Straightforward fabrication of stable white LEDs by embedding of inorganic UV-LEDs into bulk polymerized polymethyl-methacrylate doped with organic dyes

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Stable white-emitting down-converted LEDs are straightforwardly prepared by bulk polymerization of an organic dye doped polymethyl-methacrylate (PMMA) shell directly on top of a highly efficient commercial blue-emitting InGaN LED. Our optimized polymerization procedure allows for extending the form factor of achievable luminescence converter (LUCO) material beyond the conventional thin film form and to directly produce devices with light bulb design. The selected organic dyes, the blue-emitting Coumarin 30 and a red-emitting diketopyrrolopyrrole derivative, exhibit high compatibility with the free radical polymerization reaction of the PMMA matrix and ensure high stability of the final hybrid device. The control of both the thickness of the PMMA shell and the concentration of the dopant dyes allow for fine tuning of the emission color of the LUCO LED and to obtain white light with CIE chromatic coordinates x = 0.32 and y = 0.33, with rendering index as high as 80. This simple and versatile procedure is not dye-exclusive and is therefore extendable to other molecular systems for color-tunable efficient solid-state lighting sources.

The achievement of alternative lighting sources to incandescent or fluorescent lamps is a technological priority in view of a reduction of the global energy consumption. In this regard, over the last years, a massive effort has been devoted to the development of solid-state lighting devices and, in particular, white light-emitting diodes (LEDs). To date, commercially available white LEDs are fabricated using inorganic semiconductors. Despite the tremendous advancement of this technology over the years, inorganic LEDs are still expensive and limited in terms of color tunability, in the case of monochromatic LEDs, and color rendering index in the case of white-emitting devices. This is due to the fact that white light is obtained by combining narrow emission spectra which leave entire portions of the visible spectrum essentially uncovered, or through inorganic down-converters with relatively poor emission in the green and red spectral regions resulting in typically low rendering of saturated colors. In contrast, organic light-emitting diodes (OLEDs) feature high color tunability (both in terms of color coordinates and rendering index), high brightness and fast response time. Furthermore, organic materials are amenable to solution-based fabrication methods and are compatible with flexible substrates. However, the intrinsic low electrical and chemical stability of organic dyes strongly limits the lifetime of OLEDs and imposes the use of expensive preparation and encapsulation procedures that significantly raise their cost of light. The use of an efficient blue-emitting inorganic LED in combination to one or more organic luminescent converters (usually referred to as LUCO dyes) can in principle provide the best trade-off between the two technologies. In a LUCO LED, an inorganic LED performs the electroluminescence function, whilst the color correction necessary to achieve aesthetically pleasing white light is provided by organic down-converter materials featuring a high luminescence efficiency and broad emission spectra. Such hybrid strategy recently gained attention thanks to the availability of group III nitride-based LEDs that provide the blue or UV excitation source necessary to drive the white emission. Ideal organic LUCO dyes should exhibit strong absorption in the UV/blue spectral region, high luminescence efficiency and broad emission covering the whole visible spectrum. Furthermore, for the color characteristics of a LUCO LED to be stable over the whole device lifetime, it is necessary that the photochemical stability of LUCO dyes is comparable to the operational lifetime of the pump.
In this work we show that the combination of an optimized bulk polymerization procedure of PMMA and the selection of suitable fluorescent dyes - Coumarin 30 and diketopyrrolopyrrole (DPPths) shown in Fig. 1 - enable the fabrication of efficient and stable white LUCO LEDs. The described strategy is of particular practical appeal as it yields white-emitting LEDs directly in the shape of a light bulb through a fabrication protocol that can be readily scaled up to industrial mass production. This is achieved through a simple and fast immersion procedure of a commercial LED in a mold containing a partially pre-polymerized viscous mixture of MMA and fluorescent dyes. Provided that the proper mold is selected, white LEDs of any shape can be directly and cost-effectively prepared.

Results and Discussion

The main interest in the development of LUCO LEDs regards the capability to fabricate white emitters in a simple yet reliable way. In addition, this strategy offers the possibility to fine tune the emission color and rendering characteristics of an LED by controlling the thickness of the doped polymeric composite and the concentration of the dopant dyes and other quenching processes.

In particular, PMMA features good optical properties and reduced permeability to oxygen and moisture, which suppresses oxidation of the dopant dyes and other quenching processes.

