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## Which is a better fluorescent sensor: aggregation-induced emission-based nanofibers or thin-coating films?†

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**In this work, aggregation-induced emission (AIE)-based fluorescent nanofibers and thin-film coatings are fabricated using electrospinning and spin coating techniques, respectively. The two material domains are utilized as visual fluorescent sensors whose mechanism of performance is governed by the intramolecular rotation phenomenon of the AIE molecular rotors. The samples' response to humidity, temperature and organics is compared. Final results revealed that there is no all-ideal morphology. Each material works best as a visual fluorescent sensor in specific environments.**

In recent years, due to the advance in technology, several material processing techniques have emerged.<sup>1</sup> The various techniques are capable of fabricating materials with different and unique properties depending on their dimensions and sizes,<sup>2,3</sup> morphologies and structures,<sup>4–6</sup> and many other essential factors. Depending on the physical and chemical properties of the final materials (for example in polymers), after their processing, their functionality can, therefore, be modulated even with same

material type. For example, the same polymer material existing in different morphologies can show very different performances.<sup>4,7</sup> This explains why nanotechnology is at the forefront of realizing new and unique high-performance materials,<sup>8,9</sup> as it is capable of producing novel materials with various features, sizes and functions as compared to the materials obtained by traditional techniques in bulk.

The application of materials in which microscopic molecular motions can ably control the behavior of the overall material under specific environments is intriguing yet versatile. As a result, various breakthroughs have been reported utilizing this principle as in the case of molecular motors/switches/shuttles, stimuli-responsive polymers, smart fluorescent materials and many others.<sup>10,11</sup> Among them, fluorescent materials have garnered enormous attention due to their wide applications ranging from biology, security, materials science, membrane chemistry to forensic applications. It is worth noting that, fluorescent systems made up of molecular motors need the suppression of molecular motions to generate a strong fluorescence for any would-be applications.<sup>12,13</sup> This exact phenomenon is what led to the development of aggregation-induced emission (AIE).

Fluorescent AIE molecular rotors have twisted intra molecular-charge-transfer properties with the configuration of their intra-molecular motions able to determine their photophysical energy dissipation pathways (fluorescence). AIE luminogens (AIEgens) show extraordinary fluorescence at solid state due to the restriction of their rotor movements, but weak fluorescence in a solution.<sup>14</sup> Such a phenomenon is opposite to the aggregation-caused quenching (ACQ) commonly observed in traditional fluorescent molecules and materials.<sup>15</sup> These properties in AIE molecules have resulted in efforts to employ them in solid-state efficient molecular machines, sensors (probes), photodetectors, LEDs, solar concentrators and many others.<sup>16–19</sup> The current work is focused on AIE fluorescent probes realized in different morphologies.

In this work, we determine and qualify the extent of performance between aggregation-induced emission (AIE)-based nanofibers and thin-coating films as fluorescent probes. Our main aim is to understand and give an account of which AIE-based material

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system is better in these two physical forms. Briefly, we selected one typical AIEgen called tetracarboxylic-tetraphenylethylene (TPE-4COOH) which was successfully knitted onto an acrylic polymer *via* a condensation reaction. The resultant polymer-AIEgen material was further processed into fluorescent thin-coating films (FTFs) and fluorescent nanofibers mats (FNFs). With reference to the popular restriction of intramolecular rotation effect in AIEgens' intrinsic rotors, the fluorescence response properties of the obtained FTFs and FNFs were used as a guide to sensitively study the extent of performance of these materials under different environmental conditions as sensors.

*General synthesis route.* The general process of realizing FTFs and FNFs is presented in Fig. 1A. From the different acrylic resins (for acrylic synthesis see Supplementary note 1, ESI<sup>†</sup>) having the same molecular weight (but different glass transition temperature,  $T_g$ , (Fig. S1, ESI<sup>†</sup>)), one acrylic resin was selected for use in our experiments. The flexibility in  $T_g$  could be realized by adjusting the soft and hard monomers in the resin according to the Fox formula. The acrylic resin with  $T_g = \sim 30$  °C was selected as the optimum ingredient to facilitate the knitting procedure with the TPE-4COOH AIEgen in different amounts (0.05–2 wt%) (for AIEgen synthesis see Supplementary note 2, ESI<sup>†</sup>) to realize the fluorescent acrylic resin (FAR) utilized in our study. The acrylic resins not selected had either very low  $T_g$  (soft) or very high  $T_g$  (hard) and thus could not easily be appropriately used in the further process of obtaining FAR. For the synthesis of polymer-AIEgen materials (the FAR) see Supplementary note 3 (ESI<sup>†</sup>). Overall, the synthesis of AIEgen and its knitting process to the acrylic resin (such as reaction time, quantity, and conditions) were all systematically studied with the help of the restriction of intramolecular rotation mechanism which dictates the emission properties of the AIEgens (Supplementary note 4, ESI<sup>†</sup>). High emission from both the AIEgen and the polymer-AIEgen (FAR) was the target. High fluorescence of the AIEgens is expected to provide reliable

