Single excited dual band luminescent hybrid carbon dots-terbium chelate nanothermometer

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Abstract

The report introduces hybrid polyelectrolyte-stabilized colloids combining blue and green emitting building blocks, that are citrate carbon dots (CDs) and [TbL]+ chelate complexes with 1,3-diketonate derivatives of calix[4]arene. The joint incorporation of green and blue emitting blocks into the polysodium polystyrenesulfonate (PSS) aggregates is carried out through the solvent-exchange synthetic technique. The coordinative binding between Tb3+ centers and CD surface groups in initial DMF solutions both facilitates joint incorporation of [TbL]+ complexes and the CDs into the PSS-based nanobeads and affects fluorescence properties of [TbL]+ complexes and CDs, as well as their ability to temperature sensing. The variation of the synthetic conditions is represented herein as a tool for tuning the fluorescent response of the blue and green emitting blocks upon heating and cooling. The revealed regularities enable developing either dual band luminescent colloids for monitoring temperature changes within 25-50 °C through double color emission, or to transform the colloids into ratiometric temperature sensors via simple concentration variation of [TbL]+ and CDs in initial DMF solution. Novel hybrid carbon dots-
terbium chelate PSS-based nanoplatform opens an avenue for new generation of sensitive and customizable single excited dual band nanothermometers.

**Keywords:** carbon dots; terbium complex; calix[4]arene; molecular thermometer; nanothermometry; hybrid nanoparticles.

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Introduction

Combination of oppositely charged luminophoric blocks with different nature within one nanobead is a promising route to enforce or modify their luminescent response in aqueous solutions. Incorporation of the different luminescent blocks into polymeric nanobead has been widely exemplified by different polymeric matrices\textsuperscript{1–4}, including those non-covalently generated from self-aggregation of water soluble polymers\textsuperscript{5–7}. Polyelectrolytes provide very promising type of water soluble polymers due to the multitude of surface exposed ionic groups providing high affinity to oppositely charged surfaces or counterions\textsuperscript{8–10}. The counterion binding is the well-known factor facilitating their intra- and inter-molecular interactions, in turn, favoring coil-like conformation of polyelectrolytes followed by their self-assembly into specific clew-like aggregates\textsuperscript{11}. However, fundamental backgrounds underlying the use of polyelectrolyte-based aggregates as promising nanobeads for the components with opposite charge and different nature are insufficiently recognized. It is known that the high surface charge of the nanoparticles resulted from the nature of hydrophilic exterior layer and predominantly constituted by chelating anions is the reason for its high affinity to counterions, including metal ions\textsuperscript{12,13}. The coordinative binding of CDs with metal ions modifies the fluorescence of CDs, which is the reason for using them as effective optical sensors\textsuperscript{14–16}, although an impact of such binding on joint incorporation of blue emitting CDs with green emitting Tb\textsuperscript{3+} centers into one polyelectrolyte-based nanobead is not enough recognized.

Ln\textsuperscript{3+}-based species possessing narrow emission bands, large Stokes shift, and high emission quantum yields inspired many reports on Ln\textsuperscript{3+}-based thermometers with outstanding sensitivity, temperature resolution, and robustness in the last decade.\textsuperscript{17,18} In this regard, implementation of Ln\textsuperscript{3+}-based materials to hybrid nanostructures is a budding strategy for design of novel ratiometric nanothermometers. Among luminescent nanomaterials, photoluminescent carbon dots have emerged as important materials in sensing and cell imaging.\textsuperscript{19,20} However, most CD-based temperature sensors rely on fluorescence quenching processes, which cannot lead to accurate intracellular thermal sensing due to the influence of many factors such as the presence of metal ions, pH changes and nonuniform distribution of probes all quenching fluorescence intensity.\textsuperscript{21} All of the drawbacks mentioned above can be mitigated via introducing second not overlapping luminescent component through the built-in calibrations of two emission bands. Thus, for biological samples, which are nonisotropic, it is highly desirable to design novel bicomponent temperature sensors characterized with two well-resolved and intensity comparable luminescence peaks that possess both strong emission and submicron dimensions.
All chemical and biological processes are greatly influenced by temperature, which makes temperature measurements crucial in a big number of practical applications. Conventional temperature sensors, so-called contact thermometers, cannot remotely report the local temperature conditions. Moreover, the explosive growth of interest to micro and nano dimensional objects imposes increasingly stringent requirements in thermometric studies. Real-time response, high spatial resolution in addition to the constraint of having nontoxic and biocompatible devices makes nanothermometry a challenging area of research, in which various functional materials are involved. Luminescence-based temperature sensing is outstanding due to its merits of being noninvasive and accurate. Therefore, in terms of temperature sensing, luminescence studies of organic dyes, lanthanide-based nanomaterials, semiconductor nanocrystals, and carbon dots have attracted much attention.

