



Article

# Understanding of Förster Resonance Energy Transfer (FRET) in Ionic Materials

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Abstract: Herein, an ionic material (IM) with Förster Resonance Energy Transfer (FRET) characteristics is reported for the first time. The IM is designed by pairing a Nile Blue A cation (NBA<sup>+</sup>) with an anionic near-infrared (NIR) dye, IR820<sup>-</sup>, using a facile ion exchange reaction. These two dyes absorb at different wavelength regions. In addition, NBA<sup>+</sup> fluorescence emission spectrum overlaps with IR820<sup>-</sup> absorption spectrum, which is one requirement for the occurrence of the FRET phenomenon. Therefore, the photophysical properties of the IM were studied in detail to investigate the FRET mechanism in IM for potential dye sensitized solar cell (DSSCs) application. Detailed examination of photophysical properties of parent compounds, a mixture of the parent compounds, and the IM revealed that the IM exhibits FRET characteristics, but not the mixture of two dyes. The presence of spectator counterion in the mixture hindered the FRET mechanism while in the IM, both dyes are in close proximity as an ion pair, thus exhibiting FRET. All FRET parameters such as spectral overlap integral, Förster distance, and FRET energy confirm the FRET characteristics of the IM. This article presents a simple synthesis of a compound with FRET properties which can be further used for a variety of applications.

Keywords: ionic material; FRET; spectator ions; dye sensitized solar cell; near-infrared dyes



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## 1. Introduction

The sun is an obvious source of renewable and clean energy that would provide approximately 300,000 times the required energy for the global population if utilized. Among solar cells, dye-sensitized solar cells (DSSCs) are a type of third-generation photovoltaic cells that utilize dyes to parallel photosynthesis [1–3]. Typically, the production of DSSCs is economical as compared to the first- and second-generation solar cells [4]. However, DSSCs have not achieved the same high efficiencies as other types of solar devices [5]. Therefore, researchers are seeking new photosensitizers [6], electrolytes [7], electrode materials [8], etc., in enhancing the efficiency of DSSCs. This manuscript presents a new class of material that can be utilized as a potential photosensitizer in DSSCs.

Solar radiation consists of approximately 43% visible light, 49% near-infrared (NIR) light, and 7% ultra-violet light [9]. Thus, a dye with a broad absorption spectrum is one of the primary requirements to absorb maximum sunlight, which consequently affects the efficiency of DSSCs [10]. Since a large portion of UV sunlight is absorbed by the ozone layer, the absorbance of a compound in this region is not of significant importance. However, the longer wavelength NIR radiation can be exploited by utilizing NIR dyes (dyes absorbing from 750–1000 nm) in DSSCs [11,12]. Ideally, the sensitizing dye in DSSCs should absorb 80% or more of the solar spectrum (350–900 nm) [13]. The task of finding or synthesizing a single dye with this broad absorbance is quite difficult. However, two or more dyes can be combined which absorb at different wavelength regions of the electromagnetic spectrum to effectively broaden the absorption spectra that subsequently affects DSSC efficiency [14].

or more dyes can be combined which absorb at different wavelength regions of the electromagnetic spectrum to effectively broaden the absorption spectra that subsequently af-<sup>565</sup> fects DSSC efficiency [14].

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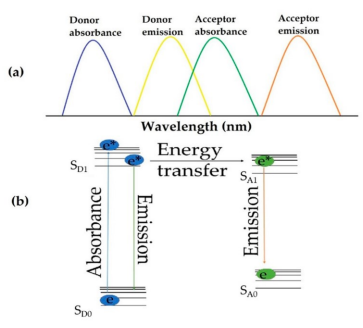


Figure 1. (a) Illustration of relationship between donor and acceptor spectra. (b) Energy diagram desdescribing epergy transfer and ecceptor spectra. (b) Energy diagram desdescribing epergy transfer and ecceptor spectra.

