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Frank R. Fronczek
Louisiana State Univ, Dept Chem

Jian Xu
Louisiana State Univ, Div Elect & Comp Engn

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Simultaneous Enhancement of Near-Infrared Emission and Dye Photodegradation in a Racemic Aspartic Acid Compound via Metal-Ion Modification

Ye Wu,† Pengfei Ou,‡ Frank R. Fronczek,§ Jun Song, Yingcheng Lin,*⊥ Hui-Min Wen,*# and Jian Xu*∥

†School of Electrical and Automation Engineering, Jiangsu Key Laboratory of 3D Printing Equipment and Manufacturing, Nanjing Normal University, Nanjing 210046, P. R. China
‡Department of Mining and Materials Engineering, McGill University, Montreal, QC H3A 0C5, Canada
§Department of Chemistry and Division of Electrical and Computer Engineering, Louisiana State University, Baton Rouge, Louisiana 70803, United States
∥School of Microelectronics and Communication Engineering, Key Laboratory of Dependable Service Computing in Cyber Physical Society (Chongqing University) of Ministry of Education, Chongqing University, Chongqing 400044, China
#College of Chemical Engineering, Zhejiang University of Technology, Zhejiang 310014, P. R. China

Supporting Information

ABSTRACT: Changing functionalities of materials using simple methods is an active area of research, as it is “green” and lowers the developing cost of new products for the enterprises. A new small molecule racemic N,N-dimethyl aspartic acid has been prepared. Its structure is determined by single-crystal X-ray diffraction. It is characterized by FTIR, XPS, 1H NMR, and mass spectroscopy. Its near-infrared luminescence can be enhanced by the combination of metal ions, including Dy3+, Gd3+, Nd3+, Er3+, Sr3+, Y3+, Zn2+, Zr4+, Ho3+, Yb3+, La3+, Pr6+/Pr3+, and Sm3+ ions. An optical chemistry mechanism upon interaction between the sensitizer and activator is proposed. Furthermore, the association of Ca2+, Sr2+, or Zr4+ ions to the molecule enhanced its photodegradation for dyes under white-light irradiation. Specifically, rhodamine 6G can be degraded by the Ca2+-modified molecule; rhodamine B, rhodamine 6G, and fluorescein sodium salt can be degraded by the Sr2+- or Zr4+-modified molecule. This surprising development opens a way in simultaneously increasing NIR luminescence and the ability of dye photodegradation for the investigated molecule.

1. INTRODUCTION

The development of materials with multifunctionalities is important, which can be used to fulfill different tasks and therefore lower the cost for the enterprises.

One type of functionality we are interested in for the materials is the near-infrared (NIR) emission. Indeed, synthesis of materials with photoluminescence in the NIR window (750–1100 nm) has been an active area of research given that their application in biomedical imaging within this region is superior to the ultraviolet–visible range.¹–⁵ Much improvement in imaging quality can be achieved due to low levels of tissue autofluorescence, diminished photon scattering, and reduced photon absorption. Several materials systems have been reported to show NIR fluorescence, including semiconductor quantum dots,⁶,⁷ metal organic complexes,⁸ small molecule dyes,⁹ polymers,¹⁰ carbon nanotubes,¹¹ metal clusters,¹² and inorganic nanoparticles.¹³–¹⁹

One challenge that is associated with the application of NIR fluorophores would be the optimization of the luminescence. Suppose that we have synthesized one NIR fluorophore, can we further increase the NIR fluorescence intensity or shift the center wavelength using some facile methods? This is extremely useful in practical clinic or industry applications where the access to the chemical hood and chemical reactors is limited. Simple and cost-effective ways for modification of the optical properties of the fluorophores are in great demand. One possible way is
using the metal ions to process the synthesized NIR fluorophore. Given that the metal ions have rich energy levels, the optical properties of the materials may be particularly tuned by incorporation of NIR-emissive metal sites such as Er\(^{3+}\), Nd\(^{3+}\), Y\(^{3+}\), La\(^{3+}\), Pr\(^{3+}\), Gd\(^{3+}\), Dy\(^{3+}\), and Yb\(^{3+}\).\(^{20-27}\)

Another type of functionality in the materials we are looking for is dye degradation. Dyes are extensively used in textile paints for coloring clothes. The dyes that are discharged into the rivers can seriously harm a lot of birds and animals. The dyes create an unsafe environment for them due to their toxicity. Dye molecules generally contain complex aromatic molecular structures that make them more stable and more difficult to biodegrade. It is critical to perform the dye pretreatment for the wastewater as these dye materials would end up in our drinking water if the wastewater is not treated properly. The dyes can be decomposed via photodegradation. Especially, white-light- or solar-light-assisted photodegradation for organic pollutants is considered to be a green technology.

