EMISSIVE ORGANIC RADICALS SUBSTITUTED BY PHOSPHORUS-
CONTAINING MOITIES: APPLICATIONS AS OLED MATERIALS

by

Ragene Thornton, B.S.

A thesis submitted to the Graduate Council of Texas State University in partial fulfillment of the requirements for the degree of Master of Science with a Major in Chemistry August 2019

Committee Members:

Todd Hudnall, Chair

William Brittain, Co-Chair

Benjamin Martin, Co-Chair
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ABSTRACT

OLEDs are organic light emitting diodes, meaning that the emitted light is produced by organic molecules rather than solid-state materials used in LEDs. Typical OLEDs undergo emission events via closed shell systems. Radicals are considered “open shell” systems due to their lack of paired electrons. The maximum internal quantum efficiency (IQE) of an open system is 100% because the presence of only doublet states removes conflict with competing electronic states. This is significantly more efficient than the expected 25% IQE seen from fluorescence events in closed shell systems. The radical of interest in this research is tris (2,4,6-trichlorophenyl) methyl or TTM. The goal of the research at hand is to explore the effects of the introduction of pnictogen heteroles into the TTM radical system. Heteroles display attractive optoelectronic applications in organic electronic devices like OLEDs, organic field-effect transistors, and organo-photovoltaics. Carbazole is a heterole that has been at the forefront of OLED chemistry; expansion of research to the rest of the pnictogen group is significantly rarer. The production and analysis of the various heteroradicals should tell us whether the wavelength of the molecule could be tuned, if the quantum yield be adjusted, and if the stability of these molecules be improved. Due to the prevalence of carbazole in OLED systems, exploration of other pnictogen atoms could prove to be groundbreaking. There was success when the cationic form of TTM was used as the replacement for the TTM radical; although there was success, the goal of the research was not fulfilled because the
product produced did not contain a radical anywhere throughout the molecule.
1. INTRODUCTION

1.1 Organic Electronics and the History of OLEDs

The field of organic electronics has grown over the years, owing its success to the desire to obtain both conductive and light-weight materials to be used in the manufacturing of flexible solar cells and more energy efficient light emitting diodes (LEDs). The three primary areas of organic electronics include organic photovoltaics (OPVs), organic field effect transistors (OFETs), and organic light emitting diodes (OLEDs). Organic photovoltaics are defined as solar cells that use organic molecules for light absorption and charge transport to produce electricity from sunlight via the photovoltaic effect. In Figure 1, the nature inspired precursor is a π-conjugated polymer that is typically found in nature as a dye. Photoexcitation of this molecule proves that the implementation of it into OPVs is not beneficial due the immense loss of energy it undergoes via internal conversion. Functionalizing the nature inspired precursor with thiophene restricts energy loss due to internal conversion and it decreased the bandwidth of the molecule, which in turn increases the absorption rate and promotes easier electron transport.

Figure 1. The use of thiophene in OPVs. Thiophene is used to copolymerize with the host materials to decrease the width of the band gap.

Field effect transistors are the building blocks of various display and integrated
circuit systems; the OFET is the organic counterpart. OFETs are composed of a transistor and an induced electric field to manipulate the electrical behavior of the device.\textsuperscript{9, 10} These systems typically contain polymers or thiophene based molecules due to their semi-conducting abilities. In Figure 2, the exploration of furan-based systems rather than thiophene-based showed that furan is just as beneficial as thiophene in OFET systems due to oxygen’s smaller Van der Waals radius.\textsuperscript{11, 12} Discoveries like these have contributed to increased interest in organo-electronics.

![Figure 2](image_url)

Figure 2. Furan is used in OFET systems due to their semiconducting abilities and dense packing structure.

OLEDs are organic light emitting diodes, where the emitted light is produced by organic molecules rather than solid-state inorganic materials used in traditional LEDs. OLEDs are the organic and subsequently inexpensive and light weight counterpart of LEDs. Moreover, OLEDs can be lighter and more flexible, and they offer the potential for improvements in picture quality when compared to conventional LEDs.\textsuperscript{13, 14} A LED, seen in Figure 3, is defined as a junction diode with a forward bias.\textsuperscript{1} A diode contains a forward bias only if the anodic electrode sustains a positive charge and the cathodic electrode sustains a negative charge. The forward bias in a LED is created by opposing junctions. The n-type (or cathodic) junction produces electrons and the p-type (or anodic)
junction produces positive points that lack electrons, which are referred to as “holes”. As current is produced in the system the electrons and holes travel to opposite electrodes. When the electrons and holes recombine in the junction, they release energy in the form of photons of light.

![Figure 3. A schematic of the electron transport event that takes place in LEDs](www.explainthatstuff.com)

In an OLED (Figure 4) the n and p type junctions are comprised of organic materials. When a voltage is applied to the electrodes in an OLED, a similar electroluminescence event occurs; however, the current flows from the cathode to the anode through multiple layers of organic materials into the emissive layer. The holes are produced at the anode/p-type junction and electrons are produced at the cathode/n-type junction. When the holes and the electrons meet in the emissive layer, they release a burst of energy called a photon.
OLEDs have a rich history. Despite their discovery in the mid-20th century, the ultimate performance of OLEDs has yet to be achieved. Emissive organic materials were first discovered in 1953 by the Bernanose group. The group witnessed light emission after doping a cellulose film with the dying agent acridine orange. The next milestone was achieved in 1963; the Pope group reported light emission from a crystal of anthracene after they induced carrier injection within a high electric field. Although there were other minor discoveries during this time, the research developed did not reveal much about the potential of the light emitting organic materials. This misunderstanding was mainly due to issues with low efficiency and luminescence in conjunction with the need for high electric fields. The next breakthrough was made by Tang and VanSlyke of Eastman Kodak in 1987. The duo reported particularly bright emission from a device with a structure different from earlier OLEDs. The Tang-VanSlyke device contained a pair of thin organic layers housed between two electrodes. The two-layer structure consisted of separate hole transporting and electron transporting layers such that
recombination and light emission occurred in the middle of the organic layer.\textsuperscript{20} This event resulted in a reduction in operating voltage and improvements in efficiency. Tang and VanSlyke reported light emission from a voltage as low as 2.5V and a power efficiency of 1.51 lm/W; in contrast they also reported an external quantum efficiency (EQE) of only 1\%.\textsuperscript{20} Despite this drawback, the device gained lots of attention, both in industrial and academic research fields. The industry grew immensely throughout the years and in 1997 the world’s first commercial OLED was created as the display for a car radio.\textsuperscript{19, 21} A year later, in 1998, phosphorescent OLEDs which utilize triplet emitters were invented.\textsuperscript{22} These discoveries led to the prevalent use of OLEDs in modern day electronics. OLEDs can be found in a variety of digital devices such as television systems, media players, handheld devices and computer monitors. Figure 5 displays the projected growth of the OLED market from 2016-2026.