Herein, we report a facile strategy for synthesis of novel hybrid polystyrenesulfonate(PSS)-coated nanoparticles consisting of carbon dots (CDs) and Tb$^{3+}$ complex as a single excited dual band luminescent nanothermometer. Ligand environment of the Tb$^{3+}$ complex should fit specific requirements, such as: (1) high complex stability, (2) incomplete coordination sphere of Tb$^{3+}$ ions, (3) efficient and temperature responsive Tb$^{3+}$-centered luminescence. The [TbL]$^+$ complex with 1,3-diketonate derivatives of calix[4]arene (L) fits well to the aforesaid requirements. The capacity of the hybrid nanoparticles to simultaneously emit in the blue and green regions of spectrum originates from two different emissive components: CDs and [TbL]$^+$ complex, correspondingly. The coordinative binding of green emitting [TbL]$^+$ complex with blue emitting CDs is highlighted as a tool for their combination within the PSS-nanoplatform as it is schematically demonstrated in Figure 1.

**Figure 1.** Schematic illustration of PSS-{CDs-[TbL]} hybrids.
The variation of the concentration conditions will be introduced as a tool to control mutual influence of the components on their sensing properties in order to develop colloids for temperature monitoring in the 25-50 °C range through the differently emitting registration channels and to transform the colloids into ratiometric temperature sensors.

**Experimental Section**

Materials: citric acid (99%), ethylenediamine (99%), Tb(NO$_3$)$_3$\(\cdot\)6H$_2$O (99.9 wt%), Triethylamine (TEA) (99%), N,N-dimethylformamide (DMF), Poly(sodium 4-styrenesulfonate) (PSS) (MWaverage = 70,000) and sodium chloride (NaCl) were purchased from Acros Organics. TRIS-buffer, acetic acid, sodium hydroxide, bovine serum albumin (BSA), DMF was twice distilled over P$_2$O$_5$. Other reagents were used without additional purification.

Tetra-acetylacetonyl derivative of calix[4]arene with hydroxyl substituents (L, 5,11,17,23-tetrakis[(acetylaceton-3-yl)methyl]-25,26,27,28-tetrahydroxy-calix[4]arene, Figure 4a) at its lower rims was synthesized in accordance with the previously reported procedure.$^{37}$

Synthesis of CDs. Carbon quantum dots have been synthesized by microwave synthesis. In a 50 ml flask, 1 g of citric acid was dissolved in 10 ml of distilled H$_2$O and 0.348 ml of ethylenediamine was added. The resulting solution was incubated for 1 minute 30 seconds in a domestic microwave oven until a red caramel residue formed. The product was dissolved in 20 ml of distilled H$_2$O, filtered from large particles through a membrane (0.22 μm) and dried at room temperature.

Synthesis of polyelectrolyte nanoparticles PSS-[TbL]. The [TbL]$^+$ complex was obtained via mixing Tb(NO$_3$)$_3$\(\cdot\)6H$_2$O (C=0.5 mM) with the ligand in a 1:1 ratio and adding triethylamine (C=3.0 mM) to deprotonate the β-diketonate groups of the calix[4]arene ligand L. PSS-[TbL] nanoparticles were synthesized by adsorption of PSS polyelectrolyte on the surface nanosized templates, obtained by precipitation of the [TbL]$^+$ complex from a DMF solution into an aqueous PSS solution (C = 1 g/L, V=2.5 ml) and NaCl (C=0,5 M). To an aqueous solution of PSS was added 0.5 ml of a complex-containing solution of DMF with vigorous stirring (2200 rpm). The resulting colloidal solution was sonicated for 30 minutes while maintaining a constant temperature of 25 °C. Excess PSS was separated by centrifugation at 4 °C and 15000 rpm for 15 minutes, followed by draining the supernatant.