Aspartic acid is an \(\alpha\)-amino acid that is used in the biosynthesis of proteins. It is industrially made by fermentative or enzymatic batch process from fumaric acid and ammonia, using aspartase as the enzyme.\(^{28}\) L-/d-Aspartate (the ionic form of aspartic acid) is reported to be present in the human brain.\(^{29}\)

It is rarely reported that aspartic acid and its derivatives are chemically synthesized without using the aspartase enzyme. No reports about them have shown NIR fluorescence as well as dye-photodegradation effects.

Herein, we first report chemical synthesis of a new type of aspartic acid single crystal: racemic \(\text{N}_2\text{N}_2\text{-dimethyl aspartic acid (compound 1)}\). The scheme is shown in Scheme 1. Here, fumaric acid is utilized as one of the raw materials for making compound 1.

\[\text{HO-CH(C\text{H}_3)\text{CH}_2-COOH}\]

**Scheme 1. Synthetic Route of Racemic \(\text{N}_2\text{N}_2\text{-Dimethyl Aspartic Acid}\)**

Compound 1 was characterized by single-crystal X-ray diffraction, X-ray photoelectron spectroscopy (XPS), \(^1\)H nuclear magnetic resonance (NMR) spectroscopy, Fourier transform infrared (FTIR) spectroscopy, and mass spectroscopy.

Compound 1 shows NIR fluorescence in the range of 800–1100 nm when it is excited by 785 nm light. It can be modified by the metal ions to increase the fluorescence intensity as well as to shift the peak wavelength. Its NIR photoluminescence corresponding to the 785 nm light excitation can be tuned by combination of different metal ions, including Dy\(^{3+}\), Gd\(^{3+}\), Nd\(^{3+}\), Er\(^{3+}\), \(\text{Sr}^{2+}\), \(\text{Ti}^{3+}\), \(\text{Y}^{3+}\), \(\text{Zn}^{2+}\), \(\text{Zr}^{4+}\), \(\text{Ho}^{3+}\), \(\text{Yb}^{3+}\), La\(^{3+}\), Pr\(^{3+}/\text{Pr}^{4+}\), Sm\(^{3+}\), Ag\(^+\), Ce\(^{3+}\), Cs\(^{+}\), In\(^{3+}\), Li\(^{+}\), Mo\(^{6+}\), Na\(^{+}\), Ni\(^{2+}\), and Rb\(^{+}\) ions. The introduction of Dy\(^{3+}\), Gd\(^{3+}\), Nd\(^{3+}\), Er\(^{3+}\), \(\text{Sr}^{2+}\), \(\text{Ti}^{3+}\), \(\text{Y}^{3+}\), \(\text{Zn}^{2+}\), \(\text{Zr}^{4+}\), \(\text{Ho}^{3+}\), \(\text{Yb}^{3+}\), La\(^{3+}\), Pr\(^{3+}/\text{Pr}^{4+}\), and Sm\(^{3+}\) to the molecule leads to an enhanced NIR photoluminescence band centered around 813 nm. Furthermore, the introduction of Nd\(^{3+}\) to this molecule leads to intensified NIR photoluminescence bands at 812.36, 868.86, 888.31, 1014.14, and 1055.68 nm.

Additionally, metal-ion modification of the studied molecule leads to the enhanced dye photodegradation. Here, Ca\(^{2+}\)-modified \(\text{N}_2\text{N}_2\)-dimethyl aspartic acid (compound 2) can be used for photodegradation of rhodamine 6G under white-light irradiation. Both Zr\(^{4+}\)-modified \(\text{N}_2\text{N}_2\)-dimethyl aspartic acid (compound 3) and Sr\(^{2+}\)-modified \(\text{N}_2\text{N}_2\)-dimethyl aspartic acid (compound 4) can be applied for photodegradation of rhodamine 6G, rhodamine B, and fluorescein sodium salt under white-light irradiation.

The significance of the metal-ion modification for compound 1 is supported by the enhancement of the NIR fluorescence and dye photodegradation.

### 2. RESULTS AND DISCUSSION

#### 2.1. Single-Crystal X-ray Diffraction

Single-crystal X-ray diffraction indicated that compound 1 crystallizes in the space group P21/c. As depicted in Figure 1, its structure is similar to L-

![Figure 1. Single-crystal structures of racemic \(\text{N}_2\text{N}_2\)-dimethyl aspartic acid (black sphere: carbon atom; pink sphere: hydrogen atom; red sphere: oxygen atom; blue sphere: nitrogen atom) are viewed from different axes: (a) a axis; (b) b axis; (c) c axis.)](image)