IMs are tunable ionic salts that share properties of ionic liquids but are solid at room IMs are tunable ionic salts that share properties of ionic liquids but are solid at room temperature [21–23]. IMs are ideal for several applications due to their enhanced photo and temperature [21–23]. IMs are ideal for several applications due to their enhanced photo and temperature [21–23]. IMs are ideal for several applications due to their enhanced photo and temperature [21–23]. IMs are ideal for several applications due to their enhanced photo and temperature [21–23]. In the several applications due to their enhanced photo and thermal stability, fluorescence quantum yields, and electrochemical characteristics [10,24,25]. These properties can be further tuned by tailoring counterions using a facile one-step ion exchange method without requiring complex organic synthesis [21]. IMs are advantageous one-step ion exchange method without requiring complex organic synthesis [21]. IMs are advantageous one-step ion exchange method without requiring complex organic synthesis [21]. IMs are advantageous one-step ion exchange method without requiring complex organic synthesis [21]. IMs are advantageous one-step ion exchange method without requiring complex organic synthesis [21]. IMs are advantageous one-step ion exchange method without requiring complex organic synthesis [21]. IMs are advantageous one-step ion exchange method without requiring complex organic synthesis [21]. IMs are advantageous one-step ion exchange method without requiring complex organic synthesis [21]. IMs are advantageous one-step ion exchange method without requiring complex organic synthesis [21]. IMs are advantageous one-step ion exchange method without requiring complex organic synthesis. To the advantage of the period of

the absorption spectra of the resulting IM for potential DSSC application. To investigate FRET between donor NBA<sup>+</sup> and acceptor IR820<sup>-</sup> dyes, the photophysical properties of Nile Blue A IR820 ([NBA][IR820]) IM, parent dyes, and the mixture of parent dyes are studied in detailed. Parent dyes as well as the mixture of parent dyes contain counterion (sulfate, SO<sub>4</sub><sup>2-</sup> and sodium, Na<sup>+</sup>), while in the IM, all counterions have been removed and

it contains only two ionic sensitizer dyes (NBA<sup>+</sup> and IR820<sup>-</sup>). This work also shines light on the effect of spectator ions in FRET mechanism between two ionic dyes.

# 2. Materials and Methods

#### 2.1. Materials

2-[2-[2-Cholro-3-[[1,3-dihydro-1,1-dimethyl-3-(4-sulfobutyl)-2H-benzo[e]indol-2-ylidene]-ethylidene-1-cyclohexen-1-yl]-ethenyl]-1,1-dimethyl-3-(4-sulfobutyl)-1H-benzo [e]indolium hydroxide inner salt (NaIR820), Nile Blue A sulfate (NBA $_2$ SO $_4$ ), deuterated dimethyl sulfoxide (d-DMSO), and Wilmad 5 mm NMR tubes were purchased from Sigma-Aldrich. Ethanol and dichloromethane (DCM) were purchased from VWR. Alumina cells for thermal stability analysis were purchased from Perkin Elmer. Distilled deionized water of 18.2 M $\Omega$  cm was obtained from ultrapure distilled water purifier (ELGA).

## 2.2. Characterization

The identity of the newly synthesized IM was confirmed using a Shimadzu IT-TOF ESI high-resolution mass spectrometer. Fragments were verified by comparing the mass to charge ratio of the ions to the parent dyes. The structure of the IM was verified using proton nuclear magnetic resonance spectroscopy (<sup>1</sup>H-NMR) recorded in d-DMSO with a 300 MHz JEOL ECX instrument. Thermogravimetric analysis (TGA) was preformed using a Mettler Toledo instrument to investigate the thermal stability of the IM. Samples were heated in air at a rate of 10 °C per minute from 25–800 °C.

## 2.3. Absorbance Studies

Samples of parent dyes (NBA<sub>2</sub>SO<sub>4</sub>, NaIR820), the IM ([NBA][IR820]), and the mixture of parent dyes (0.5 NBA<sub>2</sub>SO<sub>4</sub>:1 NaIR820) were prepared in ethanol. Since the IM contains NBA<sup>+</sup> and IR820<sup>-</sup> in 1:1 ratio, a mixture was prepared by taking 0.5 mol eq. of NBA<sub>2</sub>SO<sub>4</sub> with 1 mol eq. of NaIR820 to acquire 1:1 NBA<sup>+</sup> and IR820<sup>-</sup> in the mixture. The Shimadzu UV-Vis-NIR spectrophotometer UV-3600 was used to study their absorption characteristics. A 10 mm path length, polished two-sided quartz cuvette (Starna Cells) was used for absorbance measurements against an identical cell filled with the solvent of the sample.

