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Aggregation-induced enhanced emission (AIEE), pH sensing and selective detection of sulfuric acid of novel imidazole-based surrogates made *via* microwave-assisted synthesis[†]

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A novel series of imidazole derivatives (4a-d) was synthesized via a microwave-assisted synthesis and whose structures were elucidated using ¹H and ¹³C NMR, ESI-HRMS, and FT-IR spectroscopy. Photophysical characterization revealed absorption peaks around 305 nm and 327-365 nm, with strong photoluminescence (PL) emission maxima ranging between 435 and 453 nm. Aggregation-induced enhanced emission (AIEE) behavior was observed in THF/H₂O mixtures, where 4a-c showed maximal fluorescence at certain solvent ratios, indicating aggregate formation. 4d disclosed a wavelength shift from 408 to 460 nm along with an enhanced emission, which is attributed to restricted intramolecular rotation (RIR), as confirmed by viscosity studies. Fluorescence lifetime and dynamic light scattering (DLS) measurements further supported the aggregation process, with particle sizes between 100 nm and 720 nm. Density functional theory (DFT) calculations validated electronic conjugation, showing HOMO-LUMO bandgaps (ΔE) of approximately 2.01–2.23 eV for 4a–d. Compound 4a exhibited the largest HOMO-LUMO energy gap of 2.23 eV, indicating greater electronic stability and enhanced emission efficiency upon aggregation. In contrast, compound 4d, with the smallest energy gap of 2.01 eV, suggests higher reactivity and better sensitivity to aggregation phenomena. This characteristic renders 4d particularly advantageous for sensing applications where rapid responsiveness to environmental variations is critical. pH sensing studies demonstrated the stability of 4a-d over a broad pH range, with 4d showing a 47 nm red shift in highly acidic conditions besides a selectivity for sulfuric acid detection. Investigation of sulfuric acid detection limit was studied, revealing a capacity for 4a-d to detect an acidic concentration as low as 16.5 µM. Stability and practical applicability of 4d compound as a sensor are further confirmed through reversibility and repeatability tests which reveal the possibility to regenerate the imidazole derivative even after several uses.

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Introduction

Imidazole derivatives, which are considered among the most nitrogen-abundant heterocycles, are commonly found in natural products and pharmaceutical active ingredients.^{1,2} These compounds are also conspicuous for their biological activities allowing for their usage in myriad applications, among others, anticancer, antimicrobial, antihypertensive, and protein kinase inhibition.³⁻⁶ In addition to their biological properties, imidazole surrogates were reported as prominent light emitters, promoting them for potential applications in myriad fields, namely, metal sensing,^{7,8} biological imaging,⁹⁻¹¹ LysoTracking,¹² and as components in organic light-emitting diodes (OLEDs).¹³⁻¹⁵

Aggregation-induced enhanced emission (AIE), a riveting photophysical phenomenon, was first discovered and presented by Tang *et al.* in 2001.¹⁶ Since then, AIE has garnered increasing consideration for the development of efficient fluorogens, given its unique characteristics where, as opposed to traditional molecules that exhibit aggregation-caused quenching (ACQ), AIE fluorogens demonstrate weak or no emission in solution due to intramolecular motions but portray a pronounced fluorescence upon aggregation due to the restriction of these

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motions.¹⁷ On the other hand, aggregation-induced enhanced emission (AIEE) is a photophysical phenomenon where aggregates of a given emissive compound display an increased fluorescence intensity.¹⁸ Both AIE and AIEE fluorogens are bestowed by their structural modularity, emission tunability, and biocompatibility, therefore, leading to their use in optoelectronic devices,¹⁹ bioimaging,^{20,21} stimuli-responsive materials,²² biological applications,²³ and chemical sensing.^{18,24} The latter application has witnessed extensive research to develop AIEE sensors of various ions and molecules,²⁴ aiming to make chemosensors with high selectivity, cost-effectiveness, and ease of preparation remains an exciting area of research.²⁵

Since Sørenson's first pH measurement in 1909, the quest for more selective proton detectors continues to intrigue chemists.²⁶ pH is a crucial parameter in both chemical and biological systems, with enzymatic reactions and cellular buffering systems operating within narrow pH ranges.²⁷ Deviations in pH can signal conditions like the acidic environment of cancer cells, consequently, making pH monitoring critical in biological research.²⁸ Advances in the design of novel pH sensors have paved the way for the synthesis of intricate bioengineered and nanosized systems capable of non-invasive, *in situ* analysis.²⁹