#### 2.2. XPS, FTIR, NMR, and Mass Spectroscopy Characterization

**Figure 2a** shows the high-resolution XPS spectra of O 1s, which reveal two peaks at 531.21 and 532.57 eV. They correspond to the –C–O– and –C–O–H– bonding, respectively. For C 1s XPS spectra (Figure 2b), three peaks at 284.04, 285.19, and 288.05 eV are observed, which are attributed to –C–C–, –O–C–O–, and –C–H–N–. N 1s XPS spectra present two peaks at 399.43 and 401.19 eV (Figure 2c). The 399.43 eV peak is considered to be due to \(\text{NH}^+\)
The assigned to C−O−H stretching. The 3001 cm$^{-1}$ peak is assigned to C−N−H in-plane bending. The 1456.40, 1606.48, 1708.87, 1888.40, 2362.47, 2746.78, 2848.47, 2926.31, 2998.55, and 3118.47 cm$^{-1}$. The 676.56 cm$^{-1}$ peak is assigned to C−O−O−H in-plane bending. The 904.48 cm$^{-1}$ peak is considered as −C=−N=− stretching. The 958.48 cm$^{-1}$ peak is due to C−N=− stretching. The 1180.79 cm$^{-1}$ peak is considered to be due to N−H rocking. The 1372.95 cm$^{-1}$ peak is assigned to CH$_2$ wagging. The 1372.95 cm$^{-1}$ peak is related to CH$_2$ in-plane bending. The 1456.40 and 1606.48 cm$^{-1}$ peaks are attributed to −COO− stretching. The 1708.87 cm$^{-1}$ peak corresponds to C−O− stretching. The 1888.40, 2362.47, 2746.78, and 2848.47 cm$^{-1}$ peaks are attributed to the combinational effect of C−O stretching, C−H stretching, C−C stretching, NH in-plane bending, and OH in-plane bending. The 2926.31 cm$^{-1}$ peak is assigned to −CH$_3$ wagging. The 2998.55 cm$^{-1}$ peak is related to −OH− stretching. The 3001 cm$^{-1}$ peak is attributed to C−H stretching. The 3118.47 cm$^{-1}$ peak is due to N−H stretching.$^{13}$

$^1$H NMR spectra contain six proton lines at 3.76, 3.49, 2.88, 2.72, 2.55, and 1.22 ppm, which are assigned to the groups as labeled in Figure 3b. Mass spectra (see Figure 3c) show the [(C$_6$H$_{11}$NO$_4$-C$_2$H$_5$OH)]$^{+}$ ion (160 m/z) as the major product ion. Other product ions were due to the further loss of NH, CH$_2$, NH-CH$_3$, COOH, COOCH, CH$_2$CO, CO, and CO$_2$. The peaks and their corresponding product ions are listed in Table 2.

### 2.3. Optical Properties

The optical properties of compound 1 are studied. Its absorption presents peaks at 746.4, 877.64, and 980.73 nm (Figure 4a). The peaks at 746.4 and 877.64 nm are weak, while the 980.73 nm peak is strong.

Figure 4b shows the fluorescence spectra excited by 280, 310, 320, 340, 350, and 360 nm. When the sample is excited by 280, 310, and 320 nm light, the fluorescence shows two peaks at 383 and 523 nm, separately. When it is excited by 340 nm light, only the 523 nm peak survives. Green-light emission is observed. When it is excited by 350 and 360 nm, the green-light emission is enhanced. Figure 4c presents a fluorescence peak at 531.77 nm when it is excited by 440 nm light. When it is excited by 540 nm light, the fluorescence spectra present a peak at 617.73 nm and a shoulder around 708 nm (Figure 4d). When it is excited by 610 nm light, the fluorescence spectra reveal two peaks at 670 and 720 nm (Figure 4e). When it is excited by 785 nm light, the emission at 813.18 nm is observed.

### 2.4. Enhancement of NIR Fluorescence

NIR fluorescence spectra of compound 1 modified with Dy$^{3+}$, Gd$^{3+}$, Nd$^{3+}$, Er$^{3+}$, Sr$^{2+}$, Ti$^{3+}$, Y$^{3+}$, Zn$^{2+}$, Zr$^{4+}$, Ho$^{3+}$, Yb$^{3+}$, La$^{3+}$, Pr$^{3+}$/Pr$^{4+}$, Sm$^{3+}$, Ag$^+$, Ce$^{4+}$, Cr$^{3+}$, Cs$^+$, In$^+$, Li$^+$, Mo$^{6+}$, Na$^+$, Ni$^{3+}$, and Rb$^+$ ions have been obtained using a 785 nm diode, as shown in Figures 5 and 6. The excitation light was not intensively focused on the samples since the samples may decompose upon large excitation density. Figure 5a depicts the NIR fluorescence spectra excited by 280, 310, 320, 340, 350, and 360 nm light. The peak at 813.18 nm whose intensity is considered as $I_0$. Surprisingly, the sample that was modified by Nd$^{3+}$ ions shows several bands at 812.36, 868.86, 888.31, 1014.14, and 1055.68 nm. The 868.86 nm peak shows the highest intensity whose intensity is 2 times greater than $I_0$. The Gd$^{3+}$-ion-modified sample shows a peak at 811.25 nm whose intensity is 4.67 times greater than $I_0$. It also shows a weak peak at 1013.55 nm. The Dy$^{3+}$-ion-modified sample shows a peak at 813.18 nm whose intensity is 0.8 times higher than $I_0$. It also shows a weak peak at 813.18 nm. The Dy$^{3+}$-ion-modified sample depicts two strong peaks at 800.69 and 816.76 nm and a weak peak at 1014.14 nm. The intensity of the 816.76 nm peak is 1.38 times greater than $I_0$.

