Optical Nature of Non-Substituted Triphenylmethyl Cation: Crystalline State Emission, Thermochromism, and Phosphorescence

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Abstract: Since the discovery of the triphenylmethyl (trityl) cation 120 years ago, a variety of aromatic cations having various colors and luminescence properties have been rigorously studied. Many, differently substituted trityl cations have been synthesized and their optical properties have been elucidated. However, the optical properties of the parent, non-substituted and highly reactive trityl cation, which was observed to be very weakly luminescent, have not been subjected to detailed investigation. In the effort described herein, we explored the optical nature of non-substituted trityl hexafluorophosphate (PF₆) in the crystalline state. Trityl PF₆ was found to exist as two crystal polymorphs including a yellow (Y) and an orange (O) form. Moreover, we observed that these crystalline forms display crystalline-state emission with different colors. The results of X-ray crystallographic analysis showed that the two polymorphs have totally different molecular packing arrangements. Furthermore, an investigation of their optical properties revealed that the O-crystal undergoes a distinct color change to yellow upon cooling as a consequence of a change in the nature of the charge transfer interaction between the cation and PF₆ anion, and that both the Y and O crystal exhibit phosphorescence.

Introduction

Triphenylmethylyl (trityl) cation[1-4] is a classical carbocation that possesses three-fold symmetry and a positive charge that is delocalized over the entire molecule (Figure 1a). Trityl cation, which can be readily prepared by treatment of trityl alcohol with acid, exhibits intense absorption in the visible region. Since the time of the discovery of this cation, organic chemists have conducted extensive studies concentrating on the synthesis and exploration of derivatives including π-extended analogs[5-8] and heteroatom containing (N or O) systems, as well as on developing methods for stabilizing the cationic state toward reactions with nucleophiles, and tuning its absorption and emission properties. Through these efforts, many highly absorbing and emitting derivatives have been uncovered, including those of malachite green, cyan violet, fluorescein and rhodamine derivatives that are utilized as textile dyes, external stimuli responsiveness fluorescent probes and cell imaging reagents[9-11] (Figure 1b). Furthermore, owing to its high reactivity, the non-substituted (NS) trityl cation has been widely used as a Lewis acid catalyst (Mukaiyama-type aldol or Michael reaction), photodissociation promoter, hydride acceptor and protecting group component.[12-15] In contrast, the optical properties of NS trityl cation have not yet been fully explored likely because of its high reactivity.

Although various new aromatic compounds have been synthesized, knowledge about the fundamental properties of reactive aromatic cations is also useful in the development of cationic dye, sensors and emissive materials. It seems appropriate, that now, 120 years from the time of the first report of the trityl cation,[2,3] we have carried out an investigation exploring the absorption and luminescence features of the NS trityl cation. In this effort, we discovered that in the crystalline state NS trityl cation displays bright luminescence, unique thermochromism,[17-20] and phosphorescence[21-25] which are governed by the arrangement of molecules in the crystal.

![Figure 1. (a) Structure of NS trityl cation. (b) Trityl cation based stable dye, fluorescent probe, or emissive imaging materials containing heteroatoms.](image-url)

Results and Discussion

Synthesis and crystallization behavior

The NS trityl cation with PF₆ as the counter anion was prepared by treatment of trityl alcohol with hexafluorophosphoric acid (HPF₆). Specifically, to a solution of trityl alcohol in acetic anhydride at room temperature was added a 60% HPF₆, and the resulting mixture was let stand at room temperature while crystal growth occurred.[26] At first, yellow (Y) crystals formed (Figure 2a, b), but when the solution stood for a several ten minute period, production of orange (O) crystal began (Figure 2c, 2d). Finally, upon standing for days to weeks (Figure 2e-2g), O-crystal are formed as the major component. The single-crystal-to-single-crystal (SCSC) phase transition[27-32] from Y- to O-crystals can be effectively suppressed by keeping the solution at low temperatures.
temperature (4 °C). This phenomenon enabled selective isolation of the two crystal polymorphs.

It is known that the colors of trityl salts in the solid state depend on the nature of the counter anion. It was previously reported that the crystal color of trityl PF₆ is orange. Furthermore, trityl tetrafluoroborate, prepared to evaluate the counter anion effect on the SCSC transition, was found not to exist as crystal polymorphs but only one yellow crystalline form, which has cell parameters that are similar to those of the with Y-crystal of trityl PF₆ (Figure S5). Thus, the nature of the counter anion is crucial for the occurrence of this SCSC phase transition.