PMMA is among the most commonly used polymers in applications requiring excellent optical quality and good mechanical, thermal and chemical stability. The preparation of bulk PMMA with high optical quality is usually performed through a bulk polymerization procedure, as this method does not require the evaporation of the polymerization solvent. Fabrication procedure of our LUCO LED is explained in details in the experimental section.

Photophysical properties of the LUCO dyes in solution and embedded in PMMA. Photoluminescence QYs were measured in an integrating sphere (see experimental section).

**Table 1**

| Dye                        | Emission $\lambda_{\text{max}}$ [nm] | PL Lifetime [ns] | QY    |
|----------------------------|-------------------------------------|------------------|-------|
| Coumarin 30 in hexane      | 470                                 | 2.5 ± 0.1        | 0.5510 |
| Coumarin 30 in PMMA        | 470                                 | 2.8 ± 0.1        | 0.92  |
| DPPth in dichloromethane   | 570                                 | 6.6 ± 0.1        | 0.7420 |
| DPPth in PMMA              | 580                                 | 7.7 ± 0.1        | 1.0   |
| Coumarin 30 [5·10⁻³ M]     | 470                                 | 2.7 ± 0.1        | 0.92  |
| DPPth [5·10⁻⁴ M] in PMMA   |                                     |                  |       |

Figure 1 | Optical absorption and emission spectra of (a) Coumarin 30 in hexane (black lines) and embedded in PMMA (green lines) and (b) DPPth solved in dichloromethane (black lines) and in PMMA (red lines). The chemical structures of Coumarin 30 and DPPth are reported as insets.
efficiency. Indeed, for both dyes, we observe a significant increase of the PL lifetimes from solution to the solid state (see Table 1), indicating the effective suppression of nonradiative decay channels. As a consequence, the measured QYs of Coumarin 30 and DPPth in PMMA are as high as 92% and 100%, respectively.

For the realization of a LUCO LED, we should consider that Coumarin 30 is directly excited by the employed pump LED at 405 nm. On the contrary, DPPth is nearly transparent at 405 nm and it must be indirectly excited by Coumarin 30 through two different mechanisms: (i) absorption of Coumarin 30 fluorescence, which depends on both the DPPth molar concentration and the thickness of the doped PMMA capsule, and (ii) Förster resonant energy transfer (FRET), which depends exclusively on the concentration of DPPth. The schematic representation of the LUCO LED is reported in Fig. 2a showing the InGaN LED embedded in the doped PMMA capsule and the excitation/emission mechanisms of the dyes.

The color tuning of the LUCO LED requires a proper balance of the contributions of the two dyes to the total emission spectrum. However, the control of dye concentration is not straightforward in bulk. Indeed, during the polymerization and the subsequent thermal annealing at high temperature, a fraction of the dye molecules undergoes degradation. The process is sizeable only in the presence of high temperature, residual monomer and oxygen, in agreement with what has already been observed for PMMA-based luminescent solar concentrators. We did not detect any further degradation after complete polymerization and high temperature degassing (see below). However, this degradation process prevents the fine optimization of the FRET efficiency. Therefore, we preferred to use a DPPth concentration low enough to minimize FRET and to obtain the desired emission color by varying the thickness of the PMMA capsule. The Förster radius ($R_0$) has been calculated from the overlap between the absorption and PL spectra of the dyes in diluted solution, where the respective concentrations can be precisely determined. We obtained a Förster radius of 45.1 Å, which reflects the large superposition between Coumarin 30 emission and DPPth absorption spectra and corresponds to a FRET efficiency in PMMA of 19% for a DPPth concentration of $5 \times 10^{-4}$ M (see supplementary information for details).

The effective FRET efficiency, evaluated from the comparison of the PL lifetime of Coumarin 30 in PMMA with and without DPPth, is lower than expected (in the range 0.04–0.10, considering the accuracy of the measurements). This can be explained by a small difference between the nominal and the real molecular concentrations in PMMA. Accordingly, the integrated area of the DPPth absorption band in solution is larger than that in PMMA (Fig. 1b), despite the nominal concentrations are the same. As expected, this energy-transfer rate is not sufficient to obtain white emission exclusively

![Figure 2](https://www.nature.com/scientificreports/)