Figure 5b–e plots the fluorescence spectra of the samples modified by Er$^{3+}$, Sr$^{2+}$, Ti$^{3+}$, Y$^{3+}$, Zn$^{2+}$, Zr$^{4+}$, Ho$^{3+}$, Yb$^{3+}$, La$^{3+}$, Pr$^{3+}$/Pr$^{4+}$, and Sm$^{3+}$ ions. They show a profile similar to that of the Gd$^{3+}$-ion-modified sample. They all present a peak around 810−813 nm whose intensity is 0.8−2.8 times higher than $I_0$. The samples modified by Ce$^{4+}$, Pr$^{3+}$, Cs$^+$, In$^+$, Li$^+$, Mo$^{6+}$, Na$^+$, Ni$^{3+}$, and Rb$^+$ ions all show a suppressed fluorescence peak whose intensity is comparable to $I_0$ (Figure 6a–c). The sample modified by Ag$^+$ ions shows a suppressed fluorescence profile (Figure 6a). The

### Table 2. Mass Spectra of Product Ions for Compound 1

| peaks (m/z) | product ions |
|------------|--------------|
| 147        | [(C$_6$H$_{11}$NO$_4$-CH$_3$)$_3$]$^+$ |
| 146        | [(C$_6$H$_{11}$NO$_4$-NH)$_3$]$^+$ |
| 145        | [(C$_6$H$_{11}$NO$_4$-CH$_3$)$_2$]$^+$ |
| 143        | [(C$_6$H$_{11}$NO$_4$-H$_2$O)$_3$]$^+$ |
| 133        | [(C$_6$H$_{11}$NO$_4$-CO)$_3$]$^+$ |
| 131        | [(C$_6$H$_{11}$NO$_4$-(NH + CH$_3$)]$^+$ |
| 119        | [(C$_6$H$_{11}$NO$_4$-CH$_2$CO)$_3$]$^+$ |
| 117        | [(C$_6$H$_{11}$NO$_4$-CO)$_2$]$^+$ |
| 116        | [(C$_6$H$_{11}$NO$_4$-HN(CH$_3$)]$^+$ |
| 115        | [(C$_6$H$_{11}$NO$_4$-(H$_2$O + CO)]$^+$ |
| 104        | [(C$_6$H$_{11}$NO$_4$-COOCH)]$^+$ |
| 101        | [(C$_6$H$_{11}$NO$_4$-CH$_2$COOCH)]$^+$ |
| 87         | [(C$_6$H$_{11}$NO$_4$-(COOH + CH$_3$)]$^+$ |
| 83         | [(C$_6$H$_{11}$NO$_4$-(CH$_2$COOH + H$_2$O)]$^+$ |
| 81         | [(C$_6$H$_{11}$NO$_4$-(CH$_2$COOH + H$_2$O + H$_2$)]$^+$ |

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results prove that Ag\textsuperscript{+}, Ce\textsuperscript{4+}, Cr\textsuperscript{3+}, Cs\textsuperscript{+}, In\textsuperscript{+}, Li\textsuperscript{+}, Mo\textsuperscript{6+}, Na\textsuperscript{+}, Ni\textsuperscript{3+}, and Rb\textsuperscript{+} ions are not good for the modification of compound 1 in order to enhance the NIR fluorescence.

An optical chemistry mechanism is proposed to explain the NIR fluorescence of Nd\textsuperscript{3+}-ion-modified compound 1 (see Figure 7). It depicts the energy level transitions of Nd\textsuperscript{3+}-ion-modified compound 1 under 785 nm light excitation. In this diagram, it is supposed that the \(-\text{O}--\text{C}--\text{N}\) group in compound 1 is excited via absorbing excitation photons. The excited \(-\text{O}--\text{C}--\text{N}\) group is utilized as a sensitizer for the Nd\textsuperscript{3+} activators. This results in energy transfer from the \(-\text{O}--\text{C}--\text{N}\) group to the Nd\textsuperscript{3+} ion. As a result, the emission is forced to be

Figure 4. (a) Optical absorption of compound 1. (b) Fluorescence spectra show white-to-green light transition when the excitation light varies between 280 and 360 nm. Polychromatic fluorescence spectra are shown when excited by light sources with various wavelengths: (c) 440 nm; (d) 540 nm; (e) 610 nm; (f) 785 nm.