## 2.4. Photoluminescence and Quantum Yield Studies

The fluorescence emission spectra of all samples were recorded at their respective excitation wavelength using the HORIBA Jobin Yvon Nanolog fluorimeter. A 10 mm path length, polished four-sided quartz cuvette (Starna Cells) was used for fluorescence emission measurements. All samples were measured with the same slit width, integration time of 0.1 s, and at right angle geometry. Absolute quantum yield was recorded using a HORIBA integrating sphere at two different excitation wavelengths.

# 2.5. FRET Calculations

All the FRET parameters are calculated to assess the FRET phenomenon in IM. In order to calculate the spectral overlap integral, theoretical Förster distance, and FRET efficiency of the IM, several equations were employed.

First, the spectral overlap integral  $(J(\lambda))$  was quantified using Equation (1).

$$J(\lambda) = \frac{\int_{0}^{\infty} \varepsilon(\lambda) f(\lambda) \lambda^{4} d\lambda}{\int_{0}^{\infty} f(\lambda) d\lambda}$$
(1)

where  $\varepsilon(\lambda)$  is the molar extinction coefficient (M<sup>-1</sup>cm<sup>-1</sup>) of NaIR820 at overlap wavelength  $\lambda$ , and  $f(\lambda)$  is the normalized fluorescence emission intensity of NBA<sub>2</sub>SO<sub>4</sub> at overlap wavelength  $\lambda$  with NaIR820 when excited at 629 nm [26]. A higher  $J(\lambda)$  value supports the energy transfer between donor and acceptor via FRET mechanism.

Next, the theoretical Förster distance  $(R_0)$  was determined using Equation (2).

$$R_0 = 0.0211(n^{-4} \times k^2 \times \Phi_d \times J)^{6}$$
 (2)

where  $\Phi_d$  is quantum yield of the donor, NBA<sub>2</sub>SO<sub>4</sub>, in the absence of acceptor, NaIR820, n is refractive index of the media, J is the spectral overlap integral between NBA<sub>2</sub>SO<sub>4</sub> and NaIR820, and  $k^2$  is the dipole orientation factor [27]. The dipole orientation factor is conventionally estimated as 2/3 [19].

Finally, the FRET efficiency of the IM was calculated using Equation (3).

$$E = 1 - \frac{F_{da}}{F_d} \tag{3}$$

where  $F_{da}$  is integrated fluorescence emission of the donor in the presence of the acceptor and  $F_d$  is the integrated fluorescence emission of the donor in the absence of the acceptor [28].

# 2.6. Temperature Dependent Photophysical Characterization

To determine the high operating temperatures of a DSSC effect on the FRET parameters of the IM, temperature studies were performed. A sample was placed in a hot water bath on a Corning PC-420D hot plate. The temperature was measured with a Thomas Scientific Traceable Thermocouple. The absorption and fluorescence emission spectra were recorded at 25, 40, 50, 60, and 70 °C. These resulted spectra were used to calculate all above mentioned FRET parameters at different temperatures.

