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Spectrally tunable magnetic nanoparticles designed for distribution/recollection applications

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The comprehensive goal of this research is the synthesis and characterization of nanomaterials that are spectrally tunable in terms of their electromagnetic signal, are robust, magnetic (allowing their piloted movement), and have the potential to be functionalized for the detection of CBRNE threats. Various chemical methods were utilized for synthesis of magnetic (iron) and luminescent rare earth (RE) components, and their mixtures. Effects of integrating an iron core into RE luminescent lattices (excited by UV, emit in the VIS) were investigated. The determination of the optimum balances between magnetic and luminescent components such that the magnetism was maximized while maintaining acceptable fluorescence integrity will be discussed. The emphasis of this work is focused on developing a distributed sensor suitable for use in the terrestrial environment. The robust properties of using a RE luminescent shell would allow the particles to be resistant to photobleaching. Additionally the chemical stability of the RE shell would allow operation in a variety of pH conditions. The magnetic core will ultimately allow the distributed particles to be reculated. [doi:10.1063/1.3355900]

Multifunctional nanostructures have attracted significant interest. Specifically magnetic fluorescent nanoparticles serve as manipulative fluorescent probes with applications in nanoelectronic, biological, and catalytic applications. The simplest approach would be to decorate a magnetic nanoparticle with a fluorescent organic dye. However, due in part to the susceptibility that organic fluorophores may photobleach more elaborate measures have been taken. A variety of semiconductor coated magnetic particles have already been synthesized including Fe2O3 and FePt coated with MS or MSe (where M = Cd, Zn, Hg or Pb).1–7 Both Liu and Yu have demonstrated that at room temperature the magnetic core semiconducting shell particles exhibit significant magnetic saturation and luminescence without the use of a spacer.8,9 This result is significant since a balance exists between the magnetic and fluorescent properties.

Rare earth (RE) nanoparticles are also being considered as an alternative to semiconductors for similar applications due in part to their long fluorescent lifetimes, high quantum yields, low photobleaching, high chemical stability, and are not prone to degradation over time.10–14

We have successfully generated bright orange, yellow, and red emitting FL-Mag-NPs, blue and green emitters were more challenging. Combining a series of RE dopants into a matrix produced a high degree of tunability (unique signatures), with luminescence covering the visible spectrum. The second objective of this research focuses on the feasibility of effective distribution, recollection and/or aggregation, and spectral analysis of our synthesized FL-Mag-NPs in variable environmental (aqueous and soil) conditions. We have demonstrated the applicability of an external magnetic field (using a field portable electromagnet) as a technique for successful recollection of the Fluorescent-Magnetic-Nanoparticles (FL-Mag-NPs) out of water or soil media, bringing them to the surface for more feasible remote interrogation.

The preparation of the aqueous ferrofluid comprised of iron nanoparticles was performed by a modified polyol process.15,16 The ferrofluid is then placed into a Parr bomb. To which 8.9 mmol of Y(NO3)3 and 0.47 mmol of Eu(NO3)3 were dissolved in 30 ml of distilled water to which 7.5 mmol of Na2VO4 was added. While stirring a white precipitate formed. The bomb was then sealed and placed in a conventional microwave for 1 min. The precipitate was suspended in 40 ml of distilled water, and 9.38 mmol of 1-hydroxyethane-1,1-diphosphonic acid was added. The resulting solution was made basic by adding sodium hydroxide and the basic suspension was left to stir overnight. The resulting brown precipitate was magnetically separated, washed several times with DI water and vacuum dried. The as-prepared particles were characterized using transmission electron microscopy (TEM) (Joel JEM-1230 operated at 120 kV) and x-ray diffraction (XRD) (X’Pert Philips Materials Research diffractometer, with Cu Kα radiation). The compositions of the samples were determined using x-ray fluorescence (XRF) (Horiba Jobin Yvon XGT-7000). The powders were adhered to a glass slide using double-sided tape. The x-ray tube voltage was set to 50.0 kV, the current was set to 1.0 mA, and the sample diameter was 100 μm. Temperature and magnetic field variations in the magnetization χ of the as-prepared and treated samples were measured using a commercial superconducting quantum interference device magnetometer (Quantum Design).

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To determine if the ability to recollect the particles once distributed, four different soil types were used to determine magnetic recollectability along a moisture gradient. These soils were characterized as being sand, loamy sand, clay mixture, and loam. Since each soil had different water potentials, the moisture variables used differed among soil types and approached saturation for each. Six moisture levels were evaluated for each soil that varied between soil types, including a dry variable. On the soil surface, 50 mg of particles were distributed as evenly as possible and allowed to interact for 2 h in a sealed container. After this time the lid was removed and the samples were allowed to air dry for approximately 24 h. Magnetic recollection was accomplished holding a samarium cobalt magnet not closer than 3 mm from the surface of the soil.

The synthesis and characterization of the ferrofluid has been described in detail in other publications. Briefly, the particles consist of an iron center surrounded by an iron oxide layer. The particles measure ~10–15 nm in diameter and possess a magnetic saturation value of ~92 emu/g. The primary advantage of starting with a ferrofluid is that the magnetic particles are well isolated from each other and provide a good nucleation source for subsequent shell formation, as indicated in Fig. 1(a). Yttrium vanadate (YVO₄) was chosen as the host lattice for the luminescent shell. High luminescent quantum yield phosphors, in a variety of colors, can be produced by codeloping the yttrium vanadate lattice with RE ions. In particular Eu doped YVO₄ is an important commercial phosphor possessing a quantum yield of ~70%. It is used in color televisions, the high pressure mercury lamp, and as a scintillator in medical image detectors. It is significant to note that YVO₄ can be produced via a relatively mild hydrothermal route reducing the possibility that the magnetic core oxidizes. The XRD pattern of the resulting magnetic 5% Eu doped YVO₄ shell particles synthesized hydrothermally in the presence of ferrofluid is provided in Fig. 1(b). The diffraction pattern indicates that the YVO₄ adopts a tetragonal crystal system and matches well with intensified charge coupled device database pattern 01-072-0274. It is significant to note that no indication of Fe or Fe₂O₄ is evident in the diffraction pattern indicative that the magnetic core has a thick coating of YVO₄. Also, the absence of a peak attribute to Eu is also a good indication that it is well dispersed in the lattice. The volume-weighted average crystalline size calculated from the XRD peak width using Scherrer’s equation indicates that the crystallite size is ~23 nm. This is in agreement with the TEM micrograph shown in Fig. 1(c). While the results from XRF confirm that the Eu doping was 5.1%, the percentage of Fe was lower than expected at 10.5%. Since XRF is considered a surface technique this low value indicates that the magnetic core is well encased in a thick shell of YVO₄.

The temperature variations of $\chi$ for both the field cooled (FC) and zero field cooled cases (ZFC) of the 5% Eu doped YVO₄ is shown in Fig. 2(a). It is evident from the curves that the sample displays an overall superparamagnetic character. However, the width of the blocking temperature for the ZFC case is suggestive of a wide distribution of particle sizes. The convergence of the temperature variation of $\chi$ above the blocking temperature for the FC and ZFC data is characteristic of interacting particles. This is indicative that some of the particles might have multiple magnetic centers resembling pomegranates rather than a true core shell nature. The results of measuring the hysteresis loops at 5 and 300 K are shown in Fig. 2(b). The highly symmetrical shape of the loops indicates that other mixed phases are not present in the sample. The magnetic saturation of the magnetic 5% Eu doped YVO₄ was determined to be ~41.1 emu/g significantly less than the 92 emu/g reported for the particles making up the ferrofluid. This lower value is attributed to the diamagnetic contribution of the YVO₄ lattice. While what diminished the overall magnetic responsibility is significant enough to allow the particles to be piloted in the presence of a magnetic field as shown in Fig. 2(c).

The luminescent spectra of magnetic 5% Eu doped YVO₄ is shown in Fig. 3(a). It has been well documented that strong luminescence occurs from efficient energy transfer from the VO₄²⁻ group to Eu³⁺, specifically attributed to the charge transfer from the oxygen ligands to the central vanadium atom, and this transition is known to occur at ~300 nm. Thus, we have chosen 290 nm as the excitation wavelength. It has also been previously determined that the quantum yield shows a maximum at a doping level of ~5 mol% and slowly decreases as the concentration is increased further. Concentration quenching is observed if the mean distance between neighboring Eu ions decrease below a fixed value. The characteristic luminescence spectra shown
in an environmental recollection scenario, their behavior was observed on both moist and dry soils. Four different soil types were used that varied in pH, water potential, and composition (sand, loamy sand, loam, and clay mixture). The moisture variables for each soil were determined by their water potential and approached saturation for each different soil type. While no correlations were found between magnetic recollectability and pH, moisture, and composition were found to affect recollection. The particles were found to readily adsorb water from the soil and adhered to the soil surface, forming a crust that clung to the soil underneath it. Magnetic recollection was near 100% on all dry soils, however, on moist soil recollection was observed to depend more on the soil density, moisture, and the aggregation of the particles. Since the particles adhered to the moist soil below, recollection from denser soils (sand and loamy sand) was difficult. While recollection was more successful on lower density soils (loam and clay mixture), the recollected particles were on the soil surface.

In conclusion, we have demonstrated the synthesis of multifunctional nanoparticles that are both luminescent and magnetic at room temperature. By varying the RE dopant, it is possible to vary the emission wavelength or create unique signatures by combining them.