Synthesis of hybrid nanoparticles PSS-{CDs-[TbL]}. Hybrid PSS-{CDs-[TbL]} nanoparticles were synthesized on the basis of a DMF solution containing a TbL and CD complex. In the Synthesis_1 concentration of [TbL]$^+$ – 4.5 mM and CD – 4.5 g/L were used. The
concentrations of the components in the Synthesis_2 were 4.5 mM and 1.5 g/L for [TbL]⁺ and CD, respectively. Corresponding DMF solution ([TbL]⁺ and CD mixture) in an amount of 0.5 ml was added to an aqueous solution of PSS (C = 1 g/L, V=2.5 ml) and NaCl (C=0.5 M) with intensive stirring (2200 rpm). PSS is wound onto the resulting nanoscale [TbL]⁺–CD precipitate with formation a stable colloidal solution. Purification of nanoparticles from excess PSS was carried out by centrifugation under conditions identical to those in the synthesis of PSS-[TbL].

The pH variations at 4.5-7.0 and 7.0-8.4 were performed by the use of acetic-acetate and TRIS- buffers correspondingly.

The UV–Vis measurements were performed on Lambda 35 spectrophotometer (Perkin-Elmer, USA) using 10 mm cuvettes at room temperature. The steady-state emission and excitation spectra were recorded on a fluorescence spectrophotometer Hitachi F-7100 (Japan) with stigmatic concave diffraction grating. Excitation of samples has been performed at 330 nm, and emission detected at 545 nm with 5 nm slit width for both excitation and emission. All samples were ultrasonicated for 15 min before measurements. The dynamic light scattering (DLS) measurements was carried out by means of the Malvern Mastersize 2000 particle analyzer. A He–Ne laser operating at 633 nm wavelength and emitting vertically polarized light has been used as a light source. The measured autocorrelation functions were analyzed by Malvern DTS software and the second-order cumulant expansion methods. The effective hydrodynamic diameter (d) was calculated by the Einstein–Stokes relation from the first cumulant: \( D = \frac{k_BT}{3\pi\eta d} \), where \( D \) is the diffusion coefficient, \( k_B \) is the Boltzmann constant, \( T \) is the absolute temperature, and \( \eta \) is the viscosity. The Nano-ZS (MALVERN) with using laser Doppler velocimetry and phase analysis light scattering has been used to determine the zeta potential. The zeta potential was measured at least three times for each sample. Only multiply reproducible results have been taken into account, thereby they differed by less than 4%. The transmission electron microscopy (TEM) images have been obtained with Hitachi HT7700, Japan. The images have been acquired at an accelerating voltage of 100 kV. Samples have been sonicated in water for 30 min and then dispersed on 200 mesh copper grids with continuous formvar support films.

**Results and discussion**

The synthesis of hybrid carbon dots-terbium chelate nanothermometer owning blue and green emitting components into the PSS-based nanoplatfrom was performed through several steps: (1) synthesis of the components; (2) evaluation of their sensing ability to heating and cooling; (3) revealing of the binding between the components; (4) optimization of conditions for the combined incorporation of both components into the PSS-nanoplatform.
Synthesis, characterization and sensing properties of CDs

Carbon dots (CDs) were synthesized according to a previously reported method in a one-step microwave assisted reaction of citric acid and ethylenediamine. Transmission electron microscopy (TEM) particle characterization shows that the dots are quasi-spherical monodisperse species with an average size of (2.3±0.6) nm remaining erratically distributed on the substrate surface (Figure 2a). The CDs show no tendency to agglomerate upon drying on the formvar coated copper grid used as a substrate.

Dispersion in water at a concentration of 10 μg mL⁻¹ results in a transparent yellowish aqueous colloids stable for at least three days (Figure S1). UV-Vis absorbance spectroscopy reveals that the water solution of CDs has one distinct absorption bands at 350 nm and 270 nm (Figure 2b) assigned to n→π* and π→π* transitions of the aromatic sp² domains, respectively.