The thermal stabilities of the Y- and O-crystals were elucidated by using differential scanning calorimetry (DSC). The scan of the Y-crystal contains lower endothermic peaks at 104 °C (phase transition) and 148 °C (decomposition) and that of O-crystal contains a peak at 168 °C (decomposition) at 293 K. This result indicates that the Y-crystal phase is kinetically preferred and the O-crystal is thermally more stable in a manner that typically exemplifies the Ostwald’s step rule.

Optical properties and crystal structures

In Figure 3a are shown the UV-vis spectrum of a CHCl₃ solution of the NS trityl cation, generated by reaction of trityl alcohol with 2% TFA, and solid-state transmission UV-vis spectra of Y- and O-crystals dispersed in KBr pellets. The color of the solution of the trityl cation is yellow, corresponding to a λmax at 435 and 410 nm (ε = 45000 cm⁻¹ M⁻¹). The UV-vis spectrum of the Y-crystal is nearly identical to that of a solution of the trityl cation whereas the spectrum of the O-crystal contains a distinctively bathochromically shifted maximum above 550 nm. This finding was confirmed by inspection of the solid-state diffuse reflectance spectra of these crystals (Figure S2). Although several aromatic cations containing heteroatoms such as N or O are known to exhibit intense emission,[37,38] those composed of only hydrogen and carbon exhibit very weak or no emission in solution owing to the favorability of non-radiative decay processes.[39] In fact, the trityl cation in solution displays no emission. On the other hand, both the Y- and O-crystals of trityl PF₆ display relatively intense yellow green (λem = 525 nm) and orange (λem = 585 nm) emission, respectively (Figure 3b). The emission quantum yields of the Y- and O-crystals at 293 K are 4% and 12%, respectively. This phenomenon is a consequence of the aggregation induced emission (AIE) because radiationless decay associated with rotation of the three phenyl rings is effectively suppressed in the crystalline state. To the best of our knowledge, no hydrocarbon cations and only a few aromatic cations containing N and O atoms have been reported to exhibit AIE.

To gain information about the optical differences between the Y- and O-crystals, single crystal X-ray analysis was performed to determine crystal packing structures. The results show that the trityl cation in the Y-crystal at 170 K has D₃ symmetry with a torsion angle of the three phenyl rings being 32.9° (Figure S3). In the unit structure of the Y-crystal, four trityl cations are oriented in a tetrahedral arrangement (Figure 4a, b) and the PF₆ anions are located in a central cavity as well as in areas surrounding the tetrahedron (Figure 4c). Thus, each trityl cation in the Y-crystal is completely isolated from neighboring cations by intervening PF₆ anions and, as a result, no significant interactions exist between the trityl scaffolds. As stated above, the molecular arrangement in the Y-crystal is reflected in its UV-vis spectrum, which is similar to that of the solution state.

Figure 2. Photographs showing crystal growth and a SCSC phase transition from (a) Y-crystal to (h) O-crystal at elapsed times following initiation of crystallization. The blue arrow in (a) indicates a stir tip as a standard material. The black arrows in (c) indicate the beginning of growth of the O-crystal.

Figure 3. (a) UV-vis spectrum of trityl cation (2% TFA in CHCl₃) and solid-state UV-vis spectra of Y- and O-crystals (KBr pellet). (b) Emission spectra of Y- and O-crystals (λex = 360 nm).

Figure 4. X-ray crystallographic analysis of Y-crystal (170 K). (a) Tetrahedral arrangement of trityl cations depicting different colors without PF₆ anions for clarity. (b) Schematic drawing of the tetrahedral arrangement of trityl cations as shown in (a). (c) Drawing with PF₆ anions shown in (a). Protons are omitted for clarity. Ortep drawing is 50% probability.

Figure 5. X-ray crystallographic analysis of O-crystal (195 K). (a) Top view of trityl cation dimer in O-crystal with different colors and without PF₆ anions for clarity. (b) Side view of trityl PF₆ dimer in O-crystal with PF₆ anions. (c) One dimensional alignment of trityl PF₆ (viewing from c axis). Protons are omitted for clarity. Ortep drawing is 50% probability.
In a manner that is different from that of the in the Y-crystal, the trityl cation scaffold in the O-crystal has \( C_1 \) symmetry with the different torsion angles between the three phenyl rings being 25.4, 31.8 and 40.2° (Table S2). Two types of aggregation arrangements exist in the packing structure of the O-crystal. In one of these, two trityl cations form a face-to-face dimer with PF\(_6\) anions at exterior positions (Figure 5a, b). The intermolecular atomic distance between the central sp\(^2\) carbon of the trityl cations is 4.265 Å (195 K), which is 1.78 Å greater than the distance for typical π-π stacking as a result of steric hindrance between the phenyl rings and positive charge repulsion. In the other arrangement, the trityl cations are one dimensionally laterally aligned with a periodic distance of 8.698 Å (Figure 5c). It is noteworthy that the inter-atomic C···F distance between central sp\(^2\) carbon of the trityl cation and fluorine atom in PF\(_6\) anion is 3.101 Å (195 K) (Figure 5b), a value that is almost identical to the sum of the van der Waals (vdWs) radii of carbon (1.70 Å) and fluorine (1.47 Å) atoms.