![Figure 5. The expected increase in the OLED display market over the next couple of decades.\textsuperscript{23}](image)

**1.2 Organic Emissive Layer**

Currently, there are three types of organic materials used in OLEDs: fluorescent, phosphorescent, and thermally activated delayed fluorescent (TADF) materials.\textsuperscript{24-26} The
difference between these types of materials is the mechanism employed to achieve the emission of light. When an electron is excited from the singlet ground state, it can either form a singlet or triplet excited state. The direction of the spin of the excited electron determines whether it is singlet or triplet and transitions of a triplet exciton to the ground state is forbidden due to the Pauli Exclusion Principle. The singlet state (following the Pauli Exclusion Principle) corresponds to a state where the electron spins are paired, and a triplet state corresponds a set of spin unpaired/parallel electrons. Fluorescence occurs when the excited electron relaxes is excited to a higher energy level relaxes to a lower energy state, and then falls back to the ground state releasing a photon of light. Even though fluorescent events occur readily; they lack in efficiency, specifically internal quantum efficiency (IQE). IQE is defined as the ratio of the number of photons collected to the number absorbed. An ideal IQE is as close to 100% as possible; this implies that every photon is absorbed and subsequently collected at the electrodes. As mentioned before, a fluorescent event consists of an electron emitting light when returning to a singlet ground state from a singlet excited state. The lowest triplet excited state is very close in energy to the singlet ground state and statistically triplet states outnumber singlet states 3:1. This causes fluorescent emitters to suffer because of the prevalence of triplet states (Figure 6). Radiative decay from triplet state is spin forbidden, so the max IQE of these events is only 25%. In contrast, phosphorescence occurs when an electron in the triplet excited state relaxes radiatively to the ground state with the release of a photon of light. One form of this radiative relaxation is a phenomenon known as intersystem crossing (ISC). Intersystem crossing occurs when singlet or triplet excited states non-radiatively convert to the opposite type of state. During this time the spin of
the electron is reversed and the resultant radiative decay from the excited triplet state back down to a singlet ground state to release a photon of light is known as phosphorescence.\textsuperscript{31} This type of radiative decay is spin forbidden, however emission through this pathway is promoted through the use of heavy atoms. The presence of heavy atoms affects the photophysical and photochemical processes in a molecule through relativistic effects and spin-orbit coupling which ultimately allows forbidden spin transitions to occur radiatively.\textsuperscript{32} Because the heavy atom effect requires the use of rare and expensive metals, such as Ir or Pt,\textsuperscript{33, 34} this method of emission is also viewed as inefficient.

![Figure 6](image.png)

Figure 6. The HOMO and LUMO gaps seen in singlet and triplet state open shell systems

Thermally activated delayed fluorescence (TADF) is another emission technique that is growing in popularity. This technique involves using thermal energy to convert an electron in a triplet excited state to the singlet excited state, where it then emits in a fluorescence event.\textsuperscript{25, 26, 33, 35, 36} This process is termed reverse intersystem crossing (RISC) and it negates any spin forbidden processes.\textsuperscript{33}

### 1.2.1 Open-Shell Emitters

Most fluorescence and phosphorescence events are derived from “closed shell”
systems which contain two spin paired or unpaired electrons, respectively. In closed shell systems, there are two electrons in the highest occupied molecular orbital (HOMO). When the molecule is excited, one electron is promoted to the lowest unoccupied molecular orbital (LUMO), in the excited state while the other electron remains in the HOMO. Radical systems are considered “open shell,” as they contain a single, unpaired electron. Emission from open shell radical systems are like closed shell systems, with the exception that only doublet ground and excited states are possible (an electron is capable of having two spins, hence a “doublet”) rather than singlet and triplet excited state (Figure 7). When a radical is excited, its unpaired electron in its singly occupied molecular orbital (SOMO) is excited to the next singly unoccupied molecular orbital (SUMO). Transitions back to the SOMO are always spin allowed because there is no remaining electron to cause spin parity issues with the excited electron in the SUMO. Therefore, the theoretical maximum IQE of an open shell system is 100% because the presence of only doublet states removes conflict with competing electronic states.\textsuperscript{37, 38} Radical systems are not subject to loss of excitons through non-radiative relaxation because there are no energy states between the doublet ground and excited states.\textsuperscript{39}

![Figure 7. Fluorescence in closed shell systems vs open shell systems.](image.png)
The use of radicals as doublet emitters in OLEDs has garnered attention recently because the possibility of an IQE of 100%. This inherent phenomenon, coupled with the expected advancements in the device design of OLEDs, could lead OLEDs to be very competitive in the display market. Doublet emitters also have the ability to produce long emission wavelengths despite the lack of π conjugation.

This thesis research will focus on derivatizing the known emissive and air stable radical, tris-(2,4,6-trichlorophenyl) methyl or TTM. This compound is composed of a methyl radical stabilized by three chlorine substituted benzene rings. These bulky halogenated aryl groups act as protection for the radical causing the radical to be stable under ambient condition however it decomposes after photo-irradiation. To enhance the stability upon photolysis, and to increase the IQE, carbazole or pyridyl units have been installed into TTM radicals (Figure 8). These compounds have been incorporated into OLEDs that exhibit intense red emission and modest IQEs.

