Research Article

OPTICAL PROPERTIES PYRIMIDINE DERIVATIVES: EFFECT OF ELECTRON-DONOR/AcCEPTOR SUBSTITUENTS IN MOLECULAR TOPOLOGY

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ABSTRACT

In this work, we describe the synthetic approach of pyrimidine-substituted derivatives, photoluminescence properties and theoretical calculations of the geometry optimization and electron density distribution on the molecules in the ground and excited states. According to calculations, the location of HOMO electron density is mainly located over the molecular region with highest electron-donor character (or lower electron-acceptor). On the other hand, the electronic transition from HOMO to LUMO promotes a change in the electron density distribution on the molecule. The simulations also reveal that plane of all aryl substituent’s located at positions 2 and 6, are out of the pyrimidine ring plane and the absolute dihedral angles values between π-(p-phenyl) and pyrimidine ring planes are intrinsically dependent on the spatial HOMO distribution on the phenyl ring and p-substituent group. The higher donor character of the substituent leads to substantial changes in the absorption spectra. We also notice that the absorption changes are intrinsically related to the portion of ICT in the molecules, which are strongly dependent on polar interaction. All compounds present large Stokes shifts, higher than 3.370 cm⁻¹, and the emission spectra in solution present broad bands in the ultraviolet-visible region. On the other hand, the solid-state luminescence of 3a and 3c compounds shows emission band in the near-yellow color region 0.423, 0.475 (x, y) and 0.485, 0.542, respectively, in which the 5a and 5c compounds exhibit emission bands corresponding to the near-red color region 0.569, 0.415 and 0.608, 0.389, respectively.

Keywords: Pyrimidines, photophysical properties, theoretical study, effect of electron-donor/acceptor, solvatochromism

INTRODUCTION

Over the years, heterocyclic compounds have received attention in both academic and industry due to their optical and biological properties. Outstandingly, azaheterocycles are known to presents strong aromaticity, significant electron-deficiency, pH sensitivity and capability of nitrogen atoms act in chelation processes, resulting in lower energy band gaps. This characteristic profile proved to be suitable for several applications such as phosphor compounds, dyes, organic light emitting diodes (OLEDs), solar cells, among others. In particular, the pyrimidine ring is frequently reported as strong electron-withdrawing units in donor-acceptor π-conjugated structures (push-pull system) that results in higher optical activities. It has long been known that UV-vis/IR optical properties are intrinsically influenced by the chemical environment and different solvent types can bring out changes in position, intensity, and shape of absorption spectra. Solvatochromism studies, in these also called diazo dyes, have indicated bathochromic (positive solvatochromism) and hypsochromic (negative solvatochromism) behaviors upon increasing of the solvent polarity. However, experimental and theoretical investigations about solvent-solute interactions are one of the most difficult tasks, given the interaction complexity involved.

In the light of the above mentioned, our work report on the photoluminescence azaheterocycles derivatives series based on a 5-cyanopyrimidines core and 2,4,6-Trisubstituted periphery. Starting from the synthesized molecules are reported in literature, we decide emphasized the study into two sections. The investigation of optical properties is provided by the electron-deficiency in the pyrimidine ring in front of different substituents. In such a way, theoretical computational methods were used to investigate the bathochromic effect in different solvents.

MATERIALS AND METHODS

The photoluminescence measurements were performed in a spectrofluorometer Horiba-Jobin-Yvon Fluorolog-3 using as excitation source a xenon lamp continuous 450 W xenon lamp and UV xenon flash tube. The detection was made by a Hamamatsu R 928 P photomultiplier. The spectral correction was made by spectral response of the monochromators and using a silicon photodiode reference to monitor and compensate the power output of xenon lamp, using typical correction spectra.
provided by the manufacturer. UV-vis absorption measurements were performed in a UV-2600 from Shimadzu.

**Molecular orbital and dihedral angles calculations**

Theoretical calculations of the compounds 3a-f and 5a-c were conducted using semi empirical and Density Functional Theory (DFT). The full geometry optimization was carried out using PM3 Hamiltonian as incorporated in Gaussian 09 package. No symmetry or internal coordinate’s constraints were applied during the optimization. The ground states were calculated with the B3LYP functional employments a 6-31G* basis set provided by Games s 64 software package. Additionally, images of HOMO (Highest Occupied Molecular Orbital) and LUMO (Lowest Unoccupied Molecular Orbital) distribution were rendered using Avogadro Software.

**RESULTS AND DISCUSSION**

**Molecular orbital calculations**

Nine compounds pyrimidine derivatives, named as 3a-f and 5a-c, were successfully synthesized using the experimental conditions displayed in Scheme 1.