Figure 2. (a) TEM image of dried CDs and corresponding size distribution diagram (inset). (b) UV-Vis absorption spectrum (1), room temperature excitation (2, λ_em=450 nm) and fluorescence (3, λ_ex=360 nm) spectra of a 10 mg mL⁻¹ CD aqueous dispersion.

Aqueous colloids of CDs exhibit bright blue fluorescence with the maximum at 450 nm as a result of 360 nm xenon lamp excitation (Figure 2b). The color coordinates for luminescence of CDs in H₂O are shown on chromaticity diagram (Figure 3a). CD emission spectrum was found to be dependent from temperature changes in the range of 25-70 Celsius degrees (Figure 3b). Temperature-fluorescence intensity shows linear response (R²=0.99577 for heating; 0.99548 for cooling) over the entire analysis range and the temperature sensitivity was determined to be as high as 1.25 % °C⁻¹. This value is comparable with majority of other CD-based nanothermometers.
Temperature-dependent fluorescence of the CDs aqueous colloids is fully reversible within 25-70°C temperature range (Figure 3c). The stability of the CDs colloids under the thermal cycling experiments was evaluated by measuring fluorescence intensity recovery through multiple heating and cooling cycles from 35°C to 45°C (Figure 3d). The narrowing of the temperature range to 35-45°C is aimed at physiological relevant conditions. The thermal reversibility of the CDs was revealed for at least 6 heating-cooling cycles indicating their ability to serve as a fluorescence-based nanothermometer. However, one of the issues of single-parameter thermometers is the need of an external calibration to obtain an absolute value of the temperature through the intensity measurement.

**Figure 3.** (a) Chromaticity diagram for CDs in H₂O (b). Temperature dependence of CDs emission upon heating. (c) Changes in the fluorescence intensity of the CDs (λex=360 nm) as a function of temperature over the 25-70°C range. (d) Normalized fluorescence intensity upon six heating-cooling cycles of CDs under alternating temperature of 35°C and 45°C.

**Synthesis, characterization and sensing properties of PSS-[TbL]**

As it was previously reported the synthesis of hydrophilic colloids from the terbium complexes with cyclophanic 1,3-diketonates is based on their efficient self-assembly in the synthetic conditions of the solvent-exchange technique. The [TbL]⁺ complex derives from
efficient bis-chelating of Tb$^{3+}$ via two diketonate groups of the tetra-1,3-diketone calix[4]arene (Figures 4 a,b). The 1:1 complex is formed in the DMF solutions basified by triethylamine (TEA), which is followed by the sensitization of the Tb$^{3+}$-centered luminescence due to the efficient ligand-to-metal energy transfer. Both pronounced and reversible temperature-induced changes of the luminescence of the complex in the DMF solutions (Figure S2) are the reason for its choice as the challenging building block of the PSS-stabilized colloids.

The PSS-[TbL] colloids were synthesized through the solvent exchange procedure$^{43}$ (for more details see the Exp. Section). The synthesized colloids were characterized by means of DLS data (Table 1) and TEM images (Figure 4b). The average size revealed from the DLS measurements is about 200 nm with rather low polydispersity (PDI is below 0.2). The negative surface charge manifested by the electrokinetic potential value at -20 mV confirms that [TbL]$^{+}$-based colloids are embedded into the PSS-aggregates. The difference between the average sizes revealed by TEM and DLS is in agreement with the above-mentioned assumption that PSS-based aggregates incorporate [TbL]$^{+}$-based colloids.

Figure 4. Structure of (a) L and (b) [TbL]$^{+}$.$^{37}$ (c) TEM image of dried PSS-[TbL] nanoparticles.

The steady state luminescence of PSS-[TbL] nanoparticles is manifested by four narrow bands $^{5}D_{4} \rightarrow ^{7}F_{n}$ peculiar for Tb$^{3+}$-centered luminescence giving bright green emission with the color coordinates exemplified in Figure 5a. The temperature-dependent luminescence of
polystyrenesulfonate-coated PSS-[TbL] colloids was studied for the first time here. Gradual decrease of all of four Tb bands is observed upon heating from 25°C to 70°C (Figure 5b). The band at 547 nm ($^4D_4 \rightarrow ^7F_5$) undergo highest spectral changes with the sensitivity $S_j= 3.55 \% ^\circ C^{-1}$. Cooling the sample back to 25°C results in luminescence intensity restoration to initial values. The temperature-dependent luminescence is linear ($R^2=0.97593$ for heating; 0.98803 for cooling) and fully reversible as it can be seen from the Figure 5c, where the luminescence under heating or cooling is normalized to the maximum intensity measured at 25°C. Similarly to CDs, luminescence intensity changes of 547 nm band were recorded for PSS-[TbL] within 35-45°C range. PSS-[TbL] water colloids demonstrate reversibility for at least six cycles (Figure 5d).