![Figure 8. Carbazole substituted TTM radical.](image)

The OLED device constructed from molecule C-1 produced the electroluminescent (EL) and photoluminescent (PL) spectra seen in Figure 9. The device emits in the red region of the visible light spectrum. The Li group was able to confirm
that the emission is due to a transition from the SUMO to SOMO due to the 660nm peak seen in the PL spectra. This transition was further confirmed via the absorbance spectra. A band centered at 600nm corresponds to an energy gap of 2.07eV; which is attributed to the transition between the SUMO and SOMO. 

![Figure 9](image)

Figure 9. The PL and EL of the carabzole-TTM moiety. 

1.3 Pnictogen Heteroles

Organic π-conjugated molecules have been extensively studied during the past 30 years; the interest in these compounds lies within their ability to display metal-like qualities such as conductivities similar to metals and semi-conducting properties. A diverse subsection of organic π-conjugated molecules is the heterole. The heterole is a molecular motif that has multiple structural advantages. When compared to cyclopentadienide the introduction of various heteroatoms promotes the production of materials with tunable electronic properties. Although the introduction of heteroatoms into π-conjugated systems has obvious benefits, there were a few challenges that initially slowed progress in heavier main group element (E, where E= P, S, As, Sb, Te, etc.) based π-conjugated systems. These challenges include the high reactivity of E–C bonds due to increased bond polarity and weaker orbital overlap when large or electropositive E atoms
are present. The typical heterole is composed of a butadiene fragment coupled to main group elements such as S, O, N, etc. This coupling involves interactions between the electron delocalization of the diene fragment and the lone pair of electrons on the heteroatom. The delocalization of electrons occurs when electrons in a \( \pi \)-conjugated molecule exchange across aligned p-orbitals of all neighboring atoms. When this occurs, the butadiene fragment and the heteroatom adopt a coplanar geometry. Additionally, the resonance of electrons causes the ground state of the molecule to lower, subsequently increasing the stability. As such, many of these heteroles are regarded as 6-\( \pi \) electron Hückel aromatic molecules. The electronic properties of the heterole is dependent on the influence of the heteroatom and adjacent substituents. Examples of commonly known heteroles include in order: cyclopentadienide, pyrrole, furan, thiophene, and phosphole (Figure 10). Due to their tunable electronic properties, heteroles display attractive optoelectronic applications in organic electronic devices like OLEDs, OFETs, and OPVs.

![Figure 10. Cyclopentatdienide pyrrole, phosphole, furan and thiophene, five common heteroles.](image)

In the realm of OLEDs, there is a niche of research that focuses on the use of carbazole as the \( \pi \)-conjugated heterole moiety. OLEDs are often composed of nitrogen rich units with \( \text{sp}^2 \) or \( \text{sp}^3 \) hybridized nitrogens; as such, carbazole and its derivatives have been used extensively in OLEDs due to high thermal stabilities, high triplet energies.
and efficient transportation of holes. The efficient transportation of holes is perhaps the most attractive feature of carbazole derivatives with respect to electro- and photoactive qualities. Many molecules used to produce blue or green emitting OLEDs contain carbazole derivatives which can be used as either an electron rich host or as an emissive layer. Carbazole is considered a donor molecule and is therefore usually coupled to an electron withdrawing group (EWG) like an aromatic linking group or a π-conjugated inhibitor. Despite carbazole’s huge success in the OLED industry, not much research has been extended to other pnictogen (group 15) elements, such as phosphorus, arsenic, antimony, or bismuth.

When comparing pnictogen heteroles, it is important to note that nitrogen containing heteroles (i.e. carbazole and pyrrole) are the most unique. Since nitrogen is a second-row element, the lone pair of electrons in its 2p-orbital can overlap well with the π electrons in the butadiene fragment which are also delocalized using 2p orbitals. This overlap is what allows these heteroles to obtain aromaticity and therefore stability. Because the lone pair of electrons in nitrogen are housed in a 2p orbital, the orbital contains only one angular node and no radial nodes. The lack of radial nodes promotes easy overlap and planarization of the nitrogen and butadiene π systems. The expansion of p-orbitals of the phosphorus (3p) and arsenic (4p) atoms results in the introduction of radial nodes. These radial nodes inherent to p-orbitals of phosphorus, arsenic (and other heavy pnictogens) limit the overlap of the heteroatom lone pair with the butadiene π system by introducing increased antibonding interactions. This lack of overlap promoted by the larger atoms causes heavier pnictogen heteroles to behave more like cyclic 1,3-butadienes and/or pnictogen trihydrides (phosphine, arsines, or stibines). This behavior
causes the heavier heteroles to be much less aromatic than pyrrole. The orbital differences between pyrrole, phosphole and arsole can be seen in Figure 11. A DFT study was performed to determine the orbital compositions (B3LYP, Stuttgart RLC ECP for all atoms). For the phosphorus and arsenic analogues, a very small amount of aromaticity is reestablished through a hyperconjugation interaction between the $\pi^*$ orbital of the pnictogen and the $\sigma^*$ orbital of a substituent attached to the pnictogen. To date, the most highly explored pnictogen containing pyrrole/carbazole derivative are a family of compounds known as phospholes. Phospholes are beginning to be explored in organic electronic applications because of their high electron accepting and electron transporting capabilities. Despite the significant lack of aromaticity compared to pyrrole or carbazole, the use of phospholes in organic materials has produced results comparable to the lighter counterpart.