This observation indicates the existence of a stronger anion-nan interaction than that existing in the Y-crystal (C···F distance is 3.503 Å at 170 K). In fact, the results of natural bond orbital (NBO) analysis of trityl PF\(_6\) monomer in both Y- and O-crystals using MP2/6-311+G* reveals that a charge-transfer (CT) interaction is present between the trityl cation and PF\(_6\) anion and that the degree of the CT interaction in the O-crystal is larger than it is in the Y-crystal (Figure S8). In addition, the NBO analysis suggests that the face-to-face trityl PF\(_6\) dimeric structure in O-crystal enables a greater degree of CT interaction than that which exists in the trityl PF\(_6\) monomer (Figure S9). This likely caused by the unfavorable cation-cation repulsive energy in the face-to-face dimeric structure which enhances CT interaction from the PF\(_6\) anion to the trityl cation.

**Figure 6.** Thermochromism of the O-crystal between room temperature and 77 K. The detail of this thermochromism can be see in Movie S1 (under room light) and Movie S2 (under UV light).

**Figure 7.** (a) Excitation (dash line) and emission (solid line) spectra of O-crystal at 293 K. (b) Excitation (dash line) and emission (solid line) spectra of O-crystal at 77 K. In both excitation spectra, self-absorption from 400 to 450 nm is due to the intense absorption on the crystal surface. \( \lambda_{em} = 360 \) nm for emission measurements, \( \lambda_{em} = 555 \) nm in 293 K and \( \lambda_{em} = 560 \) nm in 77 K for excitation measurements.

**Thermochromism of O-crystal**

Additional investigations led to the discovery of other interesting properties of the O-crystal of trityl PF\(_6\). Upon cooling the temperature from room temperature to 77 K the color of O-crystal changed to yellow and the orange color reformed upon warming to room temperature (Figure 6 and Movie S1). Similar to these absorption changes, the emission color of O-crystal changes from orange to yellow upon cooling (Movie S2). This type of distinctive thermochromism does not take place with the Y-crystal (Movie S3-4, Figure S10). The thermochromic behavior of the O-crystal was evaluated by using emission and excitation spectroscopic measurements (Figure 7a, b). The excitation spectrum at room temperature contains a peak at 560 nm whereas at 77 K the maximum occurs at a wavelength shorter than 560 nm. In addition, the emission peak also shifts from 585 nm to 560 nm when the temperature is lowered from room temperature to 77 K.

To evaluate the relationship between thermochromism and structural changes, single crystal X-ray analysis was conducted on the O-crystal at 273 and 83 K (Figure S5, S6). The results show that upon cooling, no change occurs in the crystal system (monoclinic, \( P_2_1/n \)) and negligible changes take place in the torsion angles of three phenyl rings, as well as in C-C bond lengths between the central sp\(^2\) carbon and ipso-phenyl carbons (Table S2). On the other hand, the inter-atomic C···F distance between fluorine atom of the PF\(_6\) anion and the central sp\(^2\) carbon of trityl cation upon cooling shortens from 3.155 Å (273 K) to 3.047 Å (83 K) (Figure 8a and Figure S5, S6), which is shorter than the sum of vdWs radii of carbon and fluorine atoms. In addition, the distance between the central sp\(^2\) carbons in face-to-face trityl cation dimers in the crystal are slightly reduced from 4.296 Å (273 K) to 4.232 Å (83 K) (Figure 8b) upon cooling. The cause of this change is reduction of the net charge on the trityl cation by strong CT from the PF\(_6\) anion, which is confirmed by using Hirshfeld atomic population analysis (MP2/6-311+G**, Table S3).

TD-DFT calculations of trityl PF\(_6\) in the O-crystal at both 273 and 83 K were carried out using CAM-B3LYP functional with a tuned parameter method (\( \mu = 0.150, \alpha = 0.0.070, \beta = 0.9020 \)) as developed by Nakano\(^{[23]}\). The results reveal that the longer wavelength transitions (430.4, 429.6 and 424.8 nm with respective \( f = 0.0004, 0.0035 \) and 0.0116, Table S4 and Figure S11, S13) of the crystal at 273 K, containing two trityl cations arranged in a face-to-face dimer orientation, are associated with a weak CT transition from the PF\(_6\) anion to the trityl cation. The major band at 83 K corresponds to an intramolecular transition in trityl cation (422.0 nm, \( f \approx 0.0441 \)) because shortening of the C···F distance leads to strengthen of the CT transition and shifting it to higher energy (406.1, 404.3 and 402.7 nm with respective \( f = 0.0052, 0.0039 \) and 0.0438, Table S5 and Figure S12). In contrast, calculations on crystals containing a trityl PF\(_6\) monomer and a laterally arranged trityl PF\(_6\) dimer, show that little change occurs in the longer wavelength transition upon changing the temperature between 273 and 83 K (Table S6-S8 and Figure S14-20). Therefore, the face-to-face dimer structure promoted perturbation of the CT transition energy through a cation-cation repulsive interaction is the cause of the prominent color change from orange to yellow that occurs upon cooling the O-crystal.
Figure 8. Superposition of the structure generated by VT X-ray analysis of O-crystal at 273 K (grey) and 83 K (orange). (a) Depicting the structural changes between PF₆ anion and trityl cation. (b) Depicting the structural changes between face-to-face trityl cation dimer.

Phosphorescence in trityl cation

The trityl cation is isoelectronic with triphenylborane, which displays unique optical properties including thermally activated delayed fluorescence (TADF).[^6-^9] Owing to this relationship, we explored the delayed emission nature of a solid solution of trityl PF₆. The emission spectrum of a solid solution of trityl cation (5.0 x 10⁻⁵ M) in CHCl₃ with 2% TFA at 77 K contains a broad weak peak from 510 to 520 nm and an intense peak at 550 nm (Figure 9a). Delayed emission measurements (10 ms delay) show that the broad and weak 510-520 nm peak disappears and an intense emission band at 550 nm arises. It is noteworthy that the same sample at 195 K contains a broad delayed emission band 515 nm (Figure S2). The broad 510-520 nm and intense 550 nm peaks are assigned as respective fluorescence and phosphorescence of trityl cation. Thus, a remarkably small energy gap (ΔEₛₜ) of 0.15 eV exists between S₁ and T₁ of this substance. The lifetime of the phosphorescence of trityl cation is 310 ms (Figure S23) and the emission quantum yield is 40% (Table 1).

Table 1. Emission peaks, lifetime, and quantum yield of trityl cation (77 K), Y-crystal (293 and 77 K), and O-crystal (293 and 77 K).

<table>
<thead>
<tr>
<th>Sample</th>
<th>λₘₐₓ (nm)</th>
<th>τ (ms)</th>
<th>Φ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Trityl cation (77K)[a]</td>
<td>510, 550</td>
<td>310 ms (550 nm)</td>
<td>40</td>
</tr>
<tr>
<td>Y-crystal (293 K)</td>
<td>525</td>
<td>1.8 ns (525 nm)</td>
<td>4</td>
</tr>
<tr>
<td>Y-crystal (77 K)</td>
<td>510 (SS)[b]</td>
<td>340 ms (550 nm)</td>
<td>11</td>
</tr>
<tr>
<td>O-crystal (293 K)</td>
<td>585</td>
<td>9.1 ns (575 nm)</td>
<td>12</td>
</tr>
<tr>
<td>O-crystal (77 K)</td>
<td>560 (SS)[b]</td>
<td>410 ms (560 nm)</td>
<td>24</td>
</tr>
</tbody>
</table>

[^6-^9]: Triarylborane was used as a reference.

[a] Triarylborane was used as a reference.

In contrast to those of the frozen state, the phosphorescence emission properties of the Y- and O-crystals are totally different. The steady-state (SS) emission spectrum of Y-crystal at 77 K contains an intense peak at 510 nm, which is nearly the same wavelength as that in the fluorescence spectrum of frozen state of trityl cation. The phosphorescence emission spectrum of the Y-crystal contains two peaks at 500 and 585 nm (Figure 9b) associated with an emission lifetime of 340 ms, which is comparable with that of the frozen state. On the other hand, the SS emission spectrum of the O-crystal begins at 480 nm with a maximum at 560 nm. In addition, the SS emission spectrum overlaps with that of its phosphorescence spectrum (Figure 9c).[9] Although the trityl cation has a small ΔEₛₜ, the emission quantum yield of the O-crystal is enhanced by cooling (12% at 293 K to 24% at 77 K), which rules out the existence of TADF. Therefore, emission occurs from the triplet state as phosphorescence, as shown in the SS emission spectrum of the O-crystal at 77 K.