Additionally, sulfuric acid has a wide range of industrial applications, with its primary use being in the production of fertilizers.³⁰ This oxidizing agent is also employed as a key reagent in the synthesis of other chemical compounds, such as, hydrochloric acid, nitric acid, sulfate salts, detergents, and pharmaceuticals.³¹ Due to its extensive usage, the precise detection of sulfuric acid deems essential to monitor the environment, maintain the product quality and ensure safety during industrial processing. Recent advancements in optical fiber technology has significantly improved chemical sensing, making them invaluable sulfuric acid detectors.^{32,33}

In this work, we report the synthesis of a new series of imidazole derivatives **4a–d**, which were made by reacting a specially designed benzil derivative, decorated with lateral bulky groups **2**, with various commercially available mono- and di-benzaldehydes by a simple microwave-assisted one-pot synthesis. **4a–d** were subsequently explored as AIEE fluorogens, pH fluorosensors, and selective sensors of sulfuric acid. To the best of our knowledge, this is the first report which explores the employment of such imidazole compounds bearing bulky lateral groups as AIEE fluorogens and chemical sensors.

Results and discussion

Synthesis

Target imidazoles **4a–d** were made by attaching the 4-*tert*butylphenylacetylene **1** group to the extremities of the commercially available 1,2-bis(4-bromophenyl)ethane-1,2dione (DBB), through a dual Pd-catalyzed Sonogashira crosscoupling reaction, yielding 1,2-bis(4-((4-*(tert*-butyl)phenyl)) ethynyl)phenyl)ethane-1,2-dione **2** with an excellent yield of 90% (Scheme S1 of the ESI file†).^{34,35} Subsequently, a microwave-assisted, catalyst-free, one-pot reaction between compound **2** and the commercially available mono- (**3a-b**) and di-aldehydes (**3c**) at 180 °C in the presence of ammonium acetate (NH₄OAc) afforded the target imidazoles **4a–d** within 10 minutes of reaction (Scheme 1). **4a–d** exhibited high solubility in common organic solvents, including chloroform, DCM, and THF, which allowed for their seamless structural analysis using ¹H- and ¹³C-NMR spectroscopy, high-resolution mass spectrometry (ESI-HRMS), and FT-IR spectroscopy (see Fig. S1–S5, S6–S10, S11–S15 and S16–S20 in the ESI file†).

The ¹H-NMR spectra of **4a–d** exhibit a distinct, high-intensity signal in the aliphatic region between 1.28 and 1.36 ppm, thus indicating the presence of the *tert*-butyl groups. The proton signals of the aromatic regions for **4a–d** appear in the range of 7.95 and 7.32 ppm (see Fig. S1–S5 in the ESI file†). ¹³C-NMR spectra disclose the aromatic carbon signals for **4a–d** between 151.85 and 119.97 ppm, with additional characteristic signals at ~88.65, ~34.7 and ~31.1 ppm, which are attributed to the fingerprint peaks of sp and sp³ hybridized carbons of the alkynyl and tertiary butyl groups, respectively (see Fig. S6–S10 in the ESI file†).

Electrospray ionization high-resolution mass spectrometry (ESI-HRMS) further confirms the formation of **4a–d**, divulging the exact mass peaks for **4a–d** which correspond to the calculated values for each compound (see Fig. S11–S15 in the ESI file†). FT-IR spectra of **4a–d** also exhibit the anticipated stretching and bending vibrations for the key functional groups, including C=C, (2000–2219 cm⁻¹) C–H, (2951–2958 cm⁻¹; 828–836 cm⁻¹ ben) C=C, (1488–1514 cm⁻¹) and C–N, (1102–1105 cm⁻¹) therefore, further substantiating the successful synthesis of the target compounds (see Fig. S16–S20 in the ESI file†).