# 2.7. Synthesis

The ionic compound, [NBA][IR820], was synthesized by a previously reported biphasic ion exchange method [29]. Briefly, the NaIR820 parent compound was dissolved in deionized water and the NBA<sub>2</sub>SO<sub>4</sub> parent compound in DCM. The DCM solution containing the NBA2SO4 was roughly five times the volume of the aqueous NaIR820 solution and contained a 0.5:1 molee ratio of NBA<sub>2</sub>SO<sub>4</sub> to NaIR820 parent dyes, which produces 1:1 ratio of NBA<sup>+</sup> and IR820<sup>-</sup> in the IM (Equation (4)). The aqueous and organic solutions were combined in a single vessel and stirred for 72 h at room temperature. After agitation, ethanol was added to the biphasic mixture of the aqueous and the organic solvents to pull any remaining aqueous IR820 into the DCM layer. Since ethanol is miscible in both solvents, it was used as an emulsifying agent, along with intense agitation and centrifugation. This process was repeated multiple times until no color in the aqueous layer was observed, thus effectively pushing all aqueous IR820 into the DCM layer. The DCM solution was washed with deionized water multiple times to effectively remove all Na+ and  ${\rm SO_4}^{2-}$  spectator ions. The DCM was removed by evaporation in a reduced pressure environment using rotary evaporator. The remaining IM was then freeze-dried to remove any trace water impurities. The ion exchange reaction for IM synthesis is shown in the balanced chemical Equation (4), and the structures of donor NBA cation and acceptor IR820 anion are illustrated in Scheme 1. The presence of both ions in the newly designed IM was confirmed by mass spectrometry (Figure S1). The observed peaks in positive and negative ion mode were 318.1597 and 827.2946, respectively. This matched well with the calculated molecular weight of NBA+ ion (318.1601) and IR820- ion (827.2950). The structure of the IM was confirmed by <sup>1</sup>H-NMR. Figure S2 presents the <sup>1</sup>H-NMR spectra which shows the characteristic peaks of hydrogen present in IR820<sup>-</sup> and NBA<sup>+</sup> contained in the IM [30,31].

$$2\,\text{NaIR820}_{\text{(aq)}} + \text{NBA}_2\text{SO}_{4\,\text{(DCM)}} \overset{\text{stirred 72 hr RT}}{\rightarrow} 2\,\text{[NBA][IR820]}_{\text{(DCM)}} + \text{Na}_2\text{SO}_{4\,\text{(aq)}} \quad \text{(4)}$$

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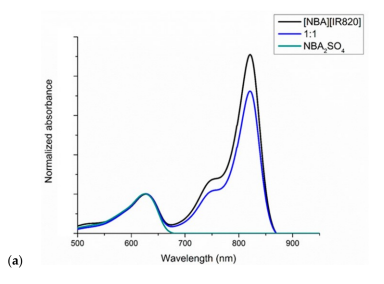
3. Results and Discussion

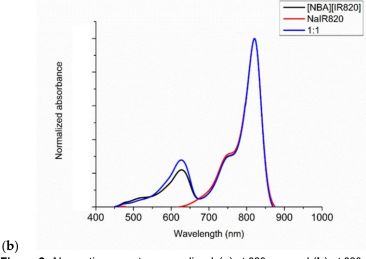
3. N.a.Marenas, Characters at iona 2 NB IR820 IR820 IN Na S (20)

(4)

The melting point of the [NBA][IR820] compound was recorded to be 240 °C. Then, further characterization of the IM was performed.

3. Rusther characterization of the IM was performed. Thermal stability of a photosensitizer is a critically important parameter when used 3.1. Materials Characterization for DSSCs application. Most organic compounds degrade at low temperatures [32,33]. This exberimentinsperior nethological examiller all momentures that the contract of further har sals in the stage of the stage o neathranal depolitions out and rewritten 25-2000 to call shown out this use so can take support and the call shows the call sh  ${
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**Figure 2.** Absorption spectra normalized (**a**) at 629 nm and (**b**) at 820 nm in ethanol. **Figure 2.** Absorption spectra normalized (**a**) at 629 nm and (**b**) at 820 nm in ethanol.

The fluorescence emission spectra of NBA<sub>2</sub>SO<sub>4</sub>, NaIR820, the mixture, and [NBA][IR820] IMake reported in the policy two different excitation visites and all policy of the po IN the manufaction althous some commentation or set to a with respect to parent which some increase of the commentation of the กท**่า**ลก็**เ**ย**ระ** ใกm. All the normalized fluorescence emission spectra with respect to parent NBA2S All BAscroptaining compounds exhibited the emission wavelength maxima at 664 nm when excited at 629 nm. Additionally, a peak at 845 nm was observed in all IR820<sup>-</sup>-containing compounds. The peak shape and fluorescence emission wavelength maxima did not significantly change in different molecules containing NBA<sup>+</sup> and IR820<sup>-</sup> ions. The NaIR820 compound did not show a fluorescence emission peak at 664 nm upon excitation at 629 nm wavelength due to the absence of NBA<sup>+</sup>. However, both the mixture and the IM exhibited the intense emission peak at 664 nm due to the presence of NBA cation. All fluorescence emission graphs were normalized with respect to the NBA2SO4 at 664 nm. It was examined that the fluorescence emission intensity of NBA+ is significantly decreased in the IM in comparison to the parent NBA<sub>2</sub>SO<sub>4</sub> compound when excited at 629 nm (NBA<sup>+</sup> excitation wavelength, Figure 3). However, the 1:1 mixture of the dyes did not show any reduction in fluorescence emission intensity regarding the parent NBA<sub>2</sub>SO<sub>4</sub> compound. Additionally, an increase in emission peak intensity at 845 nm is observed in the IM with respect to the parent NaIR820 dye when excited at 629 nm. The mixture of two dyes did not exhibit any difference in emission intensity with reference to the NaIR820 parent compound. When these compounds were excited at 820 nm (IR820 - wavelength maxima), the IM and the mixture exhibited the same intensity (Figure S5). Detailed analysis