materials if made into thin-films or nanofibers for visual fluorescence sensors.

From the prepared FAR, the FNFs and the FTFs were processed. The FNFs were obtained by electrospinning (for details see Supplementary note 5, ESI<sup>†</sup>). However, since electrospinning is a very complicated process and various factors during spinning can affect the properties of the resultant fibers such as morphology, surface area, diameter, porosity, *etc.*, thus care must be taken with systematic design of experiments. In this sense, before we obtained the optimum FNFs, the spinning conditions, the weight content of TPE-4COOH,  $T_g$  value of FAR and the molecular weight ( $M_n$ ) of acrylic resin were all varied (for detailed experiments see Supplementary note 6, ESI<sup>†</sup>). Overall the optimum condition for the best-suited FNFs (Fig. 1B, top) included using FAR with  $T_g$  of 30 °C, acrylic resin with  $M_n$  of 51 200 g mol<sup>-1</sup> and the spinning viscosity of 332 mPa s at room temperature. For comparison, a flat coating based on the same FAR was prepared by spinning coating (Fig. 1B, bottom).

*Physical and optical performances.* First, the physical and optical properties of FNFs and FTFs were compared. The scanning electron microscope (SEM) images (Fig. 1B) of the representative samples detail the morphological differences between the FNFs and FTFs. As expected, the FNFs showed random nanofibers with an average diameter of  $700 \pm 30$  nm, while the FTFs showed a smooth surface. Such morphology of the FNFs presented nanopores which can uniquely improve the surface area of the material, hence more contact and active area of interaction with the environment. Unlike FTFs, no pores are observed, and this signifies a less active area. The fluorescence emission of the samples was also captured using the UV-laser-equipped fluorescence spectroscopic tool and their respective spectra recorded together with their quantum yield (QY) values (Fig. 1C, left). A relatively broad spectral peak at  $\sim 490$  nm was observed from both samples. However, the spin-coated FTFs showed a lower QY of 44.4% as compared to electrospun FNFs' 64.1%, signifying FNFs superior emission performance than FTFs. These results are also supported by the captured UV light (365 nm) images from the samples (Fig. 1C, right). Overall the excellent emission from the two materials after knitting the AIEgen is attributed to the presence of the long polymer units (51 200 g mol<sup>-1</sup>) which consequently is capable of restricting intramolecular rotations of the rotors and hence facilitating emission.<sup>17,19,20</sup> However, the difference in intensity of emission and QY between FNFs and FTFs is due to the nature of the processing techniques used to obtain each and to their physical morphologies. Electrospinning technique involves high voltages to obtain highly oriented nanofibers.<sup>21</sup> Therefore, the polymer chains in the obtained nanofibers in the FNFs samples are under high stresses due to their traction tension involved during their flow through the high-voltage electric field.<sup>22–24</sup> This consequently stretches the molecular rotors of AIEgens, which pumps the emission from the FNFs. More still, the FNFs have a high surface area and has more material being exposed to the UV excitation source as compared to FTFs.<sup>25</sup>

*Thickness-dependent emission.* In the next series of experiments, various thicknesses of FNFs and FTFs were studied, and