Figure 5. Near-infrared fluorescence of compound 1 could be enhanced by modification with various metal ions: (a) Dy\textsuperscript{3+}, Gd\textsuperscript{3+}, and Nd\textsuperscript{3+}; (b) Er\textsuperscript{3+}, Sr\textsuperscript{2+}, and Tl\textsuperscript{3+}; (c) Y\textsuperscript{3+}, Zn\textsuperscript{2+}, and Zr\textsuperscript{4+}; (d) Ho\textsuperscript{3+} and Yb\textsuperscript{3+}; (e) La\textsuperscript{3+}, Pr\textsuperscript{3+}/Pr\textsuperscript{4+}, and Sm\textsuperscript{3+}. Here, “Control” is the fluorescence spectra of compound 1 without using any metal-ion modification.

Figure 6. Impact of modification by various metal ions for near-infrared fluorescence of compound 1 is examined: (a) Ag\textsuperscript{+}, Ce\textsuperscript{4+}, Cr\textsuperscript{3+}, and Cs\textsuperscript{+}; (b) In\textsuperscript{+}, Li\textsuperscript{+}, and Mo\textsuperscript{6+}; (c) Na\textsuperscript{+}, Ni\textsuperscript{3+}, and Rb\textsuperscript{+}. Here, “Control” is its fluorescence spectra without using any metal-ion modification.
Figure 7. Optical chemistry mechanism of the Nd\(^{3+}\)-ion-modified N,N-dimethyl aspartic acid.

The energy transfer process is described in Figure 7, which results in the emission light of 812.36 (energy level transition happens from \(^2\)H\(_{11/2}\) to \(^4\)I\(_{13/2}\)), 868.86 (\(^2\)P\(_{1/2}\) to \(^4\)F\(_{3/2}\)), 1014.14 (\(^2\)F\(_{5/2}\) to \(^4\)I\(_{11/2}\)), 1055.68 nm (\(^2\)F\(_{3/2}\) to \(^4\)I\(_{13/2}\)).\(^{36}\) The optical chemistry of compound 1 modified by other ions under 785 nm light excitation is similar. The –O–C–N– group works as a sensitizer for the specific metal ion (i.e., Dy\(^{3+}\), Gd\(^{3+}\), Er\(^{3+}\), Sr\(^{2+}\), Tl\(^{3+}\), Y\(^{3+}\), Zn\(^{2+}\), Zr\(^{4+}\), Ho\(^{3+}\), Yb\(^{3+}\), La\(^{3+}\), Pr\(^{6+}/\)Pr\(^{3+}\), Sm\(^{3+}\), Ag\(^{+}\), Ce\(^{4+}\), Ce\(^{3+}\), Cs\(^{+}\), In\(^{3+}\), Li\(^{+}\), Mo\(^{6+}\), Na\(^{+}\), Ni\(^{3+}\), and Rb\(^{+}\) ions) activators. The only difference is the emission light and the associated energy level transitions corresponding to the metal ions. The general structure of a laser contains a pumping source, gain medium, and mirrors. While the pumping source and mirrors are commercially available, the gain medium is not always conveniently accessible. The gain medium is the most important factor of the laser as it determines the operation wavelength and optical properties of the laser. Here, single crystals of compound 1 were successfully made. It shows interestingly polychromatic photoluminescence covering an ultrawide range of UV–vis–NIR optical windows. It can be selected as the gain medium to make the laser in specific optical windows. More importantly, it shows NIR fluorescence around 800–1100 nm. The NIR fluorescence intensity can be enhanced, and the emitting wavelength can be shifted when the modification using various metal ions is applied. It indicates that this small molecule can be flexibly utilized as the gain medium for making NIR luminescence lasers covering from 800 to 1100 nm range. It can be conveniently done via commercial pumping sources such as 785 nm LDs.

2.5. Enhancement of Dye Photodegradation via Metal-Ion Modification. Figure 8 compares discoloration of the dyes by using compounds 1, 2, 3, and 4. a1, a2, and a3 are the vials filled with rhodamine B (RB), rhodamine 6G (R6G), and fluorescein sodium salt (FSS) before the discoloration test. Compound 1 was then utilized for running the dye-photodegradation test. b1, b2, and b3 are the vials filled with RB, R6G, and FSS after the discoloration test. It should be noted that the colors of b1, b2, and b3 are almost the same as those of a1, a2, and a3, which indicates that the use of compound 1 is not effective for the photodegradation of RB, R6G, and FSS.