**Figure 2** | Optical properties of LUCO LEDs. (a) Schematic representation of the LUCO LED showing the InGaN LED embedded in the PMMA capsule doped with Coumarin 30 (blue) and DPPth (red). (b) Simulated PL spectra of a LUCO LED as a function of the thickness of the doped down-converting PMMA capsule. Upon increasing the capsule thickness, the emission color changes from blue of pure Coumarin 30 to red of DPPth excited via FRET (grey arrow) and by reabsorption of Coumarin 30 luminescence (blue arrow). (c) CIE chromaticity plot showing the calculated color trajectory (black line) corresponding to the spectra in ‘b’. Points A and B indicate the chromatic coordinates of the PL of pure DPPth and Coumarin 30, respectively. The color coordinates of representative LUCO LEDs with increasing capsule thickness ($T = 1.3, 2.6, 4.6, 6.3$ and $8.2$ mm) are reported in circles. (d) Emission spectra of representative LUCO LEDs corresponding to the color coordinates in ‘c’. (e) Color rendering index radial plot of a LUCO LED with capsule thickness $T = 4.6$ mm. The color scheme is the same throughout the figure.
by FRET, allowing for further fine tuning of the emission color by varying the thickness of the PMMA capsule.

DPPth emission has CIE (Commission Internationale de l’Éclairage) chromaticity coordinates: \(x = 0.56\) and \(y = 0.43\) (see point “A” in Fig. 2a) and Coumarin 30 PL spectrum corresponds to \(x = 0.16\) and \(y = 0.32\) (point “B” in Fig. 2a). By mixing the two dyes, different colors can be obtained, including white light. Given the overlap between the absorption spectrum of DPPth and the emission spectrum of Coumarin 30, even at fixed dye concentration, the emission spectrum of Coumarin 30 varies with the thickness of the PMMA composite. The color trajectory achievable with our material system was calculated using the equation reported in Monguzzi et al.\(^\text{24}\), adapted for taking into account the collection geometry. Figure 2b shows the simulated emission spectra of PMMA composites with increasing thickness. The resulting color trajectory, shown in Fig. 2c, is not a straight line and crosses the pure white region (chromatic coordinates: \(x = 0.32\), \(y = 0.33\)) for a capsule thickness of 4.60 mm (crosses in Fig. 2a). Experimental PL spectra for LUCO LEDs with different thickness are shown in Fig. 2d. As a function of the capsule thickness, the chromatic coordinates span from the green/blue to the yellow/red region of the chromaticity space, in good agreement with the theoretical predictions. Only for small thicknesses we observe a minor deviation from the expected values due to the presence of residual pump light at 405 nm.

In addition to the CIE coordinates, other parameters must be taken into account for a comprehensive qualification of a lighting source: the correlated color temperature (CCT) and the color rendering index (CRI). The color temperature of artificial white light is largely a matter of personal taste and is typically associated with the requirements of specific settings. The general trend in western countries is to use cold whites for business and medical locations (e.g., offices, factories, clinics) and warm tones of white for social ambiances that prefer a comforting atmosphere, such as hotels and houses. The CCT of our devices was evaluated by using the McCamy’s approximation algorithm\(^\text{25}\), obtaining CCT = 5980 K, corresponding to cold white light. The effect on illuminated objects with this device is the enhancement of the blues and the flattening of the reds, as required, for instance, in medical environments such as hospitals. The corresponding CRI is 80 (see supplementary information), which is an accepted standard requirement for indoor lighting sources (Fig. 2e). Only for R8 (corresponding to color test TCS08 “light reddish purple”), R3 (TCS03 “yellow green”) and R4 (TCS04 “moderate yellowish green”) the rendering is not perfect, due to the relatively low emission intensity in the region centered at about 520 nm and in the deep-red tail of the visible spectrum (\(\lambda > 635\) nm).

Finally we measured the emission efficiency and operational stability of a prototypical LUCO LED. The PL QY of the down-converter material was measured using 405 nm excitation in an integrating sphere and was found to be remarkably high, with a value of 92%, which reflects the near unity PL quantum yield of our LUCO dyes in the solid PMMA matrix. With this material we realized shells for both high- and low-power commercial UV LEDs. As expected, the obtained emission spectra are essentially independent on the employed UV LED and on the driving current (see Supplementary Fig. S1). On the contrary, the output power of the pump diode determines the luminous flux of the LUCO LED that ranges from fractions of Lumen, when the converting capsule is coupled to a LE-0503-03UV (JKL Components Corporation) LED, up to 11 Lumen when the pump is a high-power Ledengin L24-00UA00 LED (see Supplementary Fig. S1).