To understand better the photophysical properties of the proposed molecules, the optimized molecular geometries and, frontier molecular orbital energy diagram and location were calculated, and the results displayed in Figures 1. The electron donates effect were investigate for the p-substituents in the n-(p-phenyl) groups in the positions 2 and 6 of pyrimidine center; could be noted that the HOMO electron density is mainly located over the molecular region with highest electron-donor character (or lower electron-acceptor). The electronic transition from HOMO to LUMO, or vice versa, promotes a change in the electron density distribution on the molecule, relatively to HOMO distribution. In this context, we notice three different orbital behaviors in our simulations: The first one, both HOMO and LUMO presents electron density distribution fullest extended over the molecule induced by chloride substituent, as seen to the 3a, 3b and 3d molecules. For second, HOMO is mainly located above of p-phenyl substituent whereas LUMO distribution extends mainly over the pyrimidine ring. This behavior was observed for 3c, 5a and 5b. On the third one, HOMO are located over a region opposite and distinct from the LUMO region. This could be observed in Figure 1(e) e and 1(i), which shows HOMO located over the p-phenyl containing the group with the lowest electron-acceptor character. In another hand, LUMO appears under the 6-(p-phenyl) substituent.
Figure 1: Calculated spatial distributions of frontier molecular orbital (HOMO and LUMO) and their energy gap for 3a-3f (a)-(f), and 5a-5c (g)-(i) molecules.

Figure 2: Absolute dihedral angles for 3a-3f (a)-(f), and 5a-5c (g)-(i) molecules.
The simulations also reveal that all aryl substituents, located at positions 2 and 6, are out of pyrimidine ring plane. We observed that the absolute dihedral angles values (Figure 2), correlated with n-(p-phenyl) and pyrimidine ring planes (Figure 3) are intrinsically dependent of the spatial HOMO distribution on the p-phenyl, mainly on the p-substituent group. Further, lower dihedral angle was observed in the range of 14.6° until 34.9° and corresponds to the molecular region which the HOMO is predominantly located. In a surprising way the 2-(p-phenyl) position shows lower torsion angles for all compounds. In contrast, the highest dihedral angles, ranging from 49.8° to 89.0°, were related to the 6-(p-phenyl) position that clearly suffers distortion influence from the 5-carbonitrile electron density. It’s should be the point that the small dihedral angles for 5a (14.6°), 3f (15.7°) and 5b (16.6°) molecules at position 6 of pyrimidine ring exhibit a systematic dependence of character donor-acceptor and electro negativity of p-substituents. Moreover, dihedral angles between 6-(p-phenyl) and p-pyrrolone moieties (in 5a and 5b) show both similar values 51.9° and 50.9°, respectively, while the maleimide moieties and 2-(p-phenyl) presents a torsion angle of 39.2° in the 5c molecule.

To investigate the molecular energy involved, when comparing the energy gap between HOMO and LUMO we observed that electron-donor substituent in 2-(p-phenyl), aligned with the electron-acceptor an isil group at position 6, promotes a higher stabilization of molecule in detriment to other substituents. These trends are in agreement with the behavior observed to 5a and 5b molecules. As a fore mentioned, shows a strong dependence of molecular orbital distribution and energy gap between HOMO-LUMO with nature electron-donor/acceptor of the substituent at position 6 of the pyrimidine ring. This behavior, until then unusual with typical pyrimidine derivatives, can be explained mainly by the presence of amino and cyano groups at positions 4 and 5 of the pyrimidine ring, respectively, that affect their electron-acceptor character.

**Optical properties**

The experimental photophysical properties were performed for the pyrimidine derivatives 3a, 3c, 5a and 5c in solution and solid state, at room temperature. Their absorption and emission spectra in methanol, ethanol, acetonitrile, chloroform and dichloromethane were displayed in Figures 6 and 7, besides that the corresponding spectroscopic data are summarized in Table 1. The absorbance spectra of 3a, 3c, 5a and 5c compounds were obtained by UV-visible absorption spectroscopy. Considering the HOMO→LUMO transitions, the first absorption bands in the spectra of 3c and 5a could be associate to π-π* and intra molecular charge transfer (ICT) transition, while the 3a presents only an ICT transition and 5 π-π*transitions. That information is in agreement with the magnitude of the molar extinction coefficients of the lowest energy band (ε) in Table 1.
The 3c and 5c compounds exhibit different spectral profiles due to changes in solvent polarity. We notice that protic solvents are responsible for larger changes in absorption which indicate a strong sensitivity of the donor groups. In contrast, the 3a and 5a compounds (Figure 1 and 3) do not exhibit significant alterations in absorption profile, as seen in Figure 3. We also notice that the absorption changes are intrinsically related to the portion of ICT in the molecules, which is strongly dependent on the polar interaction. This behavior could be associated with 3c and 5c molecules. On the basis of the behavior of the 5a (Figure 3), we suggest that the mechanism involved in π→π* transition is most pronounced. This induces 5a to a similar behavior observed for 3a molecule, in which the ICT mechanism is absent. Table 1, 3c and 5c show systematically decrease of ε face to increase of solvent polarity. Is seen to 3c molecule a remarkable decrease from 1.7 x 10⁴ (ethanol) to 0.2 x 10⁴ L.M⁻¹.cm⁻¹ (acetanilide), whereas in 5c is from 0.2 L.M⁻¹.cm⁻¹ (acetanilide) to non-detected (acetonitrile). Moreover, 3a exhibit a slight growth of ε, rising from 1.1 x 10⁴ (methanol) to 1.6 x 10⁴ L.M⁻¹.cm⁻¹ (dichloromethane). This behavior is associated with the above-mentioned mechanism and the molecular localization of the frontier orbitals. The 5a molecule does not present a trend in the ε profile, probably correlated with an oscillation between the electronic transition mechanisms (ICT and π→π*), as mentioned before.