Compound 2 was used for the photodegradation test of R6G. c1, c2, and c3 are the vials filled with R6G after it was carried out with irradiation times of 30, 60, and 80 min. Its color was changed from dark blue to dark yellow after the photodegradation test was completed. When compound 1 was used, the color of R6G was not removed. However, R6G was found to be decolorized by compound 2. Therefore, it suggests that the introduction of Ca\(^{2+}\) ions enhances the ability of R6G photodegradation for the investigated molecule, N,N-dimethyl aspartic acid. Compounds 3 and 4 were tested for the photodegradation of RB, R6G, and FSS. d1, d2, and d3 are the vials filled with RB, R6G, and FSS after the photodegradation test via using compound 1. e1, e2, and e3 are the vials filled with R6G after photodegradation tests 30, 60, and 80 min via using compound 2. d1, d2, and d3 are the vials filled with RB after photodegradation tests for 20, 40, and 60 min via using compound 3. d4, d5, and d6 are the vials filled with R6G after photodegradation tests for 20, 40, and 60 min via compound 3. e1, e2, and e3 are the vials filled with RB after photodegradation tests 20, 40, and 80 min via compound 4. e4, e5, and e6 are the vials filled with R6G after photodegradation tests 20, 40, and 80 min via using compound 4.

Figure 8. Discoloration of the dyes using photodegradation under white-light irradiation. a1, a2, and a3 are the vials filled with RB, R6G, and FSS after the photodegradation test. b1, b2, and b3 are the vials filled with RB, R6G, and FSS after the photodegradation test via using compound 1. c1, c2, and c3 are the vials filled with R6G after photodegradation tests for 30, 60, and 80 min via using compound 2. d1, d2, and d3 are the vials filled with RB after photodegradation tests for 20, 40, and 60 min via using compound 3. d4, d5, and d6 are the vials filled with R6G after photodegradation tests for 20, 40, and 60 min via compound 3. e1, e2, and e3 are the vials filled with RB after photodegradation tests for 20, 40, and 80 min via using compound 4. e4, e5, and e6 are the vials filled with R6G after photodegradation tests for 20, 40, and 80 min via using compound 4.

The absorption and fluorescence spectra were studied for the impact of compounds 2, 3, and 4 on dye photodegradation. The absorption spectral results are shown in Figures S1–S3. The fluorescence spectral analysis is shown as follows. Figure 9a,b shows the fluorescence spectra corresponding to the photodegradation via compound 2. Figure 9a shows the fluorescence spectra of R6G after photodegradation test via compound 2. (a) Fluorescence spectra of R6G before photodegradation test. (b) Fluorescence spectral study of the influence of irradiation time. Here, the excitation wavelength is 523 nm.
fluorescence of pristine R6G, which presents a 573 nm peak upon 523 nm light excitation. Figure 9b reveals the fluorescence spectra of R6G after photodegradation processing of using compound 2 conducted in 30, 60, and 80 min. They all present a shoulder around 570–750 nm. The 573 nm peak disappears, which suggests the degradation of R6G.

Figure 10a presents the fluorescence spectra of RB before and after the photodegradation experiment of using compound 3 (the excitation wavelength is 480 nm). For pristine RB, it shows a 600 nm peak (the pink curve in Figure 10a). After the test is run for 20 min, the fluorescence spectra show a peak at 557 nm. After the photodegradation test is conducted for 40 min, a sharp peak centered at 560 nm appears. After the photodegradation test is performed for 60 min, it shows a small peak around 542 nm.

Figure 10b presents the fluorescence spectra of R6G after the photodegradation test through compound 3. The characteristic 573 nm peak of R6G (as indicated in Figure 9a) is absent in the fluorescence spectra, which means that R6G has been degraded after photodegradation processing is done after 20, 40, and 80 min. The fluorescence spectra for FSS photodegradation with respect to different times are shown in Figure 10c. The fluorescence of pristine FSS shows a characteristic peak at 524 nm when excited by 440 nm light (pink curve in Figure 10c). This peak is absent in the fluorescence spectra after the photodegradation test is run for 20, 40, and 80 min, which suggests the degradation of FSS.

The fluorescence spectra corresponding to the photodegradation test via compound 4 are revealed in Figure 11a–c. The characteristic 573 nm peak of R6G is absent in the fluorescence spectra when the sample is excited by 523 nm light (see Figure 11b). It indicates the degradation of R6G.

The fluorescence spectra for FSS photodegradation are shown in Figure 11c. The fluorescence of pristine FSS shows a characteristic peak at 524 nm when excited by 440 nm light (see the cyan curve in Figure 11c). This peak is found to be absent in the fluorescence spectra after the photodegradation test is run for 20, 40, and 60 min. This suggests the degradation of FSS.