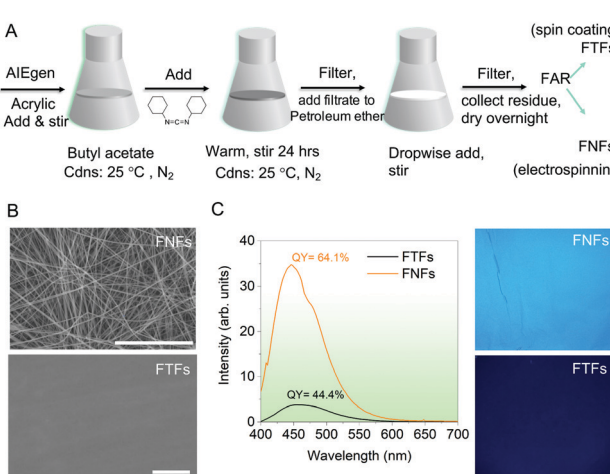


Fig. 1 Physical and optical properties of FNFs and FTFs. (A) A generalized scheme for the synthesis of FNFs and FTFs which involve first knitting the AIEgen to the acrylic polymer. (B) SEM images of the materials (scale bar is 50  $\mu$ m). (C) Fluorescence spectra captured from the FTFs and FNFs together with their QY numbers and UV light (365 nm) images.



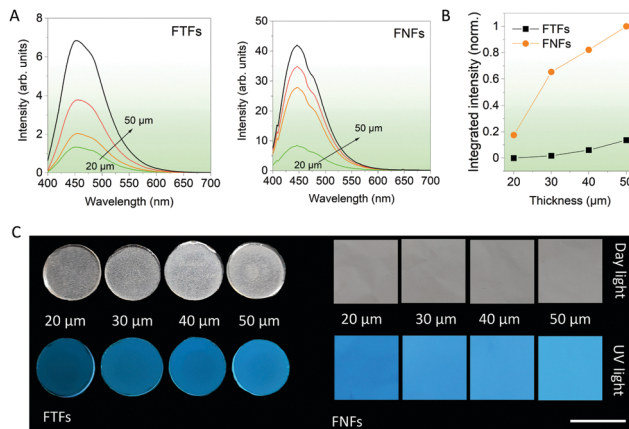


Fig. 2 Thickness-dependent emission of FTFs and FNFs. (A) Fluorescence spectra of the materials with different thickness (20–50  $\mu\text{m}$ ). (B) Summary of the spectra in (A) detailing the integrated intensity of each the samples. (C) optical and fluorescent images of the samples (FTFs and FNFs) with different thickness under room light and 365 nm UV light (scale bar is 2 cm).

their relative emission performances compared. This could help us to select the best thickness for fluorescent sensor performance comparisons in the next experiments. According to the spectral data from FTFs and FNFs (Fig. 2A), the general observed rule was “more is better”. The emission intensity of the samples generally increased with the thickness of the materials (Fig. 2B). Furthermore, the FNFs samples with 50  $\mu\text{m}$  thickness showed an excellent emission increase of  $\sim 455\%$  as compared to 20  $\mu\text{m}$  samples. In the same line, FTFs showed only  $\sim 60\%$ . The relative fluorescence images of the samples are also provided in Fig. 2C. The reason for the difference in trend between the two material samples lies in the material's AIEgen physical content properties. Samples with higher thickness have more AIEgen doped into their polymer system as compared to the thin samples. From these results, we selected the 50  $\mu\text{m}$  samples for both FNFs and FTFs for our next set of experiments. And in comparison, FNFs gave higher emission than FTFs. This explanation is clearly corroborated with the results in Fig. 1C where we observe FNFs with superior emission with a higher quantum yield. This is ascribed to the presence of high oriented polymer chains in the nanofibers in the FNFs samples which are under high stress due to their traction tension involved during their flow through the high-voltage electric field. FNFs are able to maximumly restrict the different AIE molecules in the 1D confined nanofibers space. Also, the high surface area of the FNFs is capable of interacting with to the UV excitation source as compared to FTFs.

**Response to humidity.** Now we compare the fluorescence-based sensor properties of FNFs and FTFs towards water molecules in the air. The materials were introduced to a controlled humidity environment (flow rate  $\sim 450$  sccm and relative humidity  $\sim 75\%$ ). After every interval, the fluorescence spectrum from the samples was captured (Fig. 3A). From the fluorescence spectra results, we observe that FNFs do not show any remarkable change in their fluorescence intensity. In contrast, FTFs show an increasing trend with the materials fluorescence intensity reaching maximum

