopy. Due to efficient energy transfer from the polymer and ACAC to the rare earth ion, the polymer phosphorescence intensity decreases with the increasing of the Tb³⁺ ion concentration. The luminescence spectra present narrow bands characteristic from the intraconfigurational transitions: a) emission data ⁵D₄→⁷F_J (J = 6, 5, 4, 3, 2, 1, 0) and b) excitation data ⁷F₆→⁵L₁₀ (350 nm), ⁷F₆→⁵L₁₀ (369 nm), ⁷F₆→⁵G₆ (376 nm), ⁷F₆→⁵D₃ (380 nm), ⁷F₆→⁵D₄ (488 nm). The emission arising from the emitting ⁵D₃ level was not recorded indicating that a rapid ⁵D₃→⁷F₄ radiationless transition is induced by the high vibrational frequencies of the organic ligand. High lifetime measurement (τ = 0.81 ms) typical of terbium ion suggests that the polymeric system has higher luminescence efficiency than in the Tb³⁺-hydrated compound. The concentration quenching of luminescence was verified in the polymer:[Tb(ACAC)₃]15% system. The [Tb(ACAC)₃] complex doped in epoxy resin shows high green luminescence intensity.

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TRACE AMOUNTS OF RARE EARTH ELEMENTS IN HIGH PURE GADOLINIUM OXIDE BY SECTOR FIELD INDUCTIVELY COUPLED PLASMA MASS SPECTROMETRY (SF ICP-MS)

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In recent years rare earth elements have received much attention in the fields of geochemistry and industry. Gadolinium oxide is used for many different high technology applications such as infrared absorbing automotive glass, petroleum cracking catalyst, gadolinium-yttrium garnets, used in microwave applications, and color TV tube phosphors. It can also be used in optical glass manufacturing and in the electronic industry. Rapid and accurate determinations of trace REE are increasingly required as industrial demands expand. In general, the inductively coupled plasma mass spectrometer presents some advantages for trace element analysis, due to high sensitivity and resolution, when compared with other analytical techniques. In this work, sector field inductively coupled plasma mass spectrometry (SF ICP-MS) was used. Sixteen elements (Sc, Y and 14 lanthanides) were determined selectively with the SF ICP-MS system using the concentration gradient method. The detection limits with the SF ICP-MS system were about 0.2 to 8 pg.mL⁻¹. The recovery percentage ranged from 95 to 100% for different rare earth elements. The %RSD of the methods varying between 1.5 and 2.5 % for a set of five (n=5) replicates was found for the IPEN's material and for the certificate reference sample. Determination of trace REEs in two high pure gadolinium oxide samples (IPEN and JMC) were performed. IPEN material are highly pure (>99.99%) and were successfully analyzed without spectral interference.

Keywords: Rare Earths Elements, Inductively Coupled Plasma Mass Spectrometry, Gadolinium oxide.

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