2.6. Mechanism of the Photodegradation. Ca$^{2+}$-, Zr$^{4+}$-, or Sr$^{2+}$-ion-modified aspartic acid showed better photodegradation activities, likely due to the effects of increased electron–hole recombination and reduced material defects. The proposed mechanism for the photodegradation of the dyes can be assumed as (reactions 1–6): under white-light irradiation, the excited electrons and holes diffuse to the sample surface (reaction 1) and react with surface species. With metal-ion-modified aspartic acid (MIMAA), the surface charges become more positive, leading to more OH$^-$ ions adsorbed on the material surface for charge balance. The adsorbed OH$^-$ ions can accept holes to form hydroxyl radicals (OH) (reaction 3). It should be noted that the hole sites attack the adsorbed dye molecules directly (reaction 2), while OH radicals attack the oxygenated structure (reaction 4) to form dye fragments. Subsequently, the dye fragments react with the active sites, including holes and OH radicals (reactions 5 and 6). For this reason, more absorption sites of the n–π* transition are formed at the initial stage due to the OH attack.

\[
\text{MIMAA} + \text{photon} \rightarrow \text{h}^+ + \text{e}^- \quad (1)
\]
\[
\text{dye}_{\text{adsorbed}}^- + \text{h}^+ \rightarrow \text{CO}_2 + \text{H}_2\text{O} \quad (2)
\]
\[
\text{OH}_{\text{adsorbed}}^- + \text{h}^+ \rightarrow \text{OH} \quad (3)
\]
\[
(\text{phenyl} - \text{O} - \text{R}) + \text{OH} + \text{H}^+ \rightarrow (\text{phenyl} - \text{OH}) + (\text{HO} - \text{R}) \quad (4)
\]
\[
(\text{phenyl} - \text{OH}) + \text{h}^0\text{or OH} \rightarrow \text{CO}_2 + \text{H}_2\text{O} \quad (5)
\]
\[
(\text{HO} - \text{R}) + \text{h}^0\text{or OH} \rightarrow \text{CO}_2 + \text{H}_2\text{O} \quad (6)
\]
3. CONCLUSIONS

A new type of aspartic acid single crystal, compound 1, has been synthesized.

The enhanced NIR photoluminescence of the neodymium ion has been achieved from the neodymium-ion-associated compound 1 sensitized by the =O−C=N− group upon 785 nm excitation. The results suggest that it has promising applications in neodymium-based optical materials and devices. Indirect excitation mechanisms for the investigated Nd3+-modified compound 1 are deduced mainly to be =O−C=N− group emission introduced energy transition from 2H11/2 to 4I13/2 (for 1055.68 nm light emission). Further work is in progress concerning much evidence for the optical chemistry mechanisms. Also, the intensified NIR luminescence of the Dy3+, Gd3+, Er3+, Sr2+, Tl3+, Y3+, Zr4+, Zn2+, Zr4+, La3+, Pr6+/Pr3+, Sm3+, Ag+, Ce4+, Ce3+, Cs+, In3−, Li+, Mo6+, Na+, and Ni3+-ion-modified N,N-dimethyl aspartic acid is similar to that of Zr4+-ion-modified N,N-dimethyl aspartic acid.

4. EXPERIMENTAL SECTION

4.1. Materials. All commercially available reagents and solvents were bought from Alfa Aesar and used without further purification.

4.2. Preparation of Organic Complex Solution. Solution a was made by dissolving 2-methylimidazole (2 g), terephthalic acid (5 g), fumaric acid (5 g), d-(+)-glucose (5 g), oxalic acid dihydrate (5 g), and dimethyl sulfoxide (50 mL) in deionized water (200 mL). Solution b was prepared by mixing dimethylformamide (300 mL), oleic acid (40 mL), triethylamine (40 mL), diethylene glycol (40 mL), dimethyl sulfoxide (20 mL), and methacrylic anhydride (10 mL). Solution c was made by dissolving L-cysteine (2 g) in deionized water (50 mL). Solutions a, b, and c were mixed and stirred to get the organic complex solution.

4.3. Preparation of Racemic N,N-Dimethyl Aspartic Acid. Niobium(V) chloride (0.1 g), samarium chloride hydrate (0.4 g), and sodium hydroxide (0.016 g) were dissolved in deionized water (10 mL) to get solution d. Solution d was mixed with 40 mL of organic complex solution prepared in the previous step. The acquired solution was heated with 80 °C for 2 h. The crystals were recrystallized in ethanol for single-crystal X-ray diffraction measurement.

4.4. Preparation of Metal-Ion-Modified N,N-Dimethyl Aspartic Acid. The synthesized sample of racemic N,N-dimethyl aspartic acid (0.1 g) was dissolved in methanol (50 mL) to get solution d.