Photophysical properties

The photophysical properties of compounds **4a–d** were examined using UV-vis absorption and emission spectroscopy in THF as a solvent. Fig. 1 portrays the absorbance of compounds **4a–d** which display two peaks, the first ranging between 300 and 305 nm, whereas the second is detected in the range of 327–365 nm and which corresponds to the π - π * transitions of the target compounds' aromatic cores.³⁶

Photoluminescence (PL) measurements of **4a–d** were conducted in the same solvent portraying similar emission peaks for **4a–c** with a λ_{max} in the range of 428 to 435 nm. Interestingly, **4d**, which contains a diimidazole core, exhibits two distinct emission peaks at approximately 428 and 453 nm.³⁷

Aggregation-induced enhanced emission (AIEE) studies

The newly synthesized imidazole derivatives **4a–d**, featuring multiple rotational centers, are anticipated to exhibit aggregation-induced enhanced emission (AIEE) due to revolving restriction of the peripheral phenyl rings, which diminishes luminescence in solution.³⁸ Compounds **4a–d** are highly soluble in THF but insoluble in water thus making THF/ H_2O mixtures ideal for exploring their AIEE behavior (see Fig. 2 and S21–S23 in the ESI file†). Solutions of **4a–d** in neat THF show weak emission peaks but whose fluorescence intensities increase as the water fraction (f_w) soars, suggesting the





Scheme 1 Synthesis of 4a-d.

formation of aggregates prompted by the addition of water (see Fig. 2 and S21–S23 in the ESI file†). While **4a** portrays its fluorescence maximum intensity at a THF/H₂O ratio of 40 : 60, **4b**,**c** disclose it at a ratio of 30 : 70, before the emission starts to diminish at higher f_w . The enhancement in photoluminescence (PL) intensity at 60–70% f_w confirms the AIEE effect for targets **4a–c** and which could be explained by the interlocked networks formed by the rotationally restricted aromatic groups through the pending lateral groups linkers upon aggregation.³⁹ It is worthwhile to mention that the decline in PL intensity for **4a–c**

when $f_w > 60\%$ or 70% can be explained by two factors: (i) the formation of large aggregates where only the molecules present at the surface emit light upon excitation, thus, resulting in the decrease of the overall fluorescence intensity,⁴⁰ or (ii) the solute molecules aggregate in amorphous nanoparticle forms, therefore, leading to a reduction in the emission intensity, unlike the enhancement observed with the formation of crystalline nanoparticles.⁴¹ However, in instances where higher f_w did not prompt precipitation, this could be explained by the formation of aggregates whose size remains at the nanoscale level.



Fig. 1 Normalized UV/vis absorption (A) ($C_{\rm M} = 10^{-6}$ M solid lines) and emission (B) ($C_{\rm M} = 10^{-7}$ M dashed lines) spectra of compounds 4a– d recorded in THF (absorption maxima were used as the excitation wavelengths).

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Fig. 2 (A) Emission spectra of 4d in THF/water mixtures (0–100%); (B) plot of maximum emission intensity of 4d versus water fraction; (C–F) dynamic light-scattering (DLS) spectra of 4a-d in THF/water, at specific f_w ratio.

Dynamic light scattering (DLS) analysis of **4a–c** in THF/H₂O, specifically in solutions **4a** with f_w of 60% and 90%, **4b** 70% and 90%, and **4c** 70% and 90% discloses average aggregate sizes of approximately 100 d.nm and 240 d.nm, for **4a**, 380 d.nm and 720 d.nm, for **4b**, 537 d.nm and 720 d.nm, for **4c** (Fig. 2C–E). By correlating these findings with the change in the emission

intensity observed (*c.f.* Fig. S21–S23 in the ESI file[†]), it is evident that the reduction in emission at f_w of 90% is attributed to the fact that only the molecules at the surface of large aggregates contribute in emission.⁴²

On the other hand, the fluorescence intensity of compound 4d has gradually increased from 0% to 90% in THF/H_2O mixture



Fig. 3 (A) Powder XRD pattern of the synthesized (black graph) and THF/water, f_w 90% (blue graph) of 4d and (B) fluorescence lifetime decay profiles of 4d in THF/water, f_w %; 0 (red graph) and 90 (blue graph).

in addition to the emission wavelength shifts from 408 nm to 460 nm (Fig. 2A and B) This suggests that the mechanism of the AIE activity of **4d** is mainly through restricted intramolecular rotation (RIR) which was further supported by recording the emission spectra of **4d** with different mixtures of THF and high

viscous solvent polyethylene glycol (PEG) 0–90% (with a fixed amount of THF and a gradually increasing amount of PEG) (see Fig. S24 in the ESI file†). The emission intensity was gradually enhanced by increasing the viscosity. The viscosity experiments indicates that AIEE in **4d** is due to the RIR mechanism.^{43,44}



Fig. 4 The optimized structures and molecular orbital amplitude plots of HOMO and LUMO energy levels for 4a-d computed using the B3LYP/ 6-31G* basis set.