**Figure 5.** (a) Chromaticity diagram for PSS-[TbL]. (b) Temperature dependence of PSS-[TbL] emission spectrum upon heating. (c) Changes in the fluorescence intensity of the PSS-[TbL] ($l_{ex} =360$ nm) as a function of temperature over the 25-70°C range. (d) Normalized fluorescence intensity upon six heating-cooling cycles of PSS-[TbL] under alternating temperature of 35°C and 45°C.

Relative thermal sensitivities for [TbL]$^+$ in DMF, CDs and PSS-[TbL], in aqueous solutions were plotted in Figure 6a. The comparison indicates that the thermal sensitivity of [TbL]$^+$ up to $S_j= 7.68\% ^\circ C^{-1}$ at 50°C exhibits two-fold lowering to $S_j= 3.55 \% ^\circ C^{-1}$ under the transformation of
[TbL]⁺ to PSS-[TbL] aqueous colloids. Substantial deviation in the thermal sensitivities derives from the difference between the complexes in the DMF solutions and those in the aqueous colloids.

![Graph](image)

**Figure 6.** (a) Relative thermal sensitivity for [TbL]⁺ in DMF (1), PSS-[TbL] in H₂O (2) and CDs in H₂O (3). (b) Normalized luminescence of PSS-[TbL] versus pH.

Insignificant luminescence intensity dependence of the main 545 nm band reveals high kinetic stability of PSS-[TbL] in H₂O to pH changes in physiological range (pH=4.5-8.0) (Figure 6b). Full reversibility of Iₐₜₜ upon basification with NaOH and acidification with HCl evidence no [TbL]⁺ degradation, which makes these colloids good basis for the monitoring of temperature changes in biological liquids or tissues known to have different acidity. Luminescence λ_em=450 nm band of CDs was also found to be non-sensitive to pH changes at pH=4.5-8.0 (Figure S3).

*Interactions between CDs and [TbL]⁺ in the DMF solutions*

It is worth noting that the incorporation of the positively charged complexes into the PSS-platform is electrostatically facilitated, while negative charge of CDs is the factor raising the repulsive interactions with the negatively charges nanobeads. However, the counter-ion binding of the CDs in the basified solutions of [TbL]⁺ can facilitate their joint incorporation into the PSS-nanobeads. Thus, interactions between [TbL]⁺ complexes and nanosized carbon dots in the basified DMF solutions should be revealed as the prerequisite of their joint nanoprecipitation in the aqueous-DMF solutions of PSS. The mixing of the components in the basified DMF solutions results in the dual blue-green fluorescence under the excitation by 330 nm (Figure 7a,b). The spectra represented in Figure 7b demonstrates gradual decrease of Tb-centered luminescence intensity upon addition of CDs, which amplifies their interaction with [TbL]⁺ complex. Thus, careful analysis of the fluorescence of each component under concentration variation of another one has been undertaken and demonstrated in Figure 7c,d.
As it was previously reported, the basification is aimed at shifting the complex formation equilibrium towards [TbL]$^+$, where Tb$^{3+}$ ions are bis-chelated with two 1,3-diketonate moieties. Basified conditions facilitate deprotonation of the surface groups of CDs, which, in turn, is followed by their counter-ion binding with both triethylammonium and [TbL]$^+$ cations. The fluorescence of CDs has been studied in the DMF solutions at various concentrations of TEA and Tb$^{3+}$ ions to recognize and separate their effects. Basification of CDs DMF solution by 80 μM of TEA results in 10% loss of CDs fluorescence intensity (Figure 7c). This fact points to high deprotonation extent of various groups localized on the surface of carbon dots. The addition of Tb$^{3+}$ ions to TEA rich conditions (80 μM) reveals twofold fluorescence intensity quenching of CDs 450 nm peak, indicating efficient binding with the surface of CDs, which prevents Tb$^{3+}$ ions from instantaneous hydrolysis (Figure 7c).
Figure 7. (a) Excitation ($\lambda_{em}=547$ nm) and (b) emission ($\lambda_{ex}=330$ nm) spectra of CDs and [TbL]$^+$ in DMF at different CDs to [TbL]$^+$ ratio ($C_{CD}=0.09-3.15$ mg·L$^{-1}$; $C_{[TbL]}=0.45$ mM). (c) Fluorescence intensity of CDs (0.01 g/L) versus $C_{TEA}$ (1) and versus $C_{Tb}$ (2) in the presence of 0.08 mM TEA in DMF. (d) Intensity of 450 nm (1) and 547 nm (2) emission peaks as a function of $C_{CD}$ at constant $C_{[TbL]}=0.45$ mM. Intensity of 450 nm (3) and 547 nm (4) emission peaks as a function of $C_{[TbL]}^+$ at constant $C_{CD}=4.5$ mg·L$^{-1}$ in DMF. (e) Schematic view of Tb-L interaction exemplified by Equilibrium 1. (f) Schematic view of CD-Tb-L interaction exemplified by Equilibrium 2.

The efficient quenching of the CD-centered fluorescence by the Tb$^{3+}$ ions indicates their efficient complex formation with the chelating anions localized at the surface of CDs. Thus, the
complex formation should trigger the stripping of the Tb$^{3+}$ ions from [TbL]$^+$ complex which can be introduced schematically by the by Equilibrium 1. However, both complexing ability of the CDs and incomplete saturation of the inner-coordination sphere of [TbL]$^+$ complex allow to assume the probability of the ternary complex formation due to Equilibrium 2. Both Equilibriums are designated in schemes illustrated on Figures 7e,f. The observed changes in the Tb-centered luminescence derive from the interference of equilibriums 1 and 2, since the luminescence quenching due to equilibrium (1) can be in some extent compensated by the ternary complex formation in accordance with equilibrium (2), since the excited levels of the CDs lie above the excited Tb$^{3+}$-centered level as it is schematically shown in Figure S4.

\[
\begin{align*}
[TbL] + CD &\rightarrow L + CDTb \quad (I) \\
[TbL] + CD &\rightarrow [TbLCD] \quad (2)
\end{align*}
\]

Monitoring of the steady state and time-resolved luminescence of the complex under the varied concentration of the CDs in the DMF solutions can help in choosing optimal concentration to minimize the undesirable destruction of [TbL]$^+$ complex and to adjust the intensities of the CDs- and Tb$^{3+}$-centered luminescence on the comparable levels under the same excitation wavelength (330 nm). The observed losses in the 547 nm peak intensity of Tb-centered luminescence (Figure 7d) under the increased concentration of the added CDs points to partial degradation of [TbL]$^+$ complexes. The excited state lifetime values evaluated from the time-resolved luminescence measurements reveal their insignificant changes under the growing concentrations of CDs at constant $C_{[TbL]}=0.45$ mM (Table S1). This confirms the partial stripping of Tb$^{3+}$ ions from [TbL]$^+$ complexes by the CDs as the only reason for the quenching of the luminescence. The quenching of the CD-centered fluorescence under the increased concentration of the added [TbL]$^+$ complex also argues for the coordination of either Tb$^{3+}$ ions or [TbL]$^+$ complex with the chelating anions of the CDs. The binding with Tb$^{3+}$ and [TbL]$^+$ as the counterions should induce a surface charge neutralization of the CDs, in turn, facilitating their joint incorporation with [TbL]$^+$ complex into the PSS-based aggregates.

**Synthesis and characterization of PSS-{CDs-[TbL]} colloids**

Synthesis of the hybrid PSS-{CDs-[TbL]} colloids was performed through the solvent exchange method (see Experimental Section for more details). Variation of the components concentration, CDs and [TbL]$^+$, in initial DMF solution enables to vary the intensity ratios of 450 nm and 547 nm emission peaks upon single excitation by 360 nm wavelength, correspondingly (Figure 8). Two examples of PSS-{CDs-[TbL]} hybrids have been synthesized at various [TbL]$^+$ and CDs concentrations. The synthetic conditions 1.5 mM [TbL]$^+$ and 1.5 g·L$^{-1}$ CDs for TbL and
CD, as well as 4.5 mM [TbL]$^+$ and 1.5 g·L$^{-1}$ CDs will be herein and further designated as Synthesis_1 and Synthesis_2. It is worth noting the phase separation through centrifugation as the key step of the synthesis of PSS-{CDs-[TbL]} colloids, since the colloids of CDs can’t be separated from the aqueous solutions. Thus, the appearance of the dual blue-green emitting components in the redispersed after the phase separation aqueous colloids provides clear indication of the joint incorporation of both components into PSS-{CDs-[TbL]} colloids (Figure 8). The analysis of the dual luminescence of PSS-{CDs-[TbL]} indicates that the $I_{547}/I_{450}$ bands ratio can be varied in a wide range from 1.34 to 8.02 enabling smooth tuning of green and blue contribution. It is worth noting that the concentration variation allows to change the steady state Tb$^{3+}$-luminescence intensity with remaining the excited state lifetime values unchanged (Table S2), which supports a similarity in the interactions between CDs and [TbL]$^+$ within PSS-{CDs-[TbL]} colloids.

![Figure 8](image.png)

**Figure 8.** (a) Excitation spectra of PSS-{CDs-[TbL]} in H$_2$O, ($\lambda_{em}=450$ nm and 547 nm) (b) Emission spectra of PSS-{CDs-[TbL]} synthesized according to the Synthesis_1 and Synthesis_2
(λ<sub>ex</sub>=360 nm). (c) TEM image of dried PSS-{CDs-[TbL]} colloids. (d) Enlarged black rectangle area from the Figure 8c.

The TEM image of PSS-{CDs-[TbL]} (Synthesis_1) illustrates the presence of worm-like structures formed from globular species intertwined together and layered onto each other upon drying (Figure 8c). The more layers the darker is grey color of worm-like structures formed onto light grey plain color of the formvar grid. Figure 8d represents enlarged area of black rectangle from Figure 8c selected for detailed analysis. Black 2-3 nm dots scattered within the structural element of colloidal particles visualize the presence of CDs in PSS-{CDs-[TbL]} colloids proving hybrid nature of the latter.

Hydrodynamic diameters of PSS-{CDs-[TbL]} hybrids are higher compared to single PSS-[TbL] colloids: (285.2±1.2) nm versus (193.3±1.2) nm; however, PDI values of the hybrid and initial colloids are similar (Table 1). This, along with the higher negative value of zeta potential of the hybrid colloids versus that of the initial colloidal systems (Table 1), provide one more argument for the coexistence of CDs and [TbL]<sup>+</sup> blocks within each nanoarchitecture.

Table 1. The average hydrodynamic diameters (d), electrokinetic potential values (ζ) and polydispersity indices (PDI) evaluated from the DLS measurements of PSS-{CDs-[TbL]} (Synthesis_1).

| Name             | d, nm    | PDI      | Zp, mV    |
|------------------|----------|----------|-----------|
| PSS[TbL]         | 193.3±1.2| 0.171±0.011| -26.2±0.6 |
| PSS-{CDs-[TbL]} | 285.2±1.2| 0.258±0.015| -45.7±0.7 |
| CDs              | 1648±523.1| 0.951±0.043| -39.9±3.7 |

Sensing properties of PSS-{CDs-[TbL]}

As it was described in the previous paragraphs, both CDs and [TbL]<sup>+</sup> can give explicit luminescent response on temperature variation. Fabrication of novel composite nanomaterial out of two temperature sensitive building blocks embodies the dual band nanothermometer. Different relative sensitivities of blue (450 nm) and green (547 nm) emission bands and variable ratio between the building blocks makes it possible color tuning of PSS-{CDs-[TbL]} depending on the goal of study and complexity of an investigated system. Moreover, the coordinative binding between the CDs and the complexes is the reason for modifying both blue and green luminescent response on temperature changes. In particular, the heating of PSS-{CDs-[TbL]} (Synthesis_1) up to 70° C results in the irreversible quenching of the Tb<sup>3+</sup>-luminescence, while the temperature responsivity of the CDs components is slightly decreased, but still very strong (Figure 9a). The temperature
dependence of the dual emission is illustrated by chromaticity diagram in Figure S5. Noteworthy, the green luminescence response of PSS-{CDs-[TbL]} (Synthesis_1) is reversible within 34-45°C and can be recycled for at least 6 times at least (Figure 9b). This means that two registration channels exist for the temperature sensing with the use of PSS-{CDs-[TbL]} (Synthesis_1), which can be selected by the researcher in accordance with an object of research.

**Figure 9.** (a) Emission spectra of the PSS-{CDs-[TbL]} (Synthesis_1) (λex =360 nm) as a function of temperature over the 25-50°C range. (b) Temperature dependence of 547 nm band of PSS-{CDs-[TbL]} (Synthesis_1) upon seven heating-cooling cycles under alternating temperature of 25°C and 50°C. (c) Temperature dependence of 450 nm (1) and 547 nm (2) bands of PSS-{CDs-[TbL]} (Synthesis_2) upon two heating-cooling cycles. (d) I_{547}/I_{450} value versus temperature
within the range of 25-50 Celsius degrees. (e) Emission spectra of the PSS-{CDs-[TbL]} \((Synthesis_2)\) \((\lambda_{\text{ex}} = 360\ \text{nm})\) as a function of temperature over the 25-50°C range. (f) Normalized fluorescence intensity upon four heating-cooling cycles of PSS-{CDs-[TbL]} \((Synthesis_2)\) under alternating temperature of 25°C and 50°C.

The use of PSS-{CDs-[TbL]} \((Synthesis_2)\) enables to transform the colloids to ratiometric nanothermometers for the temperature sensing applications related to inhomogeneous systems, which are majority of biological samples (blood, cells etc.). Minimizing the content of the CDs on going from the colloids \((Synthesis_1)\) to those \((Synthesis_2)\) enables to suppress dramatically the fluorescent response of the CDs to temperature changing (Figure 9e). The binding of the CDs with Tb\(^{3+}\) ions and \([TbL]^+\) complexes in greater extent is facilitated by the conditions of \(Synthesis_2\) vs \(Synthesis_1\). In turn, the binding event is the reason for disturbance of both fluorescence of the CDs (Figure 7d) and responsivity of the fluorescence on the temperature changes (Figure 9c). The green 547 nm band in PSS-{CDs-[TbL]} colloids \((Synthesis_2)\) remains highly sensitive to the changes of temperature in the range of 25-50°C, while the fluorescence of the CDs exhibit the insignificant changes within the temperature range. This enables to correlate the intensities ratio at 547 and 450 nm with the temperature values in the range of 25-50°C, which is demonstrated at Figure 9d. The relative sensitivities \(S_j = 2.89\ \%\text{C}^{-1}\) are close to each other for the colloids synthesized in both synthetic conditions. Four heating-cooling cycles performed at alternating temperature of 30°C and 50°C indicate full reversibility of the luminescent response (Figure 9f).

It is worth noting the previously reported invariance of the luminescence signal of the PSS-stabilized colloids based on the terbium complexes with calix[4]arene 1,3-diketones\(^{44}\). This provides a background of invariant luminescence of PSS-{CDs-[TbL]} colloids in biological environment, although additional studies, including cell internalization study, intracellular optical thermometry, cell viability tests will be performed in the nearest future and reported elsewhere to outline the applicability of the produced hybrid colloids for the intracellular sensing and/or marking.

**Conclusions**

The results introduce joint incorporation of green and blue emitting blocks of different nature exemplified by \([TbL]^+\) complexes and CDs into the polysodium polystyrolesulfonate (PSS) aggregates through the solvent-exchange synthetic technique. The coordinative binding between Tb\(^{3+}\) centers and the surface groups of CDs is the main driving force of their joint incorporation into the PSS-based nanobeads. The binding between the green and blue emitting components affects their fluorescence intensity, as well as the ability to give the fluorescent response to the temperature changes. The revealed regularities both enable to develop the colloids for the
monitoring of temperature changing within 25-50 °C through blue and green emitting registration channels and to transform the colloids into ratiometric temperature sensors through the easy variation of the synthetic conditions. Novel hybrid carbon dots-terbium chelate based platform exhibiting single excited dual band emission opens an avenue for new generation of sensitive and customizable nanothermometers.

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