Zirconium(IV) chloride (0.12 g) was first dissolved in water (1 mL). Then the acquired Zr4+-ion solution was added to solution d (1 mL) in a centrifuge tube, which was shaken for 5 min to get Zr4+-ion-modified N,N-dimethyl aspartic acid.

The preparation of Ca2+, Dy3+, Gd3+, Er3+, Sr2+, Tl3+, Y3+, Zr4+, Zn2+, Zr4+, La3+, Pr6+/Pr3+, Sm3+, Ag+, Ce4+, Ce3+, Cs+, In3−, Li+, Mo6+, Na+, and Ni3+-ion-modified N,N-dimethyl aspartic acid is similar to that of Zr4+-ion-modified N,N-dimethyl aspartic acid.

4.5. Characterization and Optical Study. The XPS study was performed by using a K-Alpha XPS instrument (Thermo Scientific). Single-crystal X-ray diffraction measurement was done through a Bruker APEX diffractometer. UV−vis−NIR absorption and UV−vis−fluorescence spectroscopy were collected through a SPARK spectrometer. The FTIR spectra were collected via a Thermo Fisher Nicolet IS 10 Fourier transform infrared (FTIR) spectrometer equipped with a Smart iTR diamond ATR accessory and with a deuterated triglycine sulfate detector. The mass spectra were collected through an accurate mass electrospray ionization (ESI) mass spectrometer coupled with a high-performance liquid chromatograph (HPLC). Specifically, an Agilent 1260 Infinity II quaternary liquid chromatograph coupled to an Agilent 6230 electrospray time-of-flight mass spectrometer was used for detection of analytes. NMR spectra were collected via a Bruker AV-400 MHz spectrometer. Deuterated dimethyl sulfoxide (DMSO-d6) was used as a solvent for NMR.

A commercial SPARK spectrometer was used to collect the absorption spectra. The fluorescence spectra corresponding to excitation light wavelengths of 280, 310, 320, 340, 350, 360, 440, 540, and 610 nm were also collected through the SPARK spectrometer.

For collecting NIR fluorescence, a laser diode (center wavelength = 785 nm, Thermal Scientific, Inc.) was used as the excitation light source. An optical fiber bundle was used to deliver a laser power of 20 mW. Another optical fiber bundle was moving around the sample surface to collect the NIR fluorescence. An NIR long-pass filter was set in front of an NIR spectrometer (Ocean Optics, Inc.) to get rid of the excitation light. A lens was used to focus the collected fluorescence into the inlet of the NIR spectrometer.

4.6. Photodegradation Experiment. Fluorescein sodium salt (0.1 g) was dissolved in deionized water (50 mL) to obtain the solution FSS. Rhodamine 6G (0.1 g) was dissolved in ethanol (50 mL) to acquire the R6G solution. Rhodamine B (0.53 g) was dissolved in deionized water (100 mL) to get the RB solution. Calcium chloride (1.3 g) was mixed with water (10 mL) to get the Ca2+ solution (solution 1). Compound 1 (0.2 g) was added to solution 1 and stirred to get solution 2. Solution 2 (1.5 mL) was mixed with R6G (500 μL) and stirred to get solution 3. A halogen light bulb (Model: Double Ended Quartz FCL, OSRAM) with a maximum output optical power of 500 W was utilized for white-light irradiation. The output optical power of the light bulb is controlled by a self-built controller. Solution 3 was irradiated by white light for 30, 60, and 80 min. The irradiated solution 3 was taken to an optical spectrometer for optical absorption and fluorescence measurement. The procedures of the photodegradation of RB, R6G, and FSS through compounds 3 and 4 are similar.
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■ ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsomega.9b02434.

Crystallographic data for racemic \( \text{N,N-dimethyl aspartic acid (CIF)} \)

Absorption spectra of dye solutions after the photo-degradation test (Figures S1–S3) (PDF)

Accession Codes

The X-ray crystallographic coordinates for structures reported in this article have been deposited at the Cambridge Crystallographic Data Centre (CCDC), under deposition numbers CCDC 1904827. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre via www.ccdc.cam.ac.uk/data_request/cif.

■ AUTHOR INFORMATION

Corresponding Authors

*E-mail: liny@cqu.edu.cn (Y.L.).
*E-mail: huiminwen@rjut.edu.cn (H.-M.W.).
*E-mail: jianxu1@lsu.edu (J.X.).

ORCID

Ye Wu: 0000-0002-4831-9729
Pengfei Ou: 0000-0002-3630-0385
Frank R. Fronczek: 0000-0001-5544-2779
Jun Song: 0000-0003-3675-574X

Author Contributions

Y.W., P.O., and F.R.F. contributed equally.

Notes

The authors declare no competing financial interest.

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