In order to evaluate the stability of our devices, we collected the PL spectrum of a LUCO LED, excited with the JKL Components Corp pump, every hour for a total of 72 hours of continuous operation. As shown on Fig. 3, only a negligible drop of the emission is observed. This confirms the short-term stability of the dyes after the mass polymerization process of the PMMA matrix and the subsequent thermal annealing. Long-term stability has not been tested here, but probably requires the inclusion in the PMMA matrix of proper additives as usual to avoid the degradation of all UV-exposed plastic materials.

In summary, we designed and realized a hybrid white-emitting LUCO LED with high color rendering index and efficiency of over 90%. This was achieved through rational combination of an inorganic InGaN LED with two organic dyes (Coumarin 30 and DPPth) embedded in mass polymerized PMMA. Incorporation in PMMA increases both the photostability and the efficiency of parental molecules. Aging tests confirm the operational stability of both efficiency and color purity of the device over several days of operation. Our organic LUCO materials, obtained through a new fabrication protocol that can be readily scaled up to industrial quantities, could replace conventional phosphors-based down-converters used in white LEDs, thus opening the way to better performing white solid-state lighting sources.

**Experimental section**

Measurements were performed on dyes, both in diluted solution (Coumarin 30 in hexane, and DPPth in dichloromethane), and embedded in PMMA. Coumarin 30 was purchased by Sigma Aldrich (546127-100MG), while DPPth was synthesized on purpose for this study, following reference 20.

For device fabrication, we employed a two-step polymerization protocol that involves first, the formation of a viscous solution of PMMA in monomer MMA. Such solution (usually referred to as ‘syrup’) is mixed with the fluorescent dyes and casted in the mold having the required shape, where it is finally completely polymerized to give bulk PMMA\(^\text{26}\). The nominal concentrations of the dyes in the polymeric matrix are \(5 \times 10^{-4}\) M for Coumarin 30 and \(5 \times 10^{-4}\) M for DPPth. In a typical procedure, 100 mg of 2,2′-Azobis(2-methylpropiionitrile) (AIBN) were dissolved in 100 ml of freshly distilled MMA. Solutions were placed in beakers and slowly heated until the temperature of 80°C was reached. This temperature was maintained for 2 minutes, during which the viscosity increased substantially. The solution was immediately transferred in an ice bath and cooled to 20°C. A solution of Coumarin 30 (7.00 mg, 0.02 mmol), DPPth (10.80 mg, 0.02 mmol) in freshly distilled MMA (60 ml) was added along with 150 mg of lauryl peroxide. One ml of the viscous syrup thus obtained was poured in semispherical molds (1 cm diameter) and a 405 nm emitting commercial LED (model LE-0503-03UV by JKL Components Corporation andLZ4-00UA00 by Ledengin) was immersed directly in the mold without removing its protective epoxy.
capping. The molds were sealed and immersed in a water bath at 65°C for 24 h. At the end of the thermal treatment, the syrup turns into solid PMMA blocks embedding the LEDs. The devices are then removed from the mold and further cured at 75°C for 4 h. In the end of the process, devices are still perfectly functional.

A VARIAN Cary 50 UV–Visible spectrometer was used for optical absorption measurements. Steady-state photoluminescence measurements were carried out with a home-made setup, collecting the light emitted by the LUCO LEDs with a CCD coupled to a spectrometer. For time-resolved PL measurements, samples were excited at 405 nm using a pulsed diode laser (pulse duration = 40 ps) and the signal was detected by a photomultiplier tube coupled to a monochromator (time resolution better than 0.5 ns). All measurements were performed at room temperature. QYs were measured using an integrating sphere coupled to a calibrated spectrophotometer following the procedure reported in ref. 27 in order to take into account the reabsorption effects. With the same set-up we evaluated also the external emission efficiency of the LUCO LEDs. A series of automatically collected PL spectra (1 per hour for a total of 72 hours of continuous current operation of our fabricated LED) were analyzed for aging tests.

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Author contributions

D.D., L.B., F.M. and R.T. conceived the idea of direct radical polymerization of doped PMMA on InGaN LEDs. L.B. and M.S. synthesized the materials. D.D., F.M. and S.B. performed the optical experiments. D.D., F.M. and S.B. analyzed the results and wrote the manuscript in consultation with all authors.

Additional information

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