of absorption and the fluorescence spectra demonstrate that IM photodynamic properties are significantly different than the mixture. The significant changes in the IM such as the increase in IR820<sup>-</sup> absorbance at 820 nm and decrease in emission intensity of NBA<sup>+</sup> at Sustain. Chem. 2021, 2, FOR PEER REMEMORAL as well as the enhanced fluorescence emission intensity of IR820<sup>-</sup> at 845 nm when excited at 629 nm led us to investigate the FRET phenomenon in the IM.

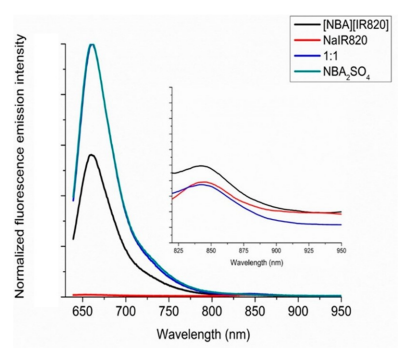


Figure 3. Normalized (with respect to parent NBASEQ 4.01564.htm) if was scarce from its in parent NBASEQ 4.01564.htm) if was scarce from its increase is in of the BASEQ AND BASEQ AND IN A BASEQ AND IN

The wiste funcascence amissing spectrated NRAs nashibit waverlapping with the 864 somptionspectified ar 820 nas. Asplayedain, Figureak Therefore, it is possible that NBAR 820serve asia denotive hereas traeque a characterismo acceptor inca ERES i compovered Jihus abarea ia and specifically control in the sign of the second control in the fluorespect of the sign of the si of NR620 (donor) as well as the enhanced absorbance and therescap esemission intensity of ureson (acceptor) in the UM to the changes in the Lyo rescence emission intensity for dragon are and we have the control of the c mentioned earlier, the spectral overlap between donor fluorescence emission and acceptor fluorescence emission graphs were normalized with respect to the NBA2SO4 at 664 nm. It absorption spectra is the primary requirement of FRET. As a result of FRET was examined that the fluorescence emission peak arising from the donor (NBA) should exhibit a decrease emission peak arising from the donor (INBA) snould exhibit a decrease in intensity due in the IM in comparison to the parent NBA<sub>2</sub>SO<sub>4</sub> compound when excited at 629 nm (NBA) In the IM in comparison to the parent NBA25O4 compound when excited at 629 nm (NBA25O4 to the energy being transferred to the acceptor mojety that consequently enhances the excitation wavelength, Figure 3). However, the 11 mixture of the dyes did not show any fluorescence emission intensity of the acceptor (IR820 ) in the FRET process [34]. This reduction in fluorescence emission intensity regarding the parent NBA25O4 compound phenomenon is observed in Figure 3. In addition to spectral overlap, the Förster distance Additionally, an increase in emission peak intensity at 845 nm is observed in the IM with between the donor and acceptor jons is a critical requirement for FRET, as previously stated the parent NaIR82O dye when excited at 629 nm. The mixture of two dyes of the parent NaIR82O dye when excited at 629 nm. The mixture of dye not exhibit any difference in emission intensity with reference to the NaIR82O parent comprevents the donor and acceptor ions from being in close proximity. However, the absence pound. When these compounds were excited at 820 pm. (IR82O) wavelength maxima), the parent parent of the counteriors are provided at 820 pm. (IR82O) wavelength maxima), the parent parent provided at 820 pm. (IR82O) wavelength maxima), the parent parent provided at 820 pm. (IR82O) wavelength maxima), the parent provided at 820 pm. racteristics. Detailed analysis of absorption and the fluorescence spectra demonstrate that IM photodynamic properties are significantly different than the mixture. The significant changes in the IM such as the increase in IR820 absorbance at 820 nm and decrease in emission intensity of NBA at 664 nm, as well as the enhanced fluorescence emission intensity of IR820- at 845 nm when excited at 629 nm led us to investigate the FRET phenomenon in the IM.

The wide fluorescence emission spectra of NBA<sup>+</sup> exhibits overlapping with the absorption spectra of IR820<sup>-</sup>, as displayed in Figure 4. Therefore, it is possible that NBA<sup>+</sup> can serve as a donor whereas IR820<sup>-</sup> can act as an acceptor in a FRET compound. Thus, there is a possibility of FRET which caused the reduction in the fluorescence emission intensity of NBA<sup>+</sup> (donor) as well as the enhanced absorbance and fluorescence emission intensity of IR820<sup>-</sup> (acceptor) in the IM. The changes in the fluorescence emission intensity for IR820<sup>-</sup>

are only observed when excited at donor (NBA+) excitation wavelength (629 nm). As men-

fluorescence emission intensity of the acceptor (IR820<sup>-</sup>) in the FRET process [34]. This phenomenon is observed in Figure 3. In addition to spectral overlap, the Förster distance between the donor and acceptor ions is a critical requirement for FRET, as previously stated. It is possible that the presence of the counterions, Na<sup>+</sup> and SO<sub>4</sub><sup>2-</sup>, in the mixture of dyes 571 prevents the donor and acceptor ions from being in close proximity. However, the absence of spectator ions in IM provides strong evidence of FRET characteristics.

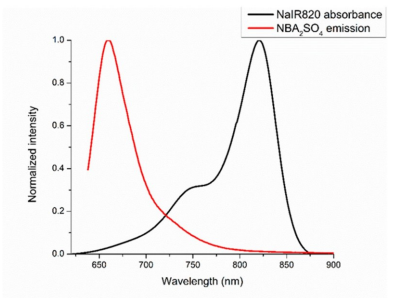


Figure 4. Normalized fluorescence emission spectra of NBA<sub>2</sub>SO<sub>4</sub> (donor) and absorption spectra of Figure 1R820 (acceptor) In ethanol.

The fluorescence emission spectra for all compounds are recorded at 820 nm excitation will veiling the assume in the cinsiper peak is hip pean and emission wavedength 820 xime (&45 nm) tion is abselved in a Re201. - Ebertaining pyck (Flighter 66) eithes reason for ength angelinat (&45 uonm) rescobse and stion in the thinger of the (Flixture 86) d The ate &20 nm reconstation and a value begin flucture and the harden and the content of the

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that FRET is occurring in IM but not in the mixture of two dyes that are used to develop the IM BALLIR 201 is the absence and the presence of two dyes that are used to develop the IM. The only difference in the free interesting that the respectively, thus indicating that the removal of counterions helps to attain the FRET mechanism in the IM, which was not observed in the mixture of parent compounds. As stated earlier, the spectral overlap between the donor 's emission and the acceptor's absorption spectra is not the only criteria for FRET phenomenon; the optimum distance between the donor and acceptor moieties is also required. In the mixture, the presence of counterions prevents the donor and acceptor moieties to come in close proximity in the solution. For this reason, the mixture does not show any possibility for the FRET mechanism to occur, while the absence of counterions in the IM aids to bring the donor and acceptor ions in close proximity (<10 nm). Previous works have confirmed that the ions in the IMs remain associated in the solution (as ion pairs) by observing conductivity inconsistent with the Nernst–Einstein approximation, indicating that there is a net zero charge for the ion pair [38]. Therefore, energy transfer between the two ionic species in an IM is possible.