Fig. 5 Emission spectra of 4d (1 \times 10⁻⁷ M) in different THF-buffer mixture, 9:1, v/v, λ_{exc} = 365 nm.

Dynamic light scattering (DLS) analysis of 4d in THF/H₂O, with f_w of 10% and 90%, discloses average aggregate sizes of approximately 505 d.nm and 177 d.nm, respectively (Fig. 2F).

Furthermore, powder X-ray diffraction (PXRD) of **4d** as synthesized and that of the 90% water fraction solutions were investigated where an aliquot of the latter was centrifuged and filtered before collecting the residue which was collected and then characterized. PXRD analysis revealed broad peaks for **4d**, indicating its amorphous nature, contrarily to the sharp and intense diffraction peaks for the isolated sample of **4d** from solution, thus, indicating microsegregation (Fig. 3A), and which further supports the RIR effect leading to an increase in the PL intensity with an increasing water concentration.

The lifetimes of compounds **4a–d** were measured at room temperature employing time-correlated single photon counting (TCSPC). Samples **4a–c**, prepared in THF/H₂O mixtures with

concentration ratios of 40:60, 30:70, and 30:70, respectively, were found to have determined lifetimes of 1.48, 1.46, and 1.14 ns, respectively (see Fig. S25 in the ESI file†). In contrast, the lifetime of **4d** measured in pure THF was 0.32 ns, while a notable increase to 1.05 ns was observed for the same derivative in a 10:90 THF/H₂O mixture, which undoubtedly indicates enhanced luminescence yield due to aggregation effects when compared to **4d** solution in pure THF (Fig. 3B).⁴⁵

Electronic structure

Computational optimization of **4a–d** using B3LYP/6-31G* basis set disclose similar orbital distribution of their HOMO–LUMO energy levels with electron conjugation majorly located in the benzene rings connected through the alkynyl bonds at the HOMO level, and which becomes more relevant in the remaining part of the target compound at the LUMO level. All



Fig. 6 (a) Emission spectra of 4d (1×10^{-6} M in THF) in various acid solutions (excitation wavelength: 365 nm). (b) Linear correlation of I_0/I vs. the concentrations of H₂SO₄ in the range 0.5 × 10⁻⁵ to 1.75 × 10⁻⁴ M.



Fig. 7 (a) Emission spectra of 4d (1×10^{-6} M in THF) in different acidic and basic solutions (excitation wavelength: 365 nm). (b) Change of the emission intensity at ~450 nm upon adding H₂SO₄ and NaOH (excitation wavelength: 365 nm).

the structures were found to be deviated from planarity, especially at the aryl-imidazole connection by a dihedral angle of around 6.2° to 6.3°, which prevents their effective conjugation (Fig. 4). Examination of the HOMO, LUMO, and energy variance values for the four samples 4a-d offers valuable insights into their properties as prominent Aggregation-Induced Emission (AIE) materials, with HOMO energies ranging from -7.99 eV to -7.77 eV, thus indicating that these compounds possess relatively high energy levels for the highest occupied molecular orbital. In contrast, the calculated LUMO energies were found at -5.76 eV to -5.77 eV. Therefore, the HOMO-LUMO energy differences are found in the range of 2.01 eV to 2.23 eV. Sample 4a has disclosed the largest HOMO-LUMO gap of 2.23 eV, suggesting larger stability and enhanced emission properties upon aggregation,⁴⁶ as such a large gap is expected to reduce non-radiative decay thus allowing for more efficient luminescence, a critical characteristic of AIE materials.⁴⁷ On the other hand, 4d which exhibits the smallest energy difference of 2.01 eV, may imply a higher reactivity or sensitivity to aggregation conditions,48 which could be advantageous for sensing applications where responsiveness to environmental changes is crucial. The slightly reduced energy gap in this sample may facilitate easier electronic transitions, allowing for real-time monitoring and detection capabilities.