Figure 11. The HOMO diagrams of the pnictogen heteroles further confirm the pyramidalization of the heavier pnictogens. As the lone pair on the heavier atoms becomes more diffuse (due to the addition of radial nodes), the main group atom becomes more pyramidalized. Pyrrole is the most planar due to its lack of radial nodes and arsole is the most pyramidalized because of its 2 radial nodes.
2. RESEARCH MOTIVATIONS

2.1 Research Motivations

Motivated by recent results which incorporate carbazole moieties into emissive TTM radicals, the goal of the proposed research is to explore the effects of the introduction of other pnictogen-containing groups into the TTM radical system. Specifically, it is the goal to determine how the structure of the pnictogen-containing group influences the photophysical properties of the TTM radical. Through this research, it is expected that the following questions can be answered: Can the wavelength and therefore color of the emission be tuned? Can the quantum yield of the emission be improved or adjusted? Can the stability of these molecules be improved?

Any novel information obtained from this research will prove to be useful in the OLED industry. The OLED industry is currently flourishing yet there is still much to be discovered. Due to the exceptional reputation of carbazole, exploration of other pnictogen atoms can prove to be very beneficial. The heavier pnictogen analogs are less likely to be researched because of increased synthetic difficulties. This research program will focus specifically on the incorporation of phosphorus-containing moieties into TTM derived radicals for potential applications as OLED materials.

2.2 Objectives

This thesis is comprised of the combination of several closely related projects, all of which involve the incorporation of pnictogen heteroles into the TTM radical system. Phosphorus derivatives were attempted first because phosphorus is the next heaviest pnictogen, and therefore, the chemistry of phosphorus should be the most similar to nitrogen in comparison to the other pnictogens. However, due to the addition of the radial
node in the 3p orbital of phosphorus atom, there are expected to be differences in their chemistry. The results can be expanded to the heavier pnictogens because each succeeding atom will contain an additional radial node that contributes to increased pyramidalization with decreasing delocalization. Any information obtained from the use of a phosphorus atom can be used to approximate information about heavier pnictogens.

Figure 12 shows the four dibenzophosphole compounds that were explored throughout the research. Compounds 1, 3 and 4 are known molecules common in literature; compound 2, however, has not been isolated. Diphenylphosphine derivatives 5-7 (Figure 12) were also incorporated into the research to serve as comparable data to the dibenzophospholes. The focus of the research will be placed mainly on molecules 2 and 5, as they are the compounds that will be implemented into the TTM radical. Compounds 1, 3 and 4 were synthesized as ways to get to compound 2 and compounds 6 and 7 were used throughout the experimental process simply for comparison reasons.

Figure 12. The dibenzophospholes and diphenylphosphines of interest

2.3 Spectroscopic and Analytical Tools Used for Characterization

The identity of the synthesized molecules was confirmed using heteronuclear
nuclear magnetic resonance (NMR) spectroscopy. Proton ($^1\text{H}$, referenced to residual solvent peak), phosphorus ($^{31}\text{P}$, referenced to external 85% $\text{H}_3\text{PO}_4$) and carbon ($^{13}\text{C}$, referenced to solvent peak) NMR were used extensively during the synthetic process because these tools yield information needed for direct structural analysis. $^{31}\text{P}$ NMR spectra were obtained as proton decoupled scans ($^{31}\text{P}{}^1\text{H}$) to prevent further complication of the spectral analysis resulting from phosphorus-proton spin coupling.

The identity of paramagnetic molecules was confirmed using electron paramagnetic resonance (EPR) spectroscopy which allowed for the determination of both the presence of unpaired electrons as well as for the elucidation of the electronic environment of the electron.
3. DIBENZOPHOSPHOLE ANALOGS AND REACTIONS

3.1 Introduction

The immense benefits of the incorporation of the nitrogen atom in carbazole into emissive organic molecules has prompted the exploration of the remaining elements in the pnictogen group. This chapter will discuss the incorporation of the dibenzophosphole moiety, the phosphorus analog of carbazole. A study performed by Chen and Huang et al., presented data on the optoelectronic properties of various substituted dibenzophosphole moieties (compounds and A-J in Figure 14).\textsuperscript{57} Included in their data are calculated HOMO and LUMO estimates (Table 1). The calculated HOMO and LUMO levels of the specified dibenzophospholes were compared to the HOMO and LUMO levels of two known carbazole units, 9-methyl-carbazole (MeN) and 9-phenyl-carbazole (PhN); both carbazole units are commonly used in optoelectronic devices.\textsuperscript{57} In comparison, the dibenzophospholes had lower HOMO and LUMO levels, but a wider range in energy gaps ($E_g$). The variation in gap levels is credited to the stronger electron donating ability of phosphorus compared to nitrogen.\textsuperscript{58} The reported triplet energies ($\ell E_g$) of these dibenzophospholes also revealed their potential as host material.\textsuperscript{12, 59}
Figure 13. Prominent dibenzophospholes with optoelectronic properties.

Table 1. The HOMO, LUMO, band gap energies ($E_g$) and triplet energies ($^3E_g$) of compounds 3, A-I as well as the MeN and PhN for comparison.  

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Chen and Huang et al. hinted at the benefits of electron withdrawing groups; a correlation between HOMO and LUMO levels showed that the incorporation of an electronegative group (in their case O, S, Se) increased the electron-transport ability of the dibenzophosphole.\textsuperscript{57, 60}

Inspired by the work of Chen and Huang, as well as the reports by Li and co-workers, the first goal of this thesis research was to incorporate a dibenzophosphole group into the TTM radical in place of carbazole. Although there is less aromaticity in the dibenzophosphole unit, it is still expected to change the optoelectronic properties of the resulting radical. The first attempt at installing the dibenzophosphole was pursued in the same fashion as the carbazole by the generation of the dibenzophospholide anion (2) (Scheme 1). The electron density on the phosphorus promotes the donation of electrons onto the para carbon of one of the three aromatic rings in the TTM system. The mechanism involves a nucleophilic aromatic substitution; a chloride atom from the para position is displaced readily allowing for the incorporation of nucleophiles.
3.2 Synthesizing dibenzophospholes

The chlorophosphole (1) was the first compound synthesized. The halo-
dibenzophosphole was chosen as the first target compound because halogen substituents
can be readily reduced to give phosphorus centered anions, making it easier to reach the
desired phospholide product. The synthesis used to make this product can be seen in
Scheme 2.
In its purest form the chlorophosphate (1-Cl) product was a pale-yellow solid with an apparent sand-like texture. After synthesizing the product, its identity was confirmed by NMR. The $^{31}$P NMR yielded a product peak at 68.15 ppm, which is consistent with the literature and falls in the expected chemical shift range for halogenated phosphorus compounds (35-200ppm).\(^{62}\)\(^{63}\) As the phosphorus atom becomes more electron deficient, the nucleus becomes deshielded and resonates downfield at a more positive chemical shift. The synthesis of 1 also results in the generation of a significant side product, namely the bromophosphate (1-Br), which exhibits an $^{31}$P NMR chemical shift of 49.21 ppm.\(^{57}\) The formation of this byproduct has been reported in the literature,\(^{61}\) and can be rationalized as the 2,2'-dibromobiphenyl starting reagent serves as a source of bromine during the synthesis of 1. These two different products form in a 3:2 chlorine to bromine ratio, yet their separation is not necessary. Both P-X (X = Cl or Br) bonds exhibit analogous reactivity; therefore, using the mixed product for further synthesis should not be detrimental. The relative purity of the product mixture was affirmed via $^1$H NMR. The compound contains 4 protons on each benzene ring making a total of 8 aromatic protons. Although the nuclear spins of phosphorus and hydrogen are the same ($I = \frac{1}{2}$), there is no significant $^1$H-$^{31}$P coupling seen in the $^1$H NMR. To prepare the phospholide anion, the mixture of compounds 1-Cl and 1-Br were treated with a potassium mirror in refluxing THF (Scheme 3). The P atom of phospholide anion is much more reactive compared to the P atom in the 1 due to its negative charge, and therefore should be more reactive in the desired nucleophilic aromatic substitution reaction with the TTM radical.
The $^1$H NMR of the phospholide anion was very similar to the spectra obtained for product 1-Cl, however there is a stark difference in the $^{31}$P NMR. The initial 68.15 ppm peak in 1-Cl was shifted upfield to -21.01 ppm. This large shift is due to the presence of the negative charge on the P atom causing it to be shielded by electron density. Although the dibenzophospholide has not been isolated and characterized in scientific literature, the $^{31}$P peak obtained correlates with the expected shift of a molecule with a phosphide anion (for example the phosphorus atom in diphenylphosphide resonates at -12.9 ppm).$^{61}$

We next focused on the synthesis of the $P$-phenyl-dibenzophosphole (3) (Scheme 4) which could also be used to generate the diphenylphospholide anion 2. The synthesis of this phosphole is reminiscent of the synthesis of product 1, the only difference being that dichlorophenylphosphine was used in place of PCl$_3$ to introduce the phosphorus moiety. 3 was identified as a sticky off-white solid, with a $^{31}$P NMR chemical shift at -10.35 ppm.$^{64}$ The influence of the substituent on the P atom is apparent when comparing the $^{31}$P NMR of 1 and 3, the less electronegative aryl group in 3 results in a less deshielded phosphorus center when compared to that in 1. The $^1$H NMR only exhibited peaks in the aromatic region; the same 8 protons from the product 1 are seen along with an additional 5 protons from the phenyl ring. The 5 protons from the phenyl group
produce peaks more upfield compared to the 8 protons in the dibenzo system because they are slightly shielded.

![Scheme 4. Synthesis of Phenyldibenzophosphole](image)

The dibenzophospholide (2) was also synthesized using 3. Instead of using potassium, the phenyl substituent was removed using lithium granules. Much like the dibenzophospholide synthesized from the 1, this lithium phospholide exhibited a $^{31}$P peak at -21 ppm. Given the negative charge on the P in the phospholide, it is reasonable to assume that the subsequent peak would occur upfield and in the negative region of the spectrum. As previously mentioned, the dibenzophospholide has never be isolated so the legitimacy of the product was further confirmed via comparison to potassium diphenylphosphide, which is a known compound that is structurally similar to the dibenzyl phospholide product. Potassium diphenylphosphide exhibits a $^{31}$P peak at -12.9 ppm, and therefore it can be assumed that the product synthesized with a $^{31}$P NMR signal at -21.29 ppm is indeed the expected phospholide anion. Although there is a large chemical shift difference between these two compounds, the phospholide anion was expected to produce a more upfield peak compared to the phosphide because the heterole confirmation promotes more electron donating from the benzene rings into the phospholide atom.
Finally, we also synthesized dibenzophosphole (4) was from 3 (Scheme 4) as an additional potential starting material for introduction into the TTM radical. The P-phenyldibenzophosphole was stirred over Li granules to make the phospholide, which was subsequently protonated using degassed water and acetic acid to give the dibenzophosphole product. \(^{45}\)

Scheme 4. Synthesis of dibenzophosphole

Much like the other phosphole products, the identity of compound 4 was confirmed through \(^{31}\)P and \(^1\)H NMR spectroscopies. The \(^1\)H NMR yielded the typical 8 aromatic protons found in the dibenzo system along with what seems like two singlets in the 5-6 ppm range. There is only one additional proton in the structure, so there should only be one additional singlet corresponding to the hydrogen attached the P atom. Interestingly, what looked like two singlets in the \(^1\)H NMR spectrum of 4 was actually one doublet with a large coupling constant of 200 Hz which was attributed to a one-bond \(^{31}\)P-\(^1\)H coupling. To confirm this assignment, the \(^1\)H coupled \(^{31}\)P spectrum also yielded a doublet with a \(^1J_{P-H} = 198\) Hz and confirmed the presence of a bond between the P atom and its H substituent. The coupling constants were measured using the following equation:

\[
J_{(Hz)} = \Delta_{ppm} \times \nu \quad (1)
\]
where $\Delta_{ppm}$ is the chemical shift difference between the peaks of interest and $\nu$ is the resonance frequency of the nucleus being interrogated.

### 3.3 Addition of Dibenzophospholide into the TTM system

The synthetic process involved in attaching the dibenzophospholide to TTM can be reviewed in Scheme 1. Theoretically, the phospholide should replace one of the chlorine atoms and assist in the stabilization of the radical. First, the TTM was obtained first through a Friedel-Crafts alkylation of trichlorobenzene using chloroform as the alkylation agent and aluminum trichloride as a Lewis acid. This reaction produces HTTM, which is the H substituted TTM molecule. The synthetic process can be seen in Scheme 5. The radical was then generated by first deprotonating the triarylmethyl C-H group using tetra-$n$-butyl ammonium hydroxide ([$n$-Bu$_4$N][OH]) followed by oxidation of the resulting triaryl anion using $p$-chloranil.

![Scheme 5. Synthesis of TTM radical](image)

The TTM radical is a bright red solid that fluoresces bright orange. Paramagnetic molecules are not NMR active because they reduce relaxation times which contribute to very broad peaks in the NMR. Consequently, the validity of the radical was confirmed via EPR. The spectrum yielded one radical peak, indicating that there were no coupling
interactions.

Once the TTM radical was synthesized and characterized, efforts were made to attach the dibenzophospholide anion to the system. Although the synthesis used was very similar to the synthesis involved in attaching carabazole to TTM, the installation of the dibenzophospholide moiety into the TTM radical was unsuccessful. All data obtained from these experiments indicated that once the phospholide was added, the TTM radical reverted into the protonated starting material, HTTM, which contains a closed shell, diamagnetic system rather than the desired radical.

As previously mentioned, the TTM radical is an air stable open shell system that produces deep red emissions. The paramagnetic radical is preferred due to the open shell system having theoretical maximum a quantum efficiency of 100%; however, due to the difficulty of adding the dibenzophospholide directly to the TTM radical, the same chemistry was attempted with the known TTM cation (TTM⁺). It was anticipated that TTM⁺ would be more reactive than the TTM radical toward nucleophilic aromatic substitution with the phospholide anion. The nucleophilic addition of phosphorus to TTM⁺ was also inspired by chemistry developed by Stephan and co-workers where dimesitylphosphine (Mes₂PH, Mes = 2,4,6-trimethylphenyl) was added to tris(pentafluorophenyl) borane (TPB) using nucleophilic aromatic substitution. Given that carbocations are isoelectronic and isolobal to boranes, it was assumed that similar chemistry could be carried out using dibenzophospholide and TTM⁺ (Scheme 6).
Scheme 6. Addition of Mes₂PH to TPB as reported by Stephan,⁶⁷ and attempted addition of dibenzophospholide to TTM⁺.

TTM⁺ was synthesized from the TTM radical by oxidation in chlorinated solvents using antimony pentachloride (SbCl₅) as an oxidant.⁶⁶ Using this synthetic methodology, the resulting cation is balanced with the counter anion SbCl₆⁻. The [TTM][SbCl₆] is a dark blue solid and its purity was confirmed through ¹H NMR before further use.

The addition of 2 to the TTM⁺ resulted in a sticky red solid with orange fluorescence. The ³¹P NMR yielded a peak at 41.40 ppm; peaks in this range indicate that the phosphorus atom in the molecule is attached to an electron withdrawing group. The phosphorus atom is bound through a phenyl ring, which are typically electron donating.
groups when unsubstituted, however in this case the phenyl ring can be viewed as electron withdrawing because of its electronegative chloro substituents and cationic nature. The \( ^{31}P \) NMR also produced another significant peak at 67.99 ppm; this peak is reminiscent of the chlorophosphole, which indicates that this byproduct may have been formed throughout the reaction. This by product is unfortunately the major component of the product mixture based on the integration of the peak (1.00) relative to the 41.40 ppm product peak (0.08).

3.4 Experimental

**Dichlorodibenzophosphole (1)** 2.49g (7.98 mmol) of dibromobiphenyl was dissolved 50 mL of anhydrous diethyl ether (Et\(_2\)O) and added to a 100 mL Schlenk flask. The solution was cooled then placed into an ice bath until it was cooled down to 0˚C. 10.0 mL of n-Butyllithium (1.6M, 16 mmol) was added dropwise to the mixture which caused the clear solution to adopt a pale-yellow color. The solution was left to stir at room temperature for approximately 1 hour before it was then cooled back down to -198˚C using liquid N\(_2\). Phosphorus trichloride (PCl\(_3\)) (5.0mL, 52.2 mmol) was added dropwise to the frozen mixture. Once all the PCl\(_3\) was added, the flask was left to warm up to room temperature while stirring. The solution was left to stir for 15 minutes, after which a white precipice began to form. Volatile materials were removed *in vacuo* and the resultant residue was left to dry under vacuum for 1 hour. The desired product was obtained after it was extracted via two toluene extractions. \(^1\)H NMR (400 MHz, CDCl\(_3\)) \( \delta \) 7.30 – 7.35 (m, 2H), 7.39 – 7.42 (m, 2H), 7.65 – 7.71 (m, 2H), 7.91-8.0 (d, 2H). \(^{31}\)P NMR (162 MHz, C\(_6\)D\(_6\)) \( \delta \) 49.21 (s), 68.15 (s).
Figure 14. Product 1 $^1$H NMR spectrum in CDCl$_3$.

Figure 15. Product 1 $^{31}$P NMR spectrum in CDCl$_3$.

**Potassium dibenzophospholide ([K][2])** In a nitrogen-filled glove box, 0.1369g (3.47mmol) of potassium metal was placed inside a 50 mL Schlenk flask. The Schlenk
flask was removed from the glove box and heated with heat gun to melt the K until it formed a mirror around the sides of the flask. The flask was taken back into the glove box and 0.640g (2.91mmol) of 1 dissolved in 20 mL of anhydrous THF was added dropwise into the Schlenk flask. The solution was refluxed at 70°C for 16 hours. The solution was then cooled and filtered to remove unreacted K metal. The solvent was removed in vacuo. The product was a pale yellow solid. $^1$H NMR (400 MHz, C$_6$D$_6$) $\delta$ 7.42 – 7.33 (m, 1H), 7.07 – 6.99 (m, 1H), 6.99 – 6.93 (m, 1H), 6.83 (dd, $J = 8.8, 7.5, 2.7, 1.6$ Hz, 1H). $^{31}$P NMR (162 MHz, C$_6$D$_6$) $\delta$ -21.93 (s).

Figure 16. Product [K][2] $^1$H NMR spectrum in C$_6$D$_6$. 
Figure 17. Product [K][2] $^{31}$P NMR spectrum in C$_6$D$_6$.

**P-phenyldibenzophosphole (3)** Approximately 0.60g of dibromobiphenyl (2 mmol) was dissolved in 10 mL of anhydrous tetrahydrofuran (THF) in a 25 mL Schlenk flask. The solution was allowed to stir for 10 minutes before it was cooled to -78°C using an acetone/dry ice bath. 1.52 ml (4 mmol, 2.57 M) of n-butyllithium was added dropwise to the cooled mixture. The solution was allowed to stir for 30 minutes. 0.34g (2 mmol) of dichlorophenyl phosphine (PhPCl$_2$) was dissolved in 1 mL of THF was then added by syringe into the di-lithiated biphenyl solution. The yellow mixture warmed up to room temperature while it was left to stir overnight. The next day the volatile solvents were removed *in vacuo* and a honey colored oil was left over. The product was extracted from this oily material with dichloromethane (DCM) and then filtered over a plug of celite and silica gel. The entire solution was concentrated under vacuum and the product was removed from an oily solution via recrystallization using degassed methanol. $^1$H NMR
(400 MHz, CDCl$_3$) $\delta$ 7.96 (d, $J = 7.9$ Hz, 1H), 7.76 – 7.63 (m, 1H), 7.47 (td, $J = 7.6$, 1.2 Hz, 1H), 7.33 (ddd, $J = 11.2$, 6.4, 1.6 Hz, 2H), 7.25 – 7.20 (m, 1H). $^{31}$P NMR (162 MHz, CDCl$_3$) $\delta$ -10.94 (s).

Figure 18. Product 3 $^1$H NMR spectrum in CDCl$_3$. 
Dibenzophosphole (4) 0.305g of 3 (1.17 mmol) was combined with 0.0081g of lithium (Li) metal in a 25 mL Schlenk flask. 10mL of THF was added to the compounds and the resultant solution was left to stir for 4.5 hours. After stirring for the allotted time, the solution was canula transferred into another Schlenk flask to separate it from unreacted Li metal. An aliquot of a degassed H₂O (1.5mL)/AcOH (0.4mL) solution was added to the reaction mixture. The entire solution was then placed under vacuum to remove volatile solvents. The product was separated from unwanted products by extracting with DCM and subsequent air-free filtration over a plug of dry alumina. The resultant white powder was further washed with anhydrous pentane. ¹H NMR (400 MHz, C6D6) δ 7.73 – 7.67 (m, 2H), 7.22 – 7.18 (m, 1H), 7.07 – 6.99 (m, 3H), 6.93 – 6.89 (m, 2H), 5.17 (d, J = 197.4 Hz, 1H). ³¹P NMR (162 MHz, CDCl₃) δ -67.98 (s).
Figure 20. Product 4 $^1$H NMR spectrum in CDCl$_3$. (Product contained excess starting material)
HTTM 1.0g (5.5 mmol) of trichlorobenzene (TCB) was placed into a 50 mL pressure flask inside a nitrogen-filled glovebox. 0.073g (0.612 mmol) of chloroform (CHCl₃) was added into the flask followed by 0.074g of anhydrous aluminium trichloride (AlCl₃). The mixture was stirred and heated to 85°C for 2 hours. The solution was then cooled to 25°C and 20 mL of 0.1N HCl was added into the solution in an ice bath. The resulting yellow solution was extracted with 50mL of CHCl₃. The organic layer was then dried over magnesium sulfate for an hour and then filtered. The subsequent solution was concentrated using a rotary evaporator. The product was identified as an off-white solid that was further washed with hexane to produce a white solid. ¹H NMR (400 MHz, C₆D₆) δ 6.99 – 6.97 (m, 1H), 6.93 – 6.91 (m, 1H), 6.76 (s, 1H).
TTM In nitrogen filled glove box, 3.0g (5.41 mmol) of HTTM was placed into a Schlenk flask previously wrapped in aluminum foil. 4.60 mL of n-butylammonium hydroxide (4.56g, 7.03 mmol) in 50 mL of THF was combined with the HTTM contained in the Schlenk flask. Upon combination the solution turned a deep red color. The solution was left to stir for 4 hours, after which 1.86g (7.57 mmol) of p-chloranil in 25 mL of THF was syringe transferred into the solution. The mixture was left to stir overnight, and the resultant purple solution was concentrated in vacuo. The dark purple solid was mixed with 1.0g of silica gel in 10 mL of DCM. The solid was again concentrated under vacuum. The product was purified via a packed silica gel column and 100% hexanes the eluent.
TTM‘SbCl₆’ (6) In a nitrogen filled glove box, a 50mL Schlenk flask covered with aluminum foil was charged with 0.5024g (0.907mmol) of TTM radical and 0.5510g (1.841mmol) of SbCl₅. The Schlenk flask was removed from the glove box and 20mL of CCl₄ was syringe transferred into the flask. The solution was left to stir over night, after which solvent was removed in vacuo. The solid obtained was dark blue. ¹H NMR (400 MHz, CDCl₃) δ 7.37 – 7.35 (m, 3H), 7.24 – 7.22 (m, 3H).
Figure 24. Product 6 $^1$H NMR spectrum in CDCl$_3$. The peaks at 7.40 ppm and 7.27 ppm represent impurities.

**TTM-P(8).** In a nitrogen filled glovebox, 0.2235 g (0.253 mmol) of product 6 was placed in a 25 mL Schlenk flask covered with aluminum foil. 10 mL of anhydrous THF was also added into the flask. 0.0513 g (0.225 mmol) of 2 was dissolved in 3 mL of THF and then subsequently added dropwise into the flask. The initially dark blue color of the solution turned bright red upon addition. The solution was allowed to stir overnight. The THF was removed *in vacuo* and the resultant product was a sticky red substance with orange fluorescence. $^1$H NMR (400 MHz, C$_6$D$_6$) δ 7.54 (d, J = 6.9 Hz, 1H), 7.32 (d, J = 7.6 Hz, 1H), 7.05 (t, J = 7.1 Hz, 1H), 6.93 (t, J = 7.3 Hz, 1H). $^{31}$P NMR (162 MHz, C$_6$D$_6$) δ 41.40 (s).
Figure 25. Product 8 $^1$H NMR spectrum in C$_6$D$_6$.

Figure 26. Product 8 $^{31}$P NMR spectrum in C$_6$D$_6$. 
4. DIPHENYLPHOSPHINE ANALOGS AND REACTIONS

4.1 Introduction

Diphenylphosphines are known compounds, all of which contain a phosphorus atom attached to two phenyl rings. These compounds can sometimes be used as precursors for ligands in catalysis.\textsuperscript{69} Although these compounds are not heteroles, they are close in structure to the dibenzophospholes described in the previous chapter to yield similar compounds with comparable properties.

4.2 Reactions with Diphenylphosphine

Much like the dibenzophosphole, the reactions with TTM radical were unsuccessful. The installation of diphenylphosphine into the TTM radical system resulted in a sticky red substance; however, the $^1$H NMR spectrum proved that the TTM radical was reduced back into the HTTM molecule throughout the reaction. The $^{31}$P NMR spectrum of this reaction yielded a peak at -40.02 ppm, which is consistent with unreacted diphenylphosphine (-40.7 ppm)\textsuperscript{70}; indicating that the phosphine did not react with the TTM radical. The mishap was also observed when potassium diphenylphosphide (5) was reacted with the TTM radical. The $^1$H NMR spectrum was indicative of only HTTM and the $^{31}$P was denoted evidence of unreacted potassium diphenylphosphide. The TTM$^+$ reactions did prove to be fruitful; and it was found that potassium diphenylphosphide could be successfully added to TTM$^+$. The $^{31}$P NMR of potassium diphenylphosphide reveals a signal for the phosphorus anion at -12.9 ppm, and any variation in the product $^{31}$P NMR would indicate a change electron shielding around the phosphorus atom.\textsuperscript{61} The $^{31}$P NMR chemical shift for the synthesized compound appeared at 41.98 ppm, which is nearly identical to what was observed from the reaction of TTM$^+$ and
dibenzophospholide compound (41.40ppm). Given this information, it can be assumed that the diphenylphosphide anion is also attached to one of the TTM rings with chlorine substituents that act as an electron withdrawing group causing the peak to show up more downfield.

4.3 Experimental

**KPPH₂-TTM⁺(10)** In a nitrogen filled glovebox, 0.1001 (0.113mmol) of potassium diphenylphosphide (5) was placed in a 25mL Schlenk flask covered with aluminum foil. 9 mL of anhydrous THF was also added into the flask. 0.0322g (0.143mmol) of 6 was dissolved in 5mL of THF and then subsequently added dropwise into the flask. The initially dark blue color of the solution turned bright red upon addition. The solution was
stirred overnight. The THF was removed in vacuo and the resultant product was a sticky red substance with orange fluorescence. $^1$H NMR (400 MHz, C$_6$D$_6$) δ 7.77 (ddd, J = 14.2, 8.2, 1.4 Hz, 3H), 7.55 – 7.47 (m, 3H), 7.02 – 6.88 (m, 10H). $^{31}$P NMR (162 MHz, C$_6$D$_6$) δ 41.98 (s).

Figure 27. Product 10 $^1$H NMR spectrum in C$_6$D$_6$. 
Figure 28. Product \textbf{10} $^{31}$P NMR spectrum in C$_6$D$_6$. 
5. CONCLUSION

Throughout the experimental process of this research, the ultimate goal was to attach a dibenzophosphole moiety to the TTM radical, a molecule with beneficial luminescence due to its open shell system. Four different dibenzophospholes were synthesized, all with the final goal of converting them to the anion so that they could subsequently be added to TTM. Difficulties arose when trying to add the phospholide anion to the TTM radical, however there was success when the TTM$^+$ was used in place of the radical. The experimental process has proven that the only successful attempts at attaching a dibenzophospholide with TTM involved using an electron-rich phosphorus anion which acts as the Lewis base and the cationic TTM that behaves as the Lewis acid. The two compounds interact in a FLP fashion to form a deep red product with an orange luminescence.

Although there was success, the goal of the research was not fulfilled because the product produced did not contain a radical anywhere throughout the molecule. The cation TTM is not a paramagnetic molecule, so it falls into the subsection of closed shell systems. This means that the fluorescence emitted from the molecule is caused by transitions between singlet states rather than doublet states. As mentioned before, this fluorescence method is not as efficient as fluorescence from doublet states. Achieving the radical from the cationic TTM can be achieved via a reduction event. This method will be explored during the next phase of experiments.

Exploration of the larger pnictogens will also be attempted. Phosphorus was expected to be a pnictogen that behaved most similarly to nitrogen in the TTM system; however, these assumptions were proven to be inconclusive given the data obtained.
According to the literature, carbazole based units can be incorporated into the TTM radical system with ease; dibenzophosphole based units did not reflect the same chemistry. The difference between the characteristics of the two is most likely attributed to the introduction of a radial node, making the lone pair on the phosphorus atom more diffuse than the lone pair on a nitrogen atom. The increase in nucleophilicity of phosphorus compared to nitrogen proved to only be beneficial in TTM cation reactions. Delving into experiments with larger pnictogens (Ar, Sb, etc) will provide a larger basis for assumptions about how pnictogen heteroles can be used in organoelectronics.
REFERENCES


68. Stephan, D. W., Frustrated Lewis Pairs. *Journal of the American Chemical Society* 2015, **137** (32), 10018-10032.


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