## 3.3. FRET Calculations

The quantum yield and other spectral data demonstrated the possibility of FRET in the IM. Therefore, all FRET calculations such as overlap integral, theoretical Förster distance between donor and acceptor, and FRET efficiency are calculated for the IM. The results for all FRET calculations are presented in Table 1. First, we observed the spectral overlap (Figure 4), and the spectral overlap integral between donor NBA<sub>2</sub>SO<sub>4</sub> dye and acceptor NaIR820 dye was calculated using Equation (1) [26]. The resultant value of spectral overlap reported in Table 1 indicates moderate overlap between donor and acceptor ions in IM. Nevertheless, it can be improved by varying the ions, which shows better spectral overlap to develop FRET-based IMs. The beauty of IM is the simplicity of tuning the properties in a compound by using task-specific ions. Their simple synthesis makes them ideal to develop more FRET-based IMs in the future for a specific application.

Table 1. FRET parameters: FRET efficiency, theoretical Förster distance, and spectral overlap integral.

% FRET efficiency	31.20%
Förster distance (R <sub>0</sub> , nm)	5.60
Spectral Overlap Integral (J(λ)/M <sup>-1</sup> cm <sup>-1</sup> nm <sup>4</sup> )	$6.83 \times 10^{15}$

Next, the Förster distance in the IM was also calculated to prove the reasoning of the FRET process in the IM in contrast with the mixture of two dyes. For an efficient energy transfer, NBA $^+$  and IR820 $^-$  photosensitizers must be within 10 nm of each other. To calculate this theoretical Förster distance ( $R_0$ ), Equation (2) is employed. The resulting value proves that donor and acceptor ions are present in close proximity in the IM.

These calculated values confirmed the FRET mechanism in the IM. The IM exhibits the possibility of the FRET mechanism when there is significant spectral overlap between two ions present, which is verified by the spectral overlap integral value (Table 1). The spectral overlap is also possible in the mixture. However, the presence of counterions in the mixture hindered the dyes from acquiring the minimum distance required for energy transfer to occur. Therefore, the FRET mechanism was not observed in the photophysical characterization results for the mixture. Thus, removing the spectator counterions is crucial to bringing the ions in close enough proximity to acquire FRET in an IM. One cannot obtain the FRET mechanism in an ionic compound just by mixing a donor compound with an acceptor compound. The removal of counterions is the key parameter to attain FRET characteristics in an ionic compound.

Finally, the FRET efficiency was calculated between donor and acceptor ions in the IM. For that, Equation (3) was employed. To determine FRET efficiency, the integrated fluorescence emission of the NBA<sub>2</sub>SO<sub>4</sub> parent compound was compared to the integrated fluorescence emission of the [NBA][IR820] IM. The fluorescence spectra of NBA<sup>+</sup> in absence (NBA<sub>2</sub>SO<sub>4</sub>) and presence of the IR820<sup>-</sup> ([NBA][IR820]) are displayed in Figure 3. The decreased fluorescence emission intensity of NBA<sup>+</sup> in [NBA][IR820] IM in comparison to the parent NBA<sub>2</sub>SO<sub>4</sub> fluorescence emission revealed that energy is transferred from NBA<sup>+</sup> to IR820 in the IM [39]. When the mixture of two dyes is compared to the NBA2SO4 parent compound, there is no observable decrease in fluorescence emission intensity. This concludes that no energy transfer is occurring in the mixture. The present FRET efficiency is tabulated in Table 1, along with the theoretical Förster distance and spectral overlap integral. All the resulting values for overlap integral and Förster distance between two ionic moieties supports the FRET in the IM. Nevertheless, the FRET efficiency is almost 32%, due to minimum spectral overlap between donor and acceptor ions, as observed in Figure 4 and obtained from Equation (1). The FRET efficiency value can be improved further by maximizing spectral overlap between the two ionic moieties in an IM that serves as a donor and an acceptor. The results obtained for the first IM-based FRET compound demonstrate that the FRET process can be achieved using a simple synthetic ion exchange

approach. This class of IMs can be exploited to develop other FRET compounds for a variety of applications by selecting task-specific ions in the IM synthesis. To the best of our knowledge, this is the first study that reports that an IM can be designed to acquire FRET characteristics using a very facile, prompt, and cost-effective synthesis approach.

