Synthesis of j-aggregating dibenz[a,j]anthracene-based macrocycles

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## Synthesis of J-Aggregating Dibenz[a,j]anthracene-Based Macrocycles

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                            | Kooi, Steven; MIT, ISN  
                            | Bulovic, V.; MIT, Electrical Engineering and Computer Science  
                            | Swager, Timothy; Mass. Inst. of Tech., Chemistry |

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Synthesis of J-Aggregating
Dibenz[\(a,j\)]anthracene-Based Macrocycles

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ABSTRACT. Several fluorescent macrocycles based on 1,3-butadiyne-bridged dibenz[\(a,j\)]anthracene sub-units have been synthesized via a multistep route. The synthetic strategy involved the initial construction of a functionalized dibenz[\(a,j\)]anthracene building block, subsequent installation of free alkyne groups on one side of the polycyclic aromatic framework, and a final cyclization based on a modified Glaser coupling under high-dilution conditions. Photophysical studies on three conjugated macrocycles revealed the formation of J-aggregates in thin films as well as in concentrated solid solutions (polyisobutylene matrix) with peak absorption and emission wavelength in the range of \(\lambda = 460 \text{ nm to } \lambda = 480 \text{ nm}\). The characteristic red-shifting of the J-aggregate features as compared to the monomer spectra, enhancement in absorption intensities, narrowed linewidths, and minimal Stokes shift values, were all observed. We demonstrate that improvements in spectral features can be brought about by annealing the films under a solvent-saturated atmosphere, where for the best films the luminescence quantum efficiency as high as 92% was measured. This class of macrocycles represents a new category
of J-aggregates that due to their high peak oscillator strength and high luminescence efficiency have the potential to be utilized in a variety of optoelectronic devices.

INTRODUCTION. Shape-persistent macrocycles have received much attention in the field of materials science, particularly in the area of nanoscale architectures.\(^1\) The first macrocycle featuring two unfunctionalized anthracenes linked by 1,3-butadiyne bridges was reported in 1960, but due to the lack of modern synthetic and characterization methods, the nature of the resulting material was not rigorously elucidated.\(^2\) Following little interest in such systems over the next four decades, reports of anthrylene-ethynylene oligomers and macrocycles have surfaced in the past 5 years.\(^3\) However, the molecular rigidity and lack of solubilizing groups resulted in the reported compounds having poor solubilities in common solvents. To create a class of molecules that could have potentially interesting photophysical and materials properties, we embarked on the design of conjugated macrocycles based on rigid dibenz[\(a, j\)]anthracene units bridged by butadiyne \(\pi\)-linkers. This was a logical choice since aryleneethynylene and 1,3-butadiyne linkages are frequently used in conjugated systems (e.g. polymers) for their ability to maintain rigidity and \(\pi\)-conjugation.\(^4\) The polycyclic aromatic motifs are commonly seen in other areas of materials science, notably in the fields of discotic liquid crystals and graphitic materials.\(^5\) By employing various modern synthetic transformations, it was possible to introduce numerous functionalities (e.g. sidechains) into the structure to give better solubility and processability. In particular, bulky 4-alkoxyphenyl substituents located near the middle of the macrocycles serve several purposes: 1) as synthetic handles to allow for the facile electrophilic cyclizations\(^6\) used to establish the dibenz[\(a, j\)]anthracene framework, 2) as solubilizing groups, and most importantly, 3) as a source of steric hindrance to bring about twisting of the \(\pi\)-system. Such distortion of the rigid framework by steric bulk has been known to induce slipped stacking arrangements,\(^7\) resulting in aggregate structures with unique optical properties. Similar slipped structures are also known in nature: for example, the arrangement of J-aggregated chlorophyll chromophores is crucial to the light-harvesting efficiency of photosynthetic systems.\(^8\) Using natural photosystems as a guide and inspiration, researchers have found...
ways to emulate this J-aggregate design in various porphyrins and perylene bisimides. More recently, the laboratories of Frank Würthner have also successfully implemented the rational synthesis of several J-aggregated systems using supramolecular design principles. Ever since their serendipitous discovery in 1936, J-aggregates have been of great theoretical interest because they display coherent, cooperative phenomena like superradiance and giant oscillator strength, a consequence of their electronic excitation being delocalized over several molecules. Besides being theoretical curiosities, J-aggregates also have a myriad of practical applications, such as their use as organic photoconductors, photopolymerization initiators, and nonlinear optical devices, as well as the emerging applications such as the recently demonstrated critically coupled resonators and strongly QED coupled microcavity LEDs.