The emission lifetime of the O-crystal at 293 K is 9.1 ns (Figure S24) whereas at 77 K it is 410 ms, which is slightly longer than those of the frozen solution state and the Y-crystal. Longer wavelength emission peaks of 500 nm are present in the phosphorescence emission spectra of the Y-crystal (λₘₐₓ = 585 nm) and O-crystal (λₘₐₓ = 617 nm). Phosphorescence spectra caused by excitation at 550 and 590 nm (for Y-crystal) and 620 nm (for O-crystal), were analyzed. Interestingly, a spectra that has a shape that is similar to that of the an absorption spectrum of trityl cation when excitation occurs at 550 nm whereas sharp longer wavelength shifted excitation spectra of both the Y- and O-crystal are promoted by using 590 and 620 nm excitation, respectively (Figure 10). Furthermore, excitation at 480 nm for Y-crystal and 500 nm for O-crystal leads to selected enhancements of the emission peaks at 580-585 nm for Y-crystal and 620 nm for O-crystal. On the other hand, the emission spectrum of the frozen state displays no dependence on excitation wavelength and concentration (Figure S22).

Therefore, these longer wavelength phosphorescence peaks are attributed from their packing structures such as the arrangement of trityl cations whereas the emission peak at 550 nm is attributed from the monomer structure of trityl cation. Inspection of the ns to μs timescale emission decay profiles of the Y- and O-crystals at 77 K shows that, in contrast to the frozen state, both Y- and O-crystal display intense emission in the nanosecond region (<240 ns) and relatively weak emission in the microsecond region (Figure S25). Owing to the weakness of emission in the microsecond region, analysis of the long time decay up to the millisecond timescale was difficult to perform using our instrumental setup and, thus, the phosphorescence quantum yield could not be determined.

Spin-orbit coupling (SOC) constants were calculated using the TD-DFT method with Tamm-Dancoff approximation (TDA) to evaluate the phosphorescence properties of the trityl cation. It is noteworthy that small SOCs occur between the Sₚ and Tₕ states as well as S₁ and T₁ states of the trityl cation alone (Figure S26) whereas significant SOCs takes place between singlet and triplet states in the ion pair of trityl cation and PF₆ anion (Figure S27-S29). This finding indicates that ISC of the trityl cation is facilitated by the presence of the counter anion. Judging from the energies of the singlet and triplet excited states of trityl PF₆ ion pair in the O-crystal at 273 K, the energies of S₁,₂, which are almost degenerated energy states, are close to those of Tₖ and a large SOC exists between these states, resulting in effective ISC and enabling subsequent internal conversion Tₖ → T₁.

The calculations also show that, although only small differences exist in the degree of SOC between trityl PF₆ ion pairs in the Y- and O-crystals as well as in the O-crystal between 273 and 83 K, a distinctive enhancement of the SOC exists in the face-to-face dimer of trityl PF₆ ion pair structure in the
Figure 10. Phosphorescence excitation spectra and excitation wavelength dependence of phosphorescence of Y-crystal (a), (b) and O-crystal (c), (d). Spectra were measured with a 10 ms delay and 8 ms integration at 77 K.

Figure 9. Steady-state emission and phosphorescence (10 ms delay, 8 ms integration) spectra at 77 K. (a) Frozen state (CHCl₃ with 2% TFA, 5.0 x 10⁻⁶ M) of trityl cation. (b) Y-crystal. (c) O-crystal. (λₑₓ = 360 nm)

Conclusion
In the investigation described above, we uncovered the unique optical nature of the non-substituted trityl cation, which is a fundamental aromatic hydrocarbon cation. The studies show that crystalline trityl PF₆ exhibits a SCSC phase transition from Y- to O-crystals, which originates from CT interactions between the trityl cation and PF₆ anion. In addition, trityl PF₆ displays crystalline state fluorescence and phosphorescence. Importantly, the molecular arrangement of the trityl cation and PF₆ anion in the crystalline state plays a dominant role in governing photophysical properties. This phenomenon was explored by using TD-DFT as well as SOC calculations, which gave a more detailed understanding of how ion arrangements in the Y- and O-crystals govern excited properties. We believe that other hydrocarbon cations have the potential of possessing these unique optical properties. It is anticipated that the results of the current effort will pave the way to an exploration of hydrocarbon cations and their use in organic luminescent devices, and in switching and sensing systems.

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Conflict of interest
The authors declare no conflict of interest.