# 3.4. Temperature Studies

Since DSSCs are exposed to sun, it is easy to see the temperature change in the solar cells. The FRET phenomenon in IM improves the spectral properties of the dye in DSSCs. However, it is very important to investigate the temperature-dependent FRET variables. Therefore, absorbance and fluorescence emission spectra of all samples were recorded at different temperatures. The results demonstrated minimal changes in intensity for the parent dyes and the IM at different temperature, as displayed in Figures S6 and S7. This finding also validates the thermal stability acquired from TGA (Figure S3), where all samples showed little degradation around ~290 °C. The insignificant degradation at an elevated temperature and stable photophysical properties demonstrate the suitability of organic dye-based IM for high-temperature applications.

As for the effect of temperature on the FRET parameters, minor changes in the FRET parameters values were observed from room temperature to 70  $^{\circ}$ C (Table 2). The spectral overlap integral values are in the same order of magnitude at different temperatures. Similarly, the  $R_0$  values are very consistent. In contrast to the Förster distance and spectral overlap integral, the FRET efficiencies on the other hand are more sporadic. The maximum FRET efficiency is attained at 50  $^{\circ}$ C while at other temperatures, these FRET efficiency values are relatively similar.

**Table 2.** FRET parameters: FRET efficiency, theoretical Förster distance, and spectral overlap integral for a range of temperatures.

	25 °C	40 °C	50 °C	60 °C	70 °C
% FRET efficiency	28.82	26.19	36.16	24.04	20.58
Förster distance (R <sub>0</sub> , nm)	5.64	5.62	5.62	5.63	5.82
Spectral Overlap Integral (J(λ)/M <sup>-1</sup> cm <sup>-1</sup> nm <sup>4</sup> )	6.97 × 10 <sup>15</sup>	6.78 × 10 <sup>15</sup>	6.76 × 10 <sup>15</sup>	6.83 × 10 <sup>15</sup>	8.39 × 10 <sup>15</sup>

## 4. Conclusions

This study presents, for the first time, a new class of compound, IM, that exhibits FRET characteristics. The IM approach to developing a compound with FRET characteristics is very simple, rapid, and economical. Furthermore, the tunable characteristics of IMs make them ideal to develop other FRET-based compounds for a variety of applications. The key to obtaining the FRET mechanism in the IM but not in the mixture is proved by comparing the photophysical properties of the IM and the mixture of two parent dyes. The absence of spectator ions is essential to attain the FRET mechanism in the ionic compounds. All FRET parameters are calculated at different temperatures, which shows that IMs exhibit very stable photophysical properties at elevated temperatures and are thermally stable. For the first time, the concept of FRET is introduced in an IM by pairing an NBA cation with an IR820 anion. All required parameters demonstrate the FRET characteristics of [NBA][IR820] IM. In the future, the simple strategy to design FRET-based IM with enhanced efficiency can further be improved by selecting different ions with greater spectral overlap.

**Supplementary Materials:** The following are available online at https://www.mdpi.com/article/10.3390/suschem2040031/s1, Figure S1: Mass Spectra of [NBA][IR820], Figure S2: NMR spectra of [NBA][IR820], Figure S3: Thermal stability curve for [NBA][IR820], NaIR820, NBA<sub>2</sub> SO<sub>4</sub>, Figure S4: Absorbance spectra of NBA<sub>2</sub> SO<sub>4</sub> and NaIR820 parent dyes, [NBA][IR820], and 1:1 mixture of dye content from parent compounds (0.5 NBA<sub>2</sub> SO<sub>4</sub>:1 NaIR820) in ethanol, Figure S5: Fluorescence emission spectra at 820 nm excitation wavelength, Figure S6 Absorbance spectra of NBA<sub>2</sub> SO<sub>4</sub> (a),

[NBA][IR820] (b), and NaIR820 (c) at various temperatures, Figure S7 Fluorescence emission spectra of NBA $_2$  SO $_4$  excited at 629 nm (a), [NBA][IR820] excited at 629 nm (b), [NBA][IR820] excited at 820 nm (c), and NaIR820 excited at 820 nm (d) at various temperatures.

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