pH sensing studies

Given that most imidazole derivatives are sensitive to acids,⁴⁹ fluorescence pH sensing studies of **4a–d** $(1 \times 10^{-7} \text{ M})$ were conducted in THF-buffer mixtures (9:1, v/v) whose pH range between 0.5 and 12.^{49,50} When excited at 365 nm, **4a–c** exhibited emission bands at 435 nm, 432 nm and 430 nm, respectively, across the pH range of 1–12. Alternatively, **4d** portrayed two emission bands at 428 nm, and 450 nm, with the former maintaining this emission maximum at a pH range of 1–12 while the latter keeping its fluorescence at pH values in the range of 2 to 12. Interestingly, the emission spectra of

compounds **4a–c** were red-shifted by up to 40 nm at a pH of 0.5, with quenching up to 44%. Whereas that of **4d** exhibited a more significant red shift of 47 nm with quenching up to 40% at pH values of 0.5 and 1.0 with a conspicuous color change from blue to pale yellow under UV illumination (Fig. 5). It is noteworthy that the hitherto mentioned observed red shifts are likely due to the protonation of the imine group in the imidazole skeleton.⁴⁹ These observations clearly demonstrate that compounds **4a–d** serve as effective fluorosensors at pH values ranging from 0.5 to 1 (see Fig. 5 and S26–S28 in the ESI file[†]).

Specific acid sensing studies

Fluorescence acid sensing experiments were performed using a 1 \times 10⁻⁶ M THF solution of 4d with 10 equivalents of four different acids, notably, trifluoroacetic acid (TFA), hydrochloric acid (HCl), acetic acid (AcOH), and sulfuric acid (H₂SO₄). A 1:1 (v/v) mixture of 4d and a given acid was excited at 365 nm, portraying no change in the emission of 4d upon using the monoprotic acids (TFA, AcOH, and HCl), unlike the diprotic sulfuric acid (H₂SO₄) which caused the emission of 4d to be red shifted by 47 nm, additionally a hypochromic shift (Fig. 6a) strongly suggests an interaction between the latter and H₂SO₄ attributed to the protonation of the imine moiety of the imidazole.51 Quantitative detection of H2SO4 was investigated employing the limit of detection (LOD), which was determined using the formula $3\sigma/K$,^{52,53} where σ represents the standard deviation obtained from 10 consecutive scans of blank samples whereas K is the slope of the calibration curve. Therefore, titration studies of 4d with H₂SO₄ were carried out utilizing a 1 \times 10– 3 M stock solution of 4d in THF and aqueous $\rm H_2SO_4$ solution concentrations ranging from 0.5 imes 10⁻⁵ M to 1.75 imes 10^{-4} M. During the titration, 50 µL of 4d and 5 µL of H₂SO₄ solutions were added and diluted to a final volume of 2 mL, revealing a decrease in the emission intensity of 4d with an increasing concentration of H₂SO₄, as expected (see Fig. S29 in the ESI file[†]). The calibration curve was obtained by plotting the

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changes in I_0/I ratio at ~450 nm against the corresponding H₂SO₄ concentrations allowing for the calculation of the LOD and which was found to be 16.48 μ M (1.62 ppm, Fig. 6b).

The stability and reproducibility of **4d** were evaluated using H_2SO_4 and NaOH. Thus, reversibility experiments were carried out by adding H_2SO_4 (1.5×10^{-4} M) and NaOH (3×10^{-4} M) successively to a 1.5×10^{-5} M solution of **4d** over four cycles and recording the fluorescence spectra during each addition step (Fig. 7). The results indicate that **4d** exhibits reversible behavior in the presence of H_2SO_4 and NaOH, thus confirming its stability.

Conclusion

In summary, a novel series of imidazole derivatives 4a-d was successfully synthesized, displaying strong aggregationinduced enhanced emission (AIEE) photophysical properties. Computational optimization of 4a-d using B3LYP/6-31G* basis set disclose compound 4a exhibited the highest HOMO-LUMO energy gap (2.23 eV), suggesting enhanced electronic stability and superior emission properties upon aggregation. In contrast, the smallest gap in 4d (2.01 eV) conferred higher sensitivity to environmental changes, making it a standout candidate for advanced sensing applications. Compounds 4ad demonstrated significant sensitivity to low pH environments, promoting them as promising candidates for pH-responsive applications. Additionally, specific acid sensing studies revealed that 4d selectively interacts with sulfuric acid (H_2SO_4) with detection limit of 16.48 µM (1.62 ppm). Furthermore, stability and reusability tests confirmed the sturdiness of 4d as a sensor even after several successive utilization cycles. These findings underscore the versatility of the reported imidazole derivatives as promising versatile materials for applications in environmental monitoring, chemical sensing, and optoelectronic technologies. The robust AIEE properties, coupled with exceptional sensitivity and selectivity, position these compounds as valuable contributors to the development of next-generation functional materials.

Data availability

The raw data required to reproduce these findings are available upon request.

Conflicts of interest

There are no conflicts to declare.

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