Herein, we report the synthesis and characterization of a series of J-aggregating macrocycles based on functionalized dibenz[a,j]anthracene fragments linked at the 6- and 8- positions by a pair of 1,3-butadiyne bridges, in which the ring interior can be viewed as an octadehydro[18]annulene system. The results of their photophysical studies are also detailed.

RESULTS AND DISCUSSION.

Synthesis. Macrocycles 6a and 6b were prepared in six steps from the previously reported dibromide 1 (Scheme 1). Subjecting the dibromide to a double Suzuki coupling with 4-alkylphenylboronic acids afforded terphenyl derivatives 2, which were then converted to the required 6,8-diiododibenz[a,j]anthracenes via a double iodonium-induced electrophilic cyclization. Numerous attempts to convert the diiodide to the bis-acetylene 5 via Sonogashira and Castro-Stephens reactions proved unsuccessful, instead resulting in complex, undefined mixtures. However, an indirect method involving a lithiation/carbonylation sequence to give 4, followed by Corey-Fuchs homologation, successfully afforded dialkyne 5. Owing to the sterically encumbered environment of the reaction centers, dialdehyde 4 was always accompanied by the formation of monoaldehyde byproduct 13. Separation of the two could however, be easily achieved by column chromatography. Finally, an
oxidative coupling utilizing conditions previously developed\(^\text{20}\) in our group was performed, furnishing macrocycles \(6\text{a}\) and \(6\text{b}\) in reasonable yields.

**SCHEME 1.** Synthesis of macrocycles \(6\text{a}\) and \(6\text{b}\).

The synthesis of macrocycle \(12\) (Scheme 2) involved a similar sequence of transformations employed in the preparation of \(6\text{a}\) and \(6\text{b}\), with the exception that the bis-alkoxyterphenyl \(7\) could only be converted to the desired diiodide \(9\) in two steps, via a skeletal rearrangement of the structurally intriguing \(8\), using modifications of known reactions.\(^\text{6, 21}\) A second alkoxy-based macrocycle bearing branched farnesol-derived sidechains was also synthesized in a manner analogous to \(12\), with its
existence confirmed by MALDI-TOF. Unfortunately, this fourth and final macrocycle could not be satisfactorily separated from a trimeric byproduct even after repeated column chromatography and attempted fractional recrystallizations. In addition to the three macrocycles, compound 15 (the acyclic analog of 6a) was also prepared to study the effect of the number of bridges on the photophysical properties. This was made in three steps (Scheme 3) starting from monoaldehyde 13, which is a byproduct isolated during the purification of dialdehyde 4a.

The abovementioned target compounds were characterized by $^1$H NMR, $^{13}$C NMR, high-resolution mass spectrometry (MALDI-TOF), UV/Vis and fluorescence spectroscopy. In the $^1$H NMR spectra of the macrocycles, the two protons located within the ring were found to be shifted downfield ($\delta \approx 9.5$ ppm) as a result of Van der Waals deshielding brought about by steric interactions. The lack of any upfield shift of those internal protons implies the absence of a ring current$^{22}$ in these systems (i.e. no diatropic effect observed). Brief polarized optical microscopy experiments were also performed on the macrocycles in hope of finding liquid crystalline behavior as well, but the compounds had extremely high melting points (between 200°C and 330°C) and were also observed to decompose and discolor at those elevated temperatures.

**SCHEME 3.** Synthesis of acyclic 15.

![Scheme 3](image.png)

**Photophysical studies.** A SPEX fluorolog, with dual monochromators, was used to collect photoluminescence (PL) and photoluminescence excitation (PLE) spectra. The instrument is wavelength and intensity calibrated and it compensates for variations in excitation intensity by monitoring the incident optical power level. In PL measurements, the 6a films were optically excited at a wavelength $\lambda = 375$ nm. For PLE spectra, emission at $\lambda = 508$ nm was collected. Figure 1 shows the UV-Vis absorption and fluorescence spectra of the four compounds in chloroform. Macrocycles 6a and
6b displayed essentially identical spectral profiles, with absorption and emission maxima occurring at around 440 nm and 455 nm respectively. Changing the peripheral alkyl groups to alