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A green approach to encapsulate proteins and enzymes within crystalline lanthanide-based Tb and Gd MOFs†

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In this work, bovine serum albumin (BSA) and *Aspergillus sp.* laccase (LC) were encapsulated *in situ* within two lanthanide-based MOFs (TbBTC and GdBTC) through a green one-pot synthesis (almost neutral aqueous solution, $T = 25\text{ }^{\circ}\text{C}$, and atmospheric pressure) in about 1 h. Pristine MOFs and protein-encapsulated MOFs were characterized through wide angle X-ray scattering, scanning electron microscopy, thermogravimetric analysis, Fourier transform infrared and Raman spectroscopies. The location of immobilized BSA molecules, used as a model protein, was investigated through small angle X-ray scattering. BSA occurs both on the inner and on the outer surface of the MOFs. LC@TbBTC, and LC@GdBTC samples were also characterized in terms of specific activity, kinetic parameters, and storage stability both in water and acetate buffer. The specific activity of LC@TbBTC was almost twice that of LC@GdBTC ($10.8\text{ }\mu\text{mol min}^{-1}\text{ mg}^{-1}$ vs. $6.6\text{ }\mu\text{mol min}^{-1}\text{ mg}^{-1}$). Both biocatalysts showed similar storage stabilities retaining ~60% of their initial activity after 7 days and ~20% after 21 days. LC@TbBTC dispersed in acetate buffer exhibited a higher storage stability than LC@GdBTC. Additionally, terbium-based MOFs showed interesting luminescent properties. Together, these findings suggest that TbBTC and GdBTC are promising supports for the *in situ* immobilization of proteins and enzymes.

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Introduction

Metal–organic frameworks (MOFs) are porous coordination networks constituted by metal ions bound together through organic linkers.¹ To date, MOFs have been investigated for several applications such as gas adsorption,² catalysis,³ sensing, drug delivery,⁴ removal of water pollutants,⁵ and other industrial applications.^{5–7} MOFs have also shown great potential as supports for enzyme immobilization.^{8–10} Among transition metals, lanthanides show some interesting properties like fluorescence and paramagnetism. Thus, lanthanide-based compounds have found applications in medicine for diagnostic and therapeutic purposes.^{11–15} This has suggested the potential use of lanthanide-based MOFs (Ln-MOFs) for the development of innovative biomedical applications.¹⁶ Recently, Zhang *et al.* developed a nanoprobe based

on luminescent terbium MOF-coated gold nanorods for photo-stimulated thermal therapy and chemotherapy.¹⁷ Gadolinium-based complexes (*i.e.* gadopentetate dimeglumine) are widely used as contrast agents for magnetic resonance imaging (MRI).¹⁸ Gd-based MOFs showed a longer *in vivo* circulation time and a higher MRI relaxation time, due to Gd content, higher than conventional Gd-based contrast agents.¹⁹ Other recent works report the coupling of Ln-MOFs with biomolecules,²⁰ such as enzymes/proteins or nucleic acids, to obtain hybrid biomaterials with improved properties for medical applications.^{21,22} Zhang *et al.* immobilized glucose oxidase (GOx) into an UiO type MOF modified with Eu^{3+} to obtain a fluorescence sensor for glucose detection in serum and urine.²³ Gao *et al.* encapsulated GOx and carbon dots within a Tb^{3+} based composite obtaining a hybrid GOx&CDs@AMP/Tb biocatalyst which displayed about double catalytic activity than free GOx. Moreover, GOx&CDs@AMP/Tb was further modified with carboxyphenylboronic acid to obtain a dual-emissive GOx&CDs@AMP/Tb-CPBA composite for the ratiometric sensing of glucose.²⁴ Although the coupling of biomolecules with Ln-MOFs is gaining interest, there are some synthetic issues to be solved. Indeed, enzymes and other biomolecules are generally immobilized post-synthesis by adsorption within the pores of solid carriers like ordered mesoporous silica and hierarchical zeolites.^{25–27} Since many

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MOFs are microporous (pore size < 2 nm), the post-synthesis immobilization would be only limited to the external surface of the material due to the larger size of enzyme molecules. Alternatively, the enzyme could be immobilized through encapsulation during the synthesis of the MOF.²⁸ However, such *in situ* immobilization is not applicable when typical MOF synthesis methods are used. Indeed, they require harsh reaction conditions, like high temperatures or pressures, the use of organic solvents and long reaction times (minimum 1–3 days and up to 1–2 weeks for diffusion methods).^{29–32} These synthetic methods, besides resulting in enzyme inactivation, are not environmentally sustainable. Recently, *in situ* enzyme encapsulation during the MOF formation, *via* a ‘one pot’ approach under biocompatible synthetic conditions, has been proposed.^{33,34} Employing this strategy, enzymes larger than the average pore size of MOF materials can be embedded within the three-dimensional MOF structure thus preventing enzyme leaching, while retaining catalytic activity and allowing free substrate diffusion.^{35,36} *In situ* encapsulation has been reported for both microporous MOFs, like ZIF-8,^{37,38} ZIF-90,³⁹ HKUST-1@Fe₃O₄,⁴⁰ and mesoporous MOFs, like ZIF-8,⁴¹ ZIF-67,⁴² and MIL-101(Cr).⁴³ However, the number of MOFs obtained through sustainable synthetic methods which allows enzyme immobilization is still low and mainly focused on divalent metals-based MOFs which show lower thermal/chemical stability compared to trivalent metals based MOFs.⁴⁴ On the other hand, the synthesis of trivalent metals-based MOFs, due to their higher reactivity (at room temperature) compared to divalent ones, might compromise the crystallinity of MOFs.⁴⁴ For example, Sanchez-Sanchez *et al.* carried out the *in situ* encapsulation of enzymes in a non-crystalline MOF based on Fe³⁺ and the tridentate linker trimesic acid in aqueous medium and mild conditions (room temperature and almost neutral pH).⁴⁵ Subsequently, the same procedure was used to encapsulate alcohol dehydrogenase, glucose oxidase, and lipase enzymes.^{9,10,46} Coming back to Ln-MOFs, the syntheses reported in the literature (*i.e.* for GdBTC and TbBTC MOFs) use microemulsion, mechanochemical, and solvo-thermal methods. All these approaches are not sustainable and incompatible with biomolecule encapsulation (*i.e.* solvo-thermal methods have long times and high reaction temperatures) or lead to the synthesis of amorphous materials (as in the cases of microemulsion and mechanochemical synthesis).^{47,48} Therefore, the sustainable synthesis and biomolecule encapsulation of trivalent (particularly lanthanides) metal ions-based crystalline MOFs is still a challenge.²⁸ In this work, two lanthanide ions, Tb³⁺ and Gd³⁺, were used to synthesize crystalline TbBTC and GdBTC MOFs in a sustainable way, *i.e.* using water as the solvent, under mild reaction conditions ($T = 25\text{ }^{\circ}\text{C}$ and atmospheric pressure). The same synthetic strategy was used to encapsulate bovine serum albumin (BSA) and *Aspergillus* sp. laccase (LC) within TbBTC and GdBTC MOFs. The pristine MOFs and the hybrid protein@MOFs were characterized through wide-angle X-ray scattering (WAXS), scanning electron microscopy (SEM), Fourier transform infrared (FTIR) and Raman spectroscopies,

and thermogravimetric analysis (TGA). The protein distribution after encapsulation within TbBTC and GdBTC MOFs was investigated through small angle X-ray scattering (SAXS) using BSA as a highly pure model protein. LC was used as a model enzyme to test the biological activity and stability of the LC@TbBTC and LC@GdBTC samples. To this purpose, LC@TbBTC and LC@GdBTC samples were characterized in terms of specific activity, kinetic parameters (K_M and V_{max}) and storage stability in water and acetate buffer. To the best of our knowledge, this is the first work regarding the *in situ* protein immobilization within crystalline lanthanide-based MOFs obtained through a sustainable synthetic approach.

Experimental section

Chemicals

Laccase from *Aspergillus* sp. (activity of $\geq 1000\text{ LAMU g}^{-1}$), 2,2'-azinobis-(3-ethylbenzothiazoline-6-sulfonate) diammonium salt (ABTS) ($\geq 98\%$), Bovine Serum Albumin (BSA) lyophilized powder ($\geq 98\%$), Gd(NO₃)₃·6H₂O (99.99%), Tb(NO₃)₃·6H₂O (99.99%), sodium hydroxide, NaOH; trimesic acid, H₃BTC (95%), sodium phosphate, monobasic NaH₂PO₄ (99%), sodium phosphate dibasic, Na₂HPO₄ (99%), Bradford reagent, HCl (37%), acetic acid (99%), sodium acetate (anhydrous) ($\geq 99.0\%$), were purchased from Sigma-Aldrich. All reagents were used as received without further purification. Milli-Q water ($\geq 18.2\text{ M}\Omega\text{ cm}$) was used to prepare all aqueous solutions.

Synthesis of TbBTC, GdBTC, protein@TbBTC, and protein@GdBTC samples

A mass of 22 mg of trimesic acid (H₃BTC) was dissolved in 5 mL of NaOH 1.06 M, then a solution containing 44 mg of Tb(NO₃)₃·6 H₂O (or 60 mg of Gd(NO₃)₃·6H₂O) in 5 mL of H₂O was added dropwise under magnetic stirring. The obtained white suspension was maintained under stirring at 25 °C for 1 h. The precipitate, TbBTC (or GdBTC) MOF was recovered by centrifugation at 4500 rpm for 5 min (three times), then extensively washed with water, and dried at room temperature (25 °C) under vacuum. The synthesis of LC@GdBTC and LC@TbBTC, was carried out following the procedure to obtain TbBTC and GdBTC MOFs except for the addition of 2 mL of LC solution (4.5 mg mL⁻¹) to the metal source solution. The resulting mix was added to the deprotonated BTC solution. The synthesis of BSA@TbBTC, and BSA@GdBTC were carried out as above by adding 1 mL of BSA solution (25 mg mL⁻¹) to the MOFs reaction mixture.

Characterization of MOFs and protein@MOFs samples

SEM analysis was performed by using a Gemini Ultra 55 microscope (Carl-Zeiss) with an acceleration voltage of 1 kV. Fibrils cross-section in MOFs was measured through Image J software [<https://www.nature.com/articles/nmeth.2019>]. More specifically, the cross-section of 30 fibers for each MOF was measured on SEM micrographs with a 50–300 kX magnification. The average and standard deviation are reported in the Results and

Discussion section. Attenuated total reflectance (ATR)-FTIR spectra were recorded using a Jasco FTIR 4100 instrument equipped with a PIKE GladiATR accessory with a single reflection diamond prism over the wavelength range, 4000–500 cm^{-1} . An in *Via* Qontor confocal microRaman (Renishaw) was used to acquire Raman spectra. The confocal microRaman is equipped with a solid-state laser (785 nm, IPS R-type NIR785, 200 mW, 1200 l mm^{-1} grating), a front-illuminated CCD camera (655×1024 px, working temperature -70 °C) and a research-grade Leica DM 2700 microscope. TGA was carried out by Discovery SDT 650 (TA Instruments) in a temperature range from 25 to 700 °C and a heating ramp of 10 °C min^{-1} , under continuous air flow (flow rate = 40 mL min^{-1}). SAXS and WAXS data were acquired with a Xeuss 3.0 HR (Xenocs) apparatus, equipped with a GeniX 3D X-ray generator, containing a high brightness Cu X-ray tube (30 W per 40 μm , $\lambda_x = 1.542$ Å) and a FOX 3D single reflection multilayer optic. The detector used to detect the scattering signal was a Eiger 2R 1 M hybrid photon counting detector (Dectris, pixel dimension of 75×75 μm^2). A powder/solid sample-holder was used, with Kapton tape as window (sample thickness: 1 mm). Measurements were performed at room temperature. 2D SAXS/WAXS images were collected at two sample-to-detector distances: 80 mm and 1800 mm to cover a scattering vector (Q) range of 0.0045–2.5 Å $^{-1}$ and a maximum 2θ of about 35°. After circularly averaging 2D images, 1D curves (Intensity vs. Q) were obtained. Here the scattering vector modulus, Q , is defined as $Q = (4\pi/\lambda_x) \sin \theta$, where 2θ is the scattering angle. The scattering camera was calibrated using silver behenate ($d = 58.38$ Å) as the ref. 49. Data were further corrected by empty cell subtraction and normalized, considering the measurement time and relative transmission factors. Finally, 1D SAXS/WAXS curves collected at the two different distances were merged in a single scattering profile and converted to absolute intensity, using glassy carbon as a secondary standard.⁵⁰ SAXS/WAXS data treatment was performed by XSACT software (Xenocs). The materials crystallinity was evaluated by the WAXS technique and quantified by the following equation:

$$\text{Crystallinity} = \frac{\text{Area of crystalline peaks}}{\text{Area of all peaks}(\text{Crystalline} + \text{Amorphous})} \times 100. \quad (1)$$

The fitting of SAXS curves was performed by using SASView software [<https://www.sasview.org/>]. The fitting model was chosen according to those reported in the literature for similar systems.^{51–53} More specifically, the fitting function is the sum of a $1/Q^n$ term (with n Porod exponent in the low- Q region, used only for BSA@MOFs and calcinated BSA@MOFs), a Guinier–Porod model (I_{GP})⁵⁴ and a background; the first two terms were weighted for two scaling factors, A and B , respectively:

$$I(Q) = \frac{A}{Q^n} + B \times (I_{\text{GP}}) + \text{bkg}. \quad (2)$$

The Guinier–Porod model, I_{GP} , consists of two terms:

$$I_{\text{GP1}}(Q) = \frac{G}{Q^s} \exp\left(\frac{-Q^2 R_g^2}{3-s}\right) \quad \text{for } Q \leq Q_1 \quad (3)$$

$$I_{\text{GP2}}(Q) = \frac{G}{R_g^{m-s}} \exp\left[\frac{-Q_1^2 R_g^2}{3-s}\right] Q_1^{m-s} Q^{-m} \quad \text{for } Q \geq Q_1 \quad (4)$$

with R_g the gyration radius of the scattering objects, m Porod exponent at higher Q , s a dimensionality parameter, that equals 0 for spherical objects, 1 for rods and 2 for platelets or lamellae and:

$$Q_1 = \frac{1}{R_g} \sqrt{(m-s)(3-s)/2}. \quad (5)$$

Immobilization efficiency and enzyme loading

Protein loading and immobilization efficiency of the immobilized LC@TbBTC and LC@GdBTC samples were obtained by means of the Bradford assay.⁵⁵ Briefly, the protein content was determined using the Bradford reagent (Bio-Rad) and BSA as the protein standard. The protein concentration in the supernatant was evaluated by measuring the absorbance ($\lambda = 595$ nm) of a mixture containing 0.5 mL of supernatant and 0.5 mL of Bradford reagent after 10 min of incubation. The protein loading ($L_{\text{protein}} = [\text{mg}_{\text{protein}} \text{g}_{\text{protein@MOF}}^{-1}]$) was determined by the equation:

$$L_{\text{protein}} = \frac{([P]_i - [P]_f)V}{m} \quad (6)$$

where, $[P]_i$ and $[P]_f$ are the initial and final protein concentrations (mg mL^{-1}), respectively; V is the volume of the protein solution (mL), and m is the mass (g) of the obtained protein@MOF sample.

Encapsulation efficiency (EE%) is the percent ratio between the amount of immobilized protein and the amount of protein in the immobilizing solution:

$$\text{EE}\% = \left(\frac{[P]_i - [P]_f}{[P]_i}\right) \times 100\%. \quad (7)$$

Protein loading and encapsulation efficiency for the BSA modified MOFs, BSA@TbBTC and BSA@GdBTC, were obtained by TGA. The mass loss over the range from 260 °C and 430 °C is ascribed to the combustion of BSA.

Measurement of specific activity and kinetic parameters of free LC, LC@TbBTC and LC@GdBTC samples

The LC active site consists of four copper centres, classified as type I, type II, and type III. Substrate oxidation at the mononuclear site containing the type I Cu generates electrons that are transferred to the trinuclear site, which contains type II and type III copper ions, where O_2 is reduced to H_2O .⁵⁶ The catalytic activity of free LC was determined through a Jasco 650 UV-Vis spectrophotometer by measuring the increase of absorbance, due the formation of the radical-ion $\text{ABTS}^{\cdot+}$, at $\lambda = 420$ nm for 5 min. One unit (U) of laccase activity is defined as

the amount of enzyme required to convert 1.0 μmol of ABTS to $\text{ABTS}^{+\cdot}$ ($\epsilon_{\text{ABTS}^{+\cdot}} = 34\,700\ \text{M}^{-1}\ \text{cm}^{-1}$) per minute at 25 °C. Briefly, 100 μL of commercially available *Aspergillus* sp. LC (diluted 1 : 500) were added to 2.7 mL of acetate buffer (pH 5, 100 mM) containing 150 μL of ABTS (5 mM) in a cuvette kept in the dark under stirring at $T = 25\ \text{°C}$. The catalytic activity of LC@TbBTC and LC@GdBTC was carried out following the aforementioned procedure but adding 100 μL of biocatalyst dispersion (2 mg mL⁻¹) instead of the free LC. The activity of TbBTC and GdBTC (with no LC) was carried out as a control experiment. No activity was detected for LC-free MOFs. The kinetic parameters, K_M and V_{max} , of free LC and LC@MOFs were obtained by measuring the initial reaction rate (V_0) as a function of ABTS concentration in the range 5–250 μM . A plot of V_0 ($\mu\text{M}\ \text{s}^{-1}$) versus ABTS concentration was built and the experimental points were fitted using the Michaelis–Menten model by mean of KaleidaGraph (4.5.4) software. All measurements were carried out at least in triplicate.

LC@MOFs storage stability in water and acetate buffer

Two LC@TbBTC and LC@GdBTC dispersions (2 mg mL⁻¹) in water and acetate buffer (pH 5, 100 mM) were prepared and stored in the fridge at 5 °C. At intervals of 7 days, the specific activity of both LC@TbBTC and LC@GdBTC samples was measured according to the procedure described above.

Results and discussion

Ln-MOF samples, *i.e.* TbBTC and GdBTC, were synthesized according to the “green” procedure described in the previous section. Then, two different model proteins, BSA and LC, were encapsulated *in situ* within TbBTC and GdBTC MOFs to obtain hybrid samples. Highly pure BSA ($\geq 98\%$) was used to assess protein encapsulation feasibility and to investigate the location of the immobilized protein within the MOF structures. Since protein immobilization may result in the loss of the biological function, laccase was used as a model enzyme to test the catalytic activity and stability. All samples, pristine MOFs (TbBTC and GdBTC), BSA modified MOFs (BSA@TbBTC and BSA@GdBTC), and LC modified MOFs (LC@TbBTC and LC@GdBTC) were characterized using WAXS to quantify their crystallinity. The WAXS patterns show sharp reflections at 10.3°, 13.6°, 17.7°, 19.4°, 20.9°, and 25.5° in agreement with the crystal structure of Tb(BTC)(H₂O)₆ (Fig. 1A and B).^{47,57} A similar result was reported by Yang *et al.* who synthesized the Tb(BTC)(H₂O)₆, which is isostructural with La(BTC)(H₂O)₆ (CCDC 290771†).^{58–60} The slight shift of the peaks between MOF and protein@MOF samples can be attributed to the structure perturbation caused by the protein/enzyme encapsulated within the framework. A similar phenomenon was observed by Al-Harbi *et al.*⁶¹

A rough estimation of the degree of crystallinity was obtained by mean of eqn (1).⁶² The enzyme/protein immobilization within TbBTC and GdBTC lead to a partial loss of crystallinity (Table 1). A similar phenomenon was observed by Qi

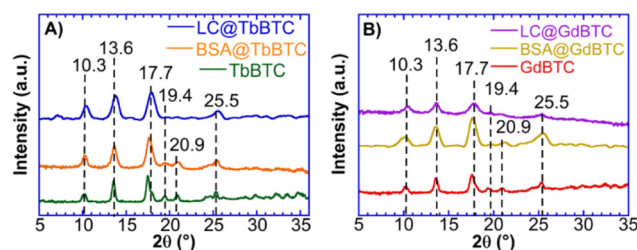


Fig. 1 WAXS patterns of (A) TbBTC, BSA@TbBTC and LC@TbBTC; (B) GdBTC, BSA@GdBTC and LC@GdBTC.

Table 1 Crystallinity (%), encapsulation efficiency (EE%) and protein loading values of MOF and protein@MOF samples

Samples	Crystallinity (%)	EE %	Loading ^a (mg g ⁻¹)	Loading ^b (mg g ⁻¹)
TbBTC	93	—	—	—
GdBTC	85	—	—	—
BSA@TbBTC	94	20.7	—	127.9
BSA@GdBTC	50	30.0	—	197.4
LC@TbBTC	70	70.2	59.3	73.2
LC@GdBTC	63	52.3	93.8	106.7

^a Determined by Bradford assay. ^b Determined by TGA.

et al. for the immobilization of *Candida rugosa* lipase within ZIF-8. Their data showed a decrease in ZIF-8 crystallinity with increasing enzyme loadings.⁶³

SEM images of TbBTC and GdBTC show a regular morphology with needle-shaped particles due to the preferred crystallization along one crystallographic direction (Fig. 2).⁶⁴ Recently, Lo Presti *et al.* found that the morphology of the lanthanide-based MOF YBTC is affected by the reaction time and temperature of the synthesis. In particular, the needle morphology was obtained for the YBTC synthesized at room temperature.⁶⁴ Owing to the encapsulation of BSA into TbBTC and GdBTC, the needle-shaped morphology changes to smaller spherical particles; the same phenomenon is observed for the immobilization of the laccase within GdBTC and TbBTC (Fig. 2). A change in morphology due to enzyme immobilization was previously obtained for *Aspergillus* sp. laccase immobilized within ZIF-zni MOF.⁸ This change might be due to the protein deposition on the MOF surface during the synthesis.⁵² However, it has also been observed that MOF morphology is affected by the type of immobilized protein. Indeed, Liang *et al.* observed the classical rhombic ZIF-8 dodecahedron crystal morphology for immobilized ribonuclease A, lipase, urease, and lysozyme, while the immobilization of ovalbumin, horseradish peroxidase, and trypsin led to leaves, flowers, and stars morphologies, respectively.⁵¹ FTIR spectra of pristine materials, BSA@MOF and LC@MOF samples are shown in Fig. 3A and B. The bands in the range of 1610 cm⁻¹–1540 cm⁻¹ and 1436 cm⁻¹–1370 cm⁻¹ are attributed to the asymmetric and symmetric C–O stretching vibration of the car-

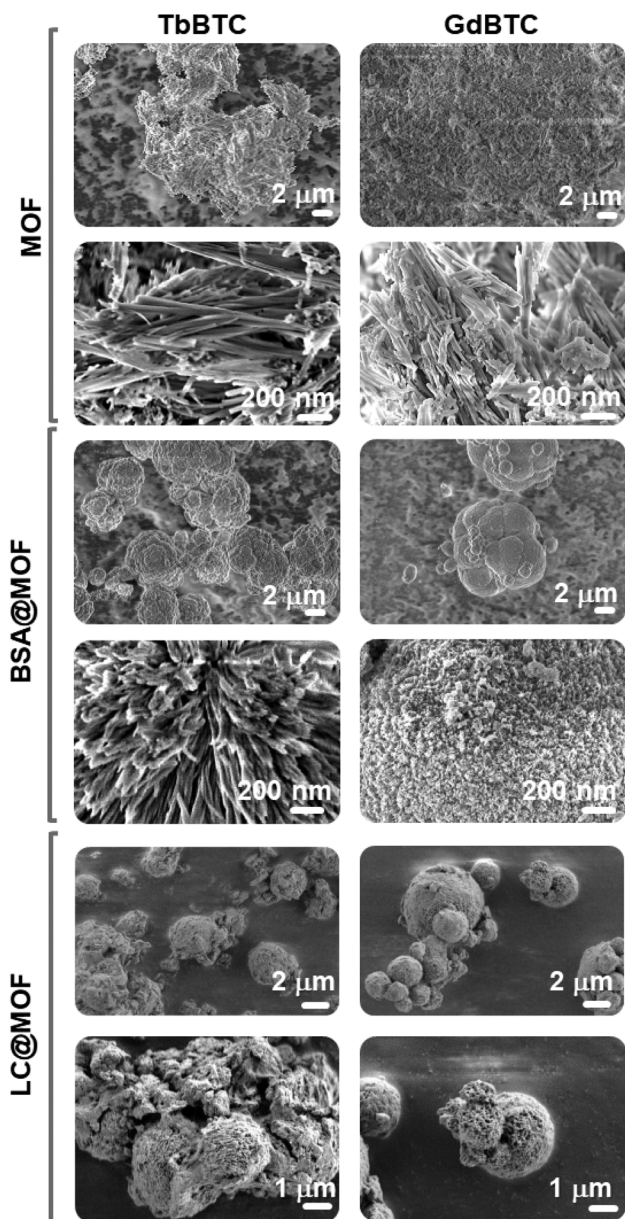


Fig. 2 SEM images of GdBTC, TbBTC, BSA@GdBTC, BSA@TbBTC, LC@GdBTC and LC@TbBTC.

boxylate group, respectively.^{65,66} The difference in wavenumber ($\Delta\nu$) between the asymmetric and symmetric stretching of the C–O gives information on the interaction between the carboxylate group and the metal ion.⁶⁷ TbBTC and GdBTC showed a $\Delta\nu$ value of $\sim 174\text{ cm}^{-1}$ and $\sim 170\text{ cm}^{-1}$, respectively, indicating that the organic linker and metal ions are coordinated *via* bidentate bridging mode.^{68–71} The two sharp peaks at 760 cm^{-1} and 710 cm^{-1} correspond to the bending of aromatic C–H bonds.^{2,72} FTIR spectra obtained for BSA@TbBTC, BSA@GdBTC, LC@GdBTC and LC@TbBTC do not show alterations if compared to pristine lanthanide-based materials demonstrating that the inclusion of LC does not alter the functional groups of the hosting MOF. The Raman spectra of

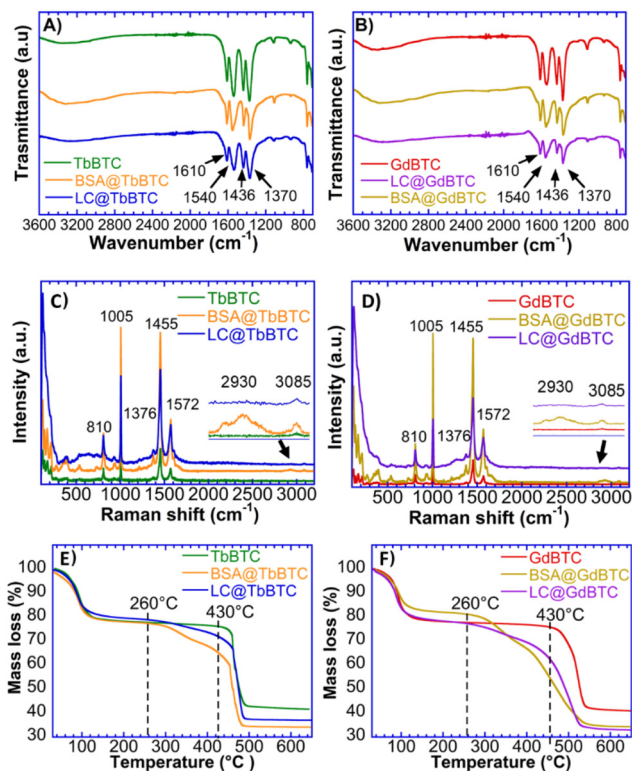


Fig. 3 Samples characterizations. FTIR spectra of (A) TbBTC, BSA@TbBTC, and LC@TbBTC (B) GdBTC, BSA@GdBTC, and LC@GdBTC. Raman spectra of (C) TbBTC, BSA@TbBTC, and LC@TbBTC; (D) GdBTC, BSA@GdBTC, and LC@GdBTC. TGA curves from 25 °C to 650 °C of (E) TbBTC, BSA@TbBTC and LC@TbBTC; (F) GdBTC, BSA@GdBTC and LC@GdBTC.

TbBTC, BSA@TbBTC, GdBTC and BSA@GdBTC samples are shown in Fig. 3C and D. Both TbBTC and GdBTC showed the same sharp bands at 536 cm^{-1} , 810 cm^{-1} , 1005 cm^{-1} , 1376 cm^{-1} , 1455 cm^{-1} and 1573 cm^{-1} assigned to the metal–O bond vibration,⁵⁷ out-of-plane CH bending, ring stretching vibrations, symmetric C–O stretching of the carboxylic group, and the asymmetric C=O stretching, respectively.⁷³ The Raman spectra obtained for BSA@TbBTC and BSA@GdBTC samples do not present substantial differences compared to the pristine Ln-MOFs, except for the Raman shift at 3085 cm^{-1} and 2930 cm^{-1} typically associated with the C–H stretching of aromatic amino acid residues, like phenylalanine or tyrosine and the symmetric stretching of CH_2 groups in aliphatic amino acid residues such as leucine, isoleucine, and valine which confirms the presence of BSA immobilization in the materials.⁷⁴ LC@TbBTC and LC@GdBTC spectra showed the same bands of BSA-modified MOFs except for the band at 2930 cm^{-1} which was not detected. This is likely due to the lower loading of laccase respect to BSA on the protein@MOF samples. Thermogravimetric profiles of TbBTC, GdBTC, BSA@TbBTC, BSA@GdBTC, LC@TbBTC and LC@GdBTC are shown in Fig. 3E and F. Both pristine materials and protein@MOF samples showed a mass loss of $\sim 15\%$ over the ranges 25 °C – 105 °C due to water desorption. TbBTC and

GdBTC exhibited a good thermal stability in air, with only a slight mass decrease between 105 °C and 430 °C. These data agree with TGA profiles of lanthanide-based MOFs obtained by green methods.⁷⁵ The final decomposition step (mass loss ~35%) occurred from 430 °C to 650 °C likely due to the decomposition of the organic moiety (trimesic acid) of the TbBTC and GdBTC. In the range from 260 °C to 430 °C, BSA@TbBTC, BSA@GdBTC, LC@TbBTC and LC@GdBTC showed a mass loss of 12.8%, 19.7%, 7.4% and 10.7%, respectively, imputable to the protein mass loss. A similar mass loss profile was observed in previous works referred to *Aspergillus* sp. laccase immobilized within FeBTC MOF.⁸ The loading of BSA and LC in the BSA@TbBTC, BSA@GdBTC, LC@TbBTC and LC@GdBTC samples quantified by TGA (mass loss in the range 260–430 °C) were 127.9 mg g⁻¹, 197.4 mg g⁻¹, 73.2 mg g⁻¹, and 106.7 mg g⁻¹, respectively (Table 1). Since the commercial *Aspergillus* sp. LC used in this work showed a low degree of purity (1% ≤ LC concentration ≤ 10%) the enzymatic loading obtained by TGA could be overestimated due to the presence of organic additives present in the commercial LC preparation.

Therefore, LC@GdBTC and LC@TbBTC loadings, were also quantified by mean of the Bradford assay. The results show a loading of 93.8 mg g⁻¹ and 59.3 mg g⁻¹ for the LC@GdBTC and LC@TbBTC samples respectively. The encapsulation efficiency (EE%), was 70.2% and 52.3% for LC@GdBTC and LC@TbBTC, respectively (Table 1).

Location of encapsulated protein within MOFs

Although protein immobilization is gaining increasing attention, particularly regarding enzymes encapsulated within MOFs, the location of the protein within the material has not been thoroughly examined. This can be done by means of SAXS technique.⁵² However, that analysis can be complicated by the fact that commercial enzyme preparations often show very low purity. Indeed, they may contain other proteins, differing from the target enzyme, coming from the biological source. Moreover, commercial enzymes are formulated to optimize catalytic activity and stability by adding additives including, carbohydrates, salts, buffers *etc.* The result is that they may have a low enzyme content with the enzyme being only a low fraction of it.⁷⁶ For these reasons, the SAXS analysis of the immobilized commercial enzymes within MOFs can be difficult. To solve these issues, BSA with high purity (>98%) was used as a model protein to investigate its location within the Ln-MOFs. More specifically, MOFs morphological changes induced by the protein loading and successive removal by calcination at 360 °C were investigated through SAXS. MOFs, BSA@MOFs and calcinated BSA@MOFs SAXS curves are shown in Fig. 4. SAXS curves were fitted according to the model reported in the Experimental section, and the fitting parameters are listed in Table 2. Pristine MOFs are characterized by inhomogeneities of 30–32 nm *ca.* (gyration radius, R_g , of 15–16 nm). Such objects size is attributable to the cross-section of MOFs fibrils, clearly visible in SEM micrographs (see Fig. 2). The average fibrils cross-section, measured

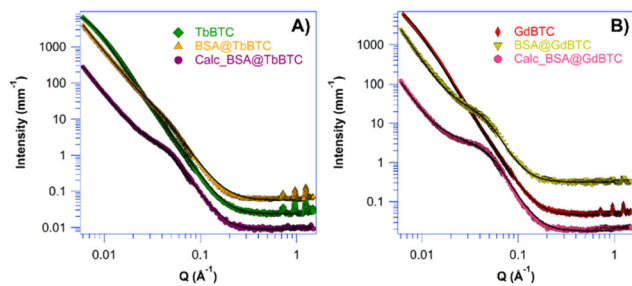


Fig. 4 SAXS profiles (full markers) and fitting (black lines) of the investigated MOFs. (A) TbBTC (green rhombus), BSA@TbBTC (yellow triangle) and calcinated BSA@TbBTC (purple circles). The curves have been shifted along the y axis for clarity. (B) GdBTC (red diamond), BSA@GdBTC (blue triangle) and calcinated BSA@GdBTC (pink circle).

through Image J software,⁷⁷ is approximately 32 ± 7 nm in GdBTC and 36 ± 9 nm in TbBTC. This is further confirmed by the SAXS fitting parameter s , that indicates the shape of the scattering objects. Being $s = 1$, the scattering signal is produced by rod-like objects. BSA loading in the investigated MOFs was found to be 197.4 mg g⁻¹ in BSA@GdBTC and 128 mg g⁻¹ in BSA@TbBTC. Therefore, the scattering signal of BSA@MOFs is expected to be mainly produced by the MOF networks. More specifically, scattering is likely produced by mesopores in which the protein is located. As a matter of fact, protein is included within the MOFs during synthesis, inducing the formation of mesopores of about 8 nm, as previously reported for other coordination networks.^{51–53} This size is compatible with monomeric BSA, with each protein molecule occupying a single mesopore.^{51,63,78} In this case the parameter s indicates scattering from globular objects, being 0–0.4 as a result of the total or partial loss of the internal elongated structure (see Fig. 2). At low Q (length scales of 50–120 nm), the Porod exponent n indicates rough surfaces of BSA@MOFs and calcinated BSA@MOFs. Surface inhomogeneities are more pronounced in BSA@TbBTC ($n = 3.3$ – 3.4), where the scattering objects are globules containing radially-arranged needles ($s = 0.44$; see Fig. 2 SEM). At high Q (length scales of 3–10 nm), $m = 3.80$ – 4.00 indicates smoother surfaces in all MOFs.

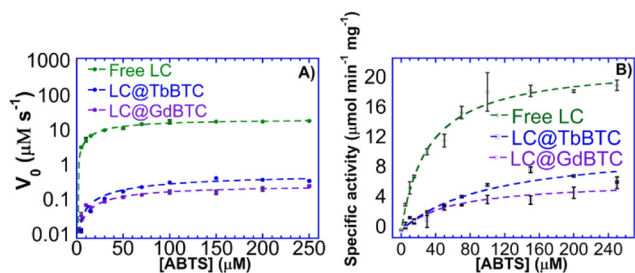
Catalytic activity and stability of the LC@TbBTC and LC@GdBTC samples

Once demonstrated that the proteins are effectively encapsulated within MOFs, it is also important to verify if they keep their biological function. For this reason, the catalytic activity of LC@MOF samples was tested. The use of enzymes immobilized within MOFs obtained by green synthesis plays a key role in the carbon-neutral economy. Indeed biocatalytic processes are green and sustainable.^{79,80} Due to their low substrate specificity and high redox potential,⁸¹ LCs can degrade several compounds possessing a phenolic structure, including lignin.⁸² Thus, LCs are promising biocatalysts for biorefinery processes. Moreover, LCs are extensively used in bioremediation processes and for biosensing.⁶ Since the laccase kinetic parameters can be affected by the immobilization process as well

Table 2 SAXS fitting parameters of MOFs, BSA@MOFs and calcinated (calc.) BSA@MOFs

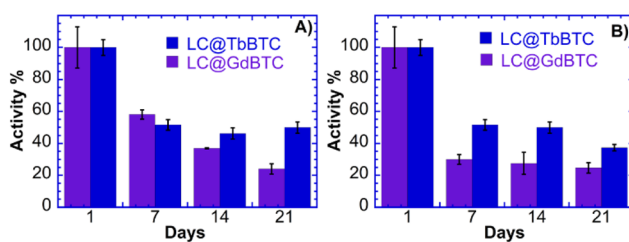
Samples	A (scale), $\times 10^{-6}$	n (Porod low-Q)	B (scale)	R_g (nm)	s	m (Porod high-Q)	bkg $\times 10^{-2}$
TbBTC	—	—	21.4 \pm 0.1	16.1 \pm 0.2	1.04 \pm 0.05	3.80 \pm 0.05	1.0 \pm 0.5
BSA@TbBTC	90.4 \pm 0.6	3.34 \pm 0.05	2.5 \pm 0.1	4.1 \pm 0.2	0.44 \pm 0.02	4.00 \pm 0.05	3.5 \pm 0.5
Calc. BSA@TbBTC	57.6 \pm 0.5	3.41 \pm 0.05	22.4 \pm 0.1	3.8 \pm 0.2	0.00 \pm 0.02	4.00 \pm 0.05	7.0 \pm 0.5
GdBTC	—	—	32.9 \pm 0.1	16.6 \pm 0.2	1.02 \pm 0.05	3.80 \pm 0.05	2.4 \pm 0.5
BSA@GdBTC	6.49 \pm 0.10	3.70 \pm 0.05	14.2 \pm 1.0	4.0 \pm 0.2	0.00 \pm 0.05	4.00 \pm 0.06	13.8 \pm 0.5
Calc. BSA@GdBTC	4.95 \pm 0.15	3.60 \pm 0.05	19.8 \pm 2.0	3.7 \pm 0.2	0.00 \pm 0.02	4.00 \pm 0.05	8.0 \pm 0.5

as by the chosen support, the effect of immobilization on the kinetic parameters of *Aspergillus* sp. laccase, used as a model enzyme, was then investigated. More precisely, the V_{\max} and K_M parameters were obtained from the Michaelis–Menten plot (Fig. 5A), measuring the initial rate, V_0 , as a function of ABTS concentration. Moreover, for a better comparison among free and immobilized enzymes, a plot specific activity vs. ABTS concentration is also shown (Fig. 5B). Both LC@GdBTC and LC@TbBTC showed an increase in K_M value, that is 1.6 and 3.0 times higher than that of the free laccase, respectively, suggesting that ABTS substrate has a lower affinity once LC is immobilized in TbBTC and GdBTC. *Aspergillus* sp. LC showed a 3.4- and 2-fold loss in specific activity once immobilized within GdBTC and TbBTC, respectively (Table 3). LC@TbBTC showed a higher specific activity (10.8 $\mu\text{mol min}^{-1} \text{mg}^{-1}$) than LC@GdBTC (6.6 $\mu\text{mol min}^{-1} \text{mg}^{-1}$). Recently, *Aspergillus* sp. laccase immobilized *in situ* within FeBTC showed a specific activity of only 0.17 $\mu\text{mol mg}^{-1} \text{min}^{-1}$.⁸ These data suggest that *Aspergillus* sp. laccase is better adapted to TbBTC and GdBTC MOFs rather than FeBTC. Similar modifications of kinetic parameters after enzyme immobilization have been previously observed. Indeed, Patil *et al.* obtained a 1.5 fold increase in K_M value, and a 1.2 fold decrease in V_{\max} due to the immobilization of the LC from *Trametes hirsuta* within ZIF-8.⁸³ Li *et al.* immobilized *in situ* a LC in a bimetallic CoCu-MOF finding that the V_{\max} decreased 3.86 times and the K_M value increased of 6 times compared to the free LC.⁸⁴ Zhang *et al.* immobilized *in situ* *Bacillus subtilis* laccase within CuBTC obtaining a K_M increase from 0.108 mM to 0.159 mM due to the LC immobilization.⁸⁵ Several factors such as the changes in protein conformation and the accessibility of the enzyme's active site, may lead to a change in kinetic parameters between free and immobilized enzymes.⁵²

**Fig. 5** Michaelis–Menten plots of the (A) V_0 vs. [ABTS] and (B) specific activity vs. [ABTS] of free LC, LC@TbBTC and LC@GdBTC.**Table 3** Kinetic parameters of the free LC, LC@TbBTC and LC@GdBTC

Samples	K_M (μM)	V_{\max} ($\mu\text{M s}^{-1}$)	Specific activity ($\mu\text{mol min}^{-1} \text{mg}^{-1}$)
Free LC	32.1	27.2	22.3
LC@TbBTC	97.9	0.71	10.8
LC@GdBTC	59.7	0.34	6.6

The large-scale application of laccases is often hindered by their low stability, which increases their cost of use.⁸⁶ Generally, compared with free laccases, immobilized laccases show higher operational and storage stabilities, moreover, immobilized LCs can be easily separated from the reaction medium.⁸⁷ The storage stability of LC@TbBTC and LC@GdBTC samples dispersed in water were hence studied. LC@TbBTC storage stability (Fig. 6A) decreased up to $\sim 50\%$ and the catalyst retained a constant activity up to 3 weeks. LC@GdBTC retained $\sim 60\%$ of its initial activity at the 7th day and about $\sim 20\%$ of its initial activity at the 21st day. A similar result was reported by Birhanlı *et al.* who encapsulated LC from *T. troglia* within cobalt- and copper-based MOFs retaining more than 58% of their initial activity over 28 days.⁸⁸ A recent interesting work by Magner and coworkers showed that buffer composition and concentration play a key role in the stability of the MOFs.⁸⁹ Their results indicated that the trimesic acid-based MOF (Fe-BTC, Cu-TMA, Co-TMA, Ni-TMA) are unstable in the presence of citrate buffer whilst the acetate is considered a promising storage medium. Hence, the stability of LC@TbBTC and LC@GdBTC stored in acetate buffer was also investigated. Both the LC@TbBTC stored in acetate buffer showed the same trend as the LC@TbBTC stored in water up to the 14th day. In the third week, LC@TbBTC stored in acetate buffer retained 37% of its initial activity, that is $\sim 10\%$ of

**Fig. 6** Storage stability (A) LC@GdBTC and LC@TbBTC; (B) storage stability on acetate buffer LC@GdBTC and LC@TbBTC.

activity lower than LC@TbBTC stored in water (Fig. 6B) whilst the LC@GdBTC stored in acetate buffer (Fig. 6B) decreased its specific activity more rapidly than LC@GdBTC stored in water (30% vs. 24% after 1 week). From the 1st week it decreased up to ~70% of its initial specific activity and retained a constant activity up to 3 weeks. LC@TbBTC showed a higher stability in acetate buffer rather than in water, whereas LC@GdBTC had an opposite behaviour. The *in situ* immobilization at room temperature within TbBTC and GdBTC MOFs, could be a valuable strategy to enhance the stability of *Aspergillus* sp. laccase. The results demonstrated that the MOF served as a protective host carrier that could maintain long storage stability.

Luminescent properties of LC@TbBTC

Besides application for green processes, enzyme@MOFs could find application in medicine. Indeed, some Ln-based MOFs display luminescent properties that have interesting potential for sensing applications.⁹⁰ Moreover some LCs (*i.e.* *A. cylindracea* and *Pleurotus cornucopiae* laccase) displayed antiproliferative activity toward cancer cells and inhibitory activity toward HIV-1 reverse transcriptase.^{91,92} Recently, Zhou *et al.* encapsulated a LC within ZIF-8 MOFs to obtain a laccase@ZIF-8-prodrug system with GSH redox cycle able to induce tumor cell apoptosis.⁹³ Hence, the properties of laccases could be combined with those of Ln-MOFs for theranostic purposes.^{94,95} In this regards, the solid-state luminescence properties of TbBTC, BSA@TbBTC and LC@TbBTC were investigated at room temperature, whereas the GdBTC MOFs were not analyzed since they are not luminescent. Upon excitation at 366 nm, both pristine TbBTC and protein@TbBTC samples show emission peaks at 488, 544, 583, and 620 nm, which could be assigned to the $^5D_4 \rightarrow ^7F_J$ ($J = 6, 5, 4,$ and 3) transitions of Tb³⁺ ions (Fig. 7). The strongest transition was the $^5D_4 \rightarrow ^7F_5$ band *ca.* 544 nm, which is responsible for the characteristic green color of Tb³⁺ emission.⁹⁶ Protein@TbBTC could be used for sensing applications. For example, Wang *et al.* developed a biosensor based on a bimetallic Tb-Fe-MOF linked with an aptamer biomolecule able to bind to ovarian

cancer biomarkers. The detection sensitivity of the obtained aptamer biosensor toward the targeted antigen was evaluated by electrochemical impedance spectroscopy, whereas the detection of living cancer cells was carried out by confocal laser spectroscopy due to the intrinsic fluorescence of Tb-Fe-MOF.⁹⁷ Although the synthesis of luminescent MOFs is widely described in the literature, luminescent MOFs are typically obtained under harsh reaction conditions posing a challenge for the encapsulation of proteins within them. For example, Chen *et al.* synthesized a luminescent TbMOF capable of selectively detecting Al³⁺ and Fe³⁺ ions in water, however their synthesis was carried out at 80 °C for 72 h under autogenous pressure.⁹⁸ The facile and green synthesis to obtain TbBTC reported in this work is more sustainable from the environmental and economic point of view compared to the conventional Tb-based MOFs synthesis.

Conclusions

In this work, BSA protein and *Aspergillus* sp. LC enzyme were immobilized within two lanthanide-based crystalline MOFs *in situ* under mild reaction conditions, that is using water as the solvent in 1 h at 25 °C and atmospheric pressure, in agreement with the 3rd, 5th, and 6th principles of Green Chemistry.⁹⁹ Pristine MOFs and protein@MOF samples were characterized employing WAXS, SEM, TGA, Raman and FTIR techniques. The enzyme location within MOFs significantly impacts their catalytic activity and stability. The immobilization of high-purity model proteins, like BSA, allows the investigation of the protein location in detail. SAXS analysis beside, confirming BSA encapsulation within MOFs structure, revealed the occurrence of mesopores of about 8 nm corresponding to the cavities created by encapsulated BSA molecules.⁵² SEM micrographs revealed a needle-shaped morphology of pristine MOFs with a cross-section of *ca.* 35 nm, also confirmed by SAXS data analysis. BSA loading changes MOFs morphology, and needle-shaped objects are partially retained only in BSA@TbBTC sample. Immobilized laccase samples, LC@TbBTC and LC@GdBTC, kept their catalytic activity although showing a slight decrease in V_{max} and an increase in K_M compared to the free LC most likely due to the mass transfer or enzyme conformational change upon MOFs immobilization. Indeed, the V_{max} as well the storage stability of *Aspergillus* sp. laccase was improved significantly upon immobilization within TbBTC and GdBTC in comparison with other MOFs like FeBTC.⁸ Furthermore, the immobilization of proteins/enzymes in a luminescent or a paramagnetic MOF, such as TbBTC and GdBTC, obtained through a 'green' approach, could lead to the sustainable development of biomedical devices. The luminescent properties of terbium-based MOFs combined with biomolecule activity suggest their applicability for theranostic purposes. For example, the laccase@ZIF-8-prodrug system with a GSH redox cycle developed by Zhou *et al.* to induce tumor cell apoptosis could be further improved by utilizing luminescent MOFs. Indeed, the intrinsic fluo-

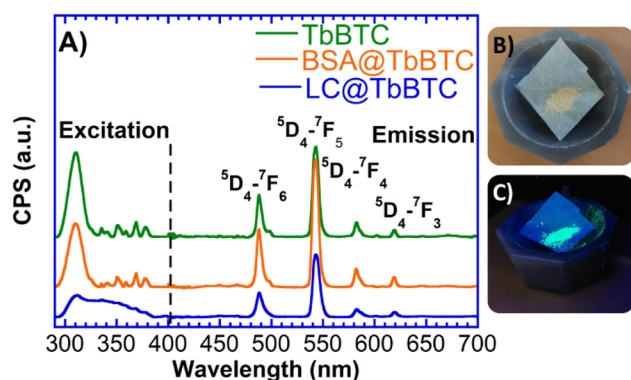


Fig. 7 (A) Solid state fluorescence spectra of TbBTC, BSA@TbBTC and LC@TbBTC samples. (B) LC@TbBTC sample and (C) LC@TbBTC sample under UV light (366 nm).

rescence of luminescent MOFs, compared to non-luminescent ones, could allow the detection of cancer cells by confocal laser spectroscopy, making the system a theranostic one.⁹³ Further work is needed to investigate the paramagnetic features and theranostic potential of enzymes encapsulated within gadolinium-based MOFs. The immobilization of proteins within crystalline MOFs synthesized under mild reaction conditions proposed in this work, paves the way for the realization of innovative devices which could contribute to the sustainable evolution of biomedical technologies.

Author contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

There are no conflicts of interest to declare.

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