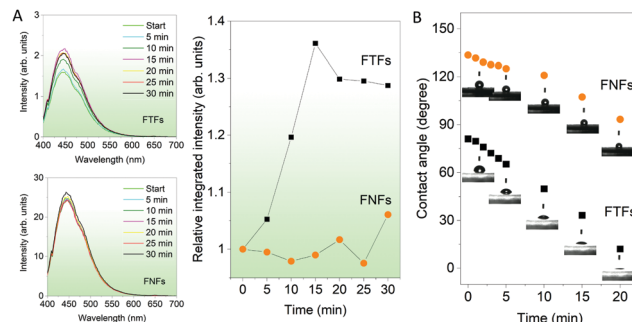
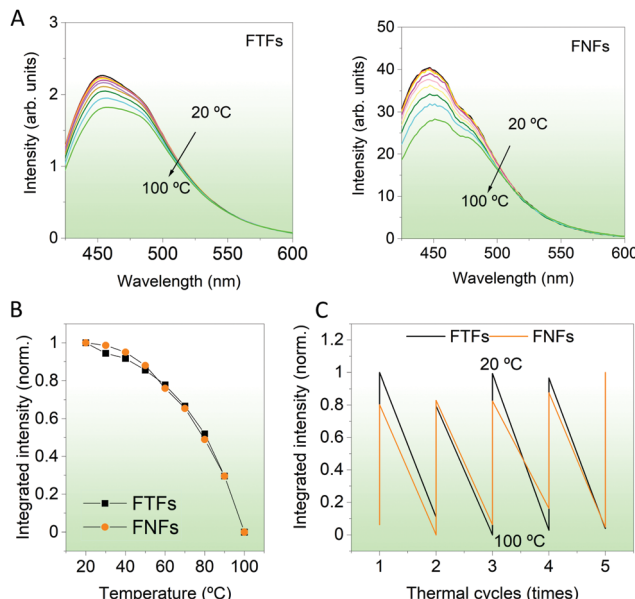


Fig. 3 Humidity response behaviors of FTFs and FNFs. (A) Effect of water molecules on the surfaces of the materials probed for half an hour in the intervals of 5 min. Left: Fluorescence spectra captured from FTFs and FNFs. Right: Summary comparing the performance response of the two materials during the 30 min duration. (B) The nature of change in the contact angle of the water molecules on the surface of the material for a 20 min duration.

intensity after 15 min. After 15 min, the intensity starts to fall progressively. Nevertheless, first, in order to account for the increased emission in FTF with humidity, the following explanation is used. Water is known to be a poor solvent for AIEgen.<sup>14,18</sup> Therefore, the intramolecular motions of TPE rotors in the AIEgen can efficiently be inhibited further with the presence of water molecules. Thus more absorbed energy can be released in the form of fluorescence radiation.<sup>26,27</sup> To explain the observations in the fluorescence intensity of the FTFs, and FNFs, we employed the contact angle measurements (Fig. 3B). The FNFs present hydrophobic material surfaces with high water contact angles (above  $100^\circ$ ). This is due to their nano-porous morphologies capable of effecting the lotus effect.<sup>28,29</sup> With the prolonged exposure of the FNFs, we observe the hydrophobicity decreasing, but overall the contact angle remained high (above  $60^\circ$ ). This means that the water molecules do not interact well with the surface and the materials, hence no change in the fluorescence. On the contrary, in the FTFs, we observe the water contact angle decreasing rapidly. Looking back at the nature of the change of the fluorescence intensity in the FTFs, we observed a rapid increase in the first 15 min, which then started to fall. In the contact angle measurements, after 15 min, we see that the contact angle was almost zero, and the FTFs surfaces acted super hydrophilic. Herein, we show observations up to 30 min because of the following reason; in FTFs the intensity remained steady with unparallel minute decreases and increases (overall, we can take it as steady) after 30 min. For FNFs the trend remained similar and steady in exact comparison to what was observed in the first 30 min. This means that the presence of water starts to deteriorate the material. Overall, it is logical to conclude that FTFs are better humidity sensors than FNFs. The FNFs cannot be used as humidity sensors. Moreover, when the FTFs are used as humidity sensors, they should be used within the active response time of 15 min or else their sensing properties start to deteriorate.