Table 1: Photophysical data of 3a, 3c, 5a and 5c in different solvents at room temperature

<table>
<thead>
<tr>
<th>Compound</th>
<th>Solvent</th>
<th>Absorbance</th>
<th>Emission</th>
<th>Stokes shift/cm²</th>
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<tr>
<td></td>
<td></td>
<td>λ₆₃0/nm</td>
<td>λ₆₃1/nm</td>
<td>λ₆₃2/nm</td>
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<td></td>
<td></td>
<td>(ε²/10⁴M⁻¹.cm⁻¹)</td>
<td>(ε²/10⁴M⁻¹.cm⁻¹)</td>
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</tr>
<tr>
<td>3a</td>
<td>Methanol</td>
<td>340 (1.1)</td>
<td>296 (1.5)</td>
<td>384</td>
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<td></td>
<td>Ethanol</td>
<td>342 (1.2)</td>
<td>298 (1.5)</td>
<td>414</td>
</tr>
<tr>
<td></td>
<td>Acetonitrile</td>
<td>338 (1.4)</td>
<td>297 (1.6)</td>
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<tr>
<td></td>
<td>Chloroform</td>
<td>343 (1.6)</td>
<td>302 (1.3)</td>
<td>*</td>
</tr>
<tr>
<td></td>
<td>Dichloromethane</td>
<td>341 (1.6)</td>
<td>301 (1.5)</td>
<td>*</td>
</tr>
<tr>
<td>3c</td>
<td>Methanol</td>
<td>345 (2.6)</td>
<td>279 (1.0)</td>
<td>410</td>
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<tr>
<td></td>
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<td></td>
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<td>274 (0.8)</td>
<td>*</td>
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<td>Chloroform</td>
<td>332 (0.2)</td>
<td>270 (1.0)</td>
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</tr>
<tr>
<td></td>
<td>Dichloromethane</td>
<td>327 (0.1)</td>
<td>271 (0.7)</td>
<td>*</td>
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<tr>
<td>5a</td>
<td>Methanol</td>
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<td>275 (5.0)</td>
<td>460</td>
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<tr>
<td></td>
<td>Ethanol</td>
<td>337 (0.8)</td>
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<td>Acetonitrile</td>
<td>332 (0.9)</td>
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<td>Chloroform</td>
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<td></td>
<td>Dichloromethane</td>
<td>331 (1.0)</td>
<td>280 (3.2)</td>
<td>*</td>
</tr>
<tr>
<td>5c</td>
<td>Methanol</td>
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<tr>
<td></td>
<td>Ethanol</td>
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<td>254 (1.5)</td>
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<tr>
<td></td>
<td>Acetonitrile</td>
<td>*</td>
<td>258 (0.4)</td>
<td>*</td>
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<tr>
<td></td>
<td>Chloroform</td>
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<td>257 (0.2)</td>
<td>*</td>
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<tr>
<td></td>
<td>Dichloromethane</td>
<td>*</td>
<td>256 (0.4)</td>
<td>*</td>
</tr>
</tbody>
</table>

- First excited state (lowest-energy absorption maximum); a*: maximum fluorescence excited at the lowest-energy absorption maximum; b. Second excited state; c. Not detected due to very weak emission; d. No were performed
Figure 4: UV-visible absorption of 3a (a), 3c (b), 5a (c) and 5c (d) ([C] = 1.25x10^{-5} M) in methanol, ethanol, Acetonitrile, chloroform and dichloromethane at room temperature.

Figure 5: Normalized absorption (red and green line), excitation (black line) and emission (blue line) spectra of 3a in methanol (a) and ethanol (b), 3c in methanol (c) and ethanol (d), 5a in methanol (e) and ethanol (f), and 5c in methanol (g) and ethanol (h).

In Figure 6, we investigated the optical properties of compounds 3a, 3c, 5a and 5c in methanol and ethanol using photoluminescence spectroscopy. The emission spectra of all molecules’ present broad bands in the ultraviolet-visible region (UV-vis) with maximum centered peaks ranging from 360 to 460 nm. The excitation spectra were acquired by monitoring emission at the maximum of emission and emission spectra was performed by exciting at a maximum of the first absorption band (Table 1). The absence of fell high energy absorption bands in excitation spectra in all molecules suggests a higher incidence of non-radioactive process, promoting a deactivation of the molecular electronic excited states by mechanisms such molecular collision, cross-relaxation, among others. All molecules present large Stokes shifts, higher than 3.370 cm^{-1} (Table 1). The 3a presents the lowest value that was associated to the low stabilization of the molecular excited state by the solvent, furthering the observed shift. On the other hand, 5a compound shows the higher values of Stokes shift (8.112 cm^{-1} -methanol and 7.696 cm^{-1} -ethanol) and presents the large stabilization for polar solvents in the excited state (Figure 1, Table 1 and Figure 6).
Also, should be noted that the shift values are more prominent for samples measured in ethanol, this because the interactions of LUMO electronic orbitals with the hydrophobic group on solvent molecules leads to a better stabilization effect. In contrast, 3a shows a disparate behavior once that the LUMO distribution allows H-bond interactions with the OH group of the solvent, increasing the stability effect. In Figure 7, the solid-states spectroscopic measurements show lower band gap values in the excited state than the samples in the solvent. Excitation spectra were recorded by monitoring emission at 550, 580, 610 and 600 nm, respectively to 3a, 3c, 5a and 5c molecules. Emission spectra of 3a, 3c, 5a and 5c molecules were performed by excitation at 435, 405, 465 and 515 nm, respectively. The samples 3a and 3c shows emission band in the near-yellow color region and presents the respective color coordinates 0.423, 0.475 and 0.485, 0.542 (x, y) in the chromaticity diagram coordinate in accordance with International Commission of Illumination (CIE) in 1931. Moreover, 5a and 5c compounds exhibit an emission band corresponding to the near-red color region and CIE coordinates as 0.569, 0.415 and 0.608, 0.389, respectively. This change in the photoluminescence color between 3a, c and 5a, c materials are correlated to increased conjugated π bonds with the change of p-substituent.

CONCLUSION

The investigation of the effect of electron donates groups at the positions 2 and 6 of the n-(p-phenyl) ring reveal that the HOMO electron density is mainly located over the molecular region with highest electron-donor character (or lower electron-acceptor). On the other hand, the electronic transition from HOMO to LUMO promotes a change in the electron density distribution on the molecule. The simulations also reveal that all planes of aryl substituents, located at positions 2 and 6, are out of the pyrimidine ring plane. We observed that the absolute dihedral angles values between n-(p-phenyl) and pyrimidine ring planes are intrinsically dependent on the spatial HOMO distribution on the phenyl ring and p-substituent group. It is worth pointing that the small dihedral angles for 5a (14.6°), 3f (15.7°) and 5b (16.6°) molecules at position 6 of pyrimidine ring exhibit a systematic dependence of character donor-acceptor and electro negativity of p-substituents. Comparing the energy gap between HOMO and LUMO we observed that electron-donor substituent in 2-(p-phenyl), aligned with the electron-acceptor anisil group at position 6, promotes a higher stabilization of molecule in detriment to other substituents. The higher donor character of the substituent, leads to substantial changes in the absorption spectra, as seen in Figure 3. We also notice that the absorption changes are intrinsically related to the portion of ICT in the molecules, which are strongly dependent on polar interaction. However, we suggest that this mechanism involved π→π* transition is most pronounced. All compounds present large Stokes shifts, higher than 3.370 cm⁻¹, and their emission spectra present broad bands in the ultraviolet-visible region. Also, the shift values are more prominent for samples measured in ethanol due to the interactions of LUMO electronic orbitals with the hydrophobic groups which solvent molecules leads to a better stabilization effect. The solid-state luminescence of 3a and 3c compounds shows emission band in the near-yellow color region 0.423, 0.475 (x, y) and 0.485, 0.542, respectively, in which the 5a and 5c compounds exhibit emission bands corresponding to the near-red color region 0.569, 0.415 and 0.608, 0.389, respectively. This change in the photoluminescence color between 3a, c and 5a, c materials are correlated to increased conjugated π bonds with the changing of p-substituent.
Figure 7: Normalized excitation (black line) and emission (green and red line) spectra of 3a (a), 3c (b) and 5a (c) and 5c (d) in solid state, and chromaticity diagram (e) with corresponding color coordinates of 3a (1), 3c (2) and 5a (3) and 5c (4)

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