**Temperature responsivity.** Then, we compared the response of the materials to environment temperature. Temperatures between  $20\text{--}100$   $^\circ\text{C}$  were probed. During then, the FTFs and FNFs samples were exposed to different temperatures and then their fluorescence spectra were captured. From Fig. 4A, we observe the fluorescence intensity of both materials decreasing

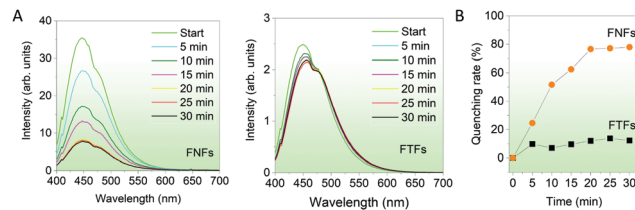




**Fig. 4** Temperature response behaviors FNFs and FTFs. (A) Fluorescence spectra of the materials when exposed to different testing temperature. (B) Summarized fluorescence results from (A) with the integrated intensity of each sample normalized. (C) Thermal recycling behaviors of the sample when continuously exposed to high (100 °C) and then cooled to low temperatures (20 °C).

progressively. The normalized integrated fluorescence data summarizing the spectral data in Fig. 4A is provided in Fig. 4B. The response of both materials to the environment temperature was similar. The decreasing trend in fluorescence of both samples is explained as follows; at low temperature (20 °C), the acrylic polymer chains are still rigid, and these chains limit the intramolecular rotation of TPE molecules,<sup>25</sup> which will then enhance the fluorescence intensity of FTFs and FNFs. However, with the increasing testing temperature, the polymer chains gradually become weak. Hence, the materials lose their originally knitted bonds between TPE and the polymer, thus promoting its intramolecular rotation. Such behavior is known to consume the absorbed energy from UV light, consequently reducing the samples' fluorescence. This is also the mechanism on which temperature sensing of AIEgens-based FTFs and FNFs work. We also performed a temperature cycling test between 20 °C and 100 °C to determine if the samples can be reused (Fig. 4C). Both samples were very stable with their fluorescence-temperature dependence response spectra being repeatable. In summary, both the FTFs and FNFs showed similar and excellent thermosensitive sensor properties with repeatable usage.

*Response to organic solvents.* Finally, we compared the fluorescent sensing properties of the FTFs and FNFs on dichloromethane (DCM) in a controlled environment. FTFs and FNFs samples were exposed to DCM for different durations (5–30 min) and their respective fluorescence spectra captured (Fig. 5A). It was observed that the fluorescence intensity of the FNFs decreased significantly with the exposure time, as compared to that of FTFs. The fluorescence quenching rate in the FTFs was only ~10% as compared to the unexposed material and kept almost stable, yet



**Fig. 5** Response behaviors of FTFs and FNFs on DCM. (A) Fluorescence spectra of the materials when exposed to DCM for 30 min, with fluorescence spectra being taken from the materials every after 5 min. (B) Summary of the quenching behaviors in (A).

for FNFs reached to 80% (Fig. 5B). The mechanism of quenching can also be substantiated with the restriction of intramolecular phenomena of AIEgen rotors. DCM is an excellent solvent and therefore, can readily interact with both the acrylic and the AIEgens. This weakens the grip effect of polymer chains with TPE molecules, hence restoring the intramolecular motions of AIEgens, which consequently results in the massive consumption of the absorbed energy from UV light and thus reduced fluorescence. The different response between FNFs and FTFs to DCM is ascribed to the FNFs nanofibrous porous morphology and high specific surface area which benefit the rapid penetration of DCM. However, for the FTFs, the reduced surface area tends to restrict DCM interaction with material and hence the same trend across all the reaction times. This also explains why in FNFs after 20 min, the fluorescence becomes stable, signalling saturation point of the FNFs samples, where they start to act as thin films. Overall, FNFs are excellent organic liquid sensors than FTFs in this regard.

In summary, AIE-based polymers are excellent visual fluorescent sensors whose working principle is governed by the AIEgen rotors' intramolecular rotation, which changes with the material environment. The change in material's morphology such as thin films, nanofibers (as in the case in this article) consequently changes the responsivity of the final material to a specific environment, rendering the same material species active or inactive (depending on the chemical and physical means the AIE rotors are affected). In this work, we have demonstrated that FTFs are better humidity sensors than FNFs, whereas FNFs are excellent organic liquid sensors than FTFs. Moreover, both the FTFs and FNFs show similar and excellent thermos-sensitive sensor properties with repeatable usage. It is therefore paramount to understand which response is needed and the tailor the existing AIEgen-polymer system chemically and physically using the different material and chemical processing techniques. There is no ideal morphology or approach, and performance lies in the art to play with the rotors.

## Conflicts of interest

There are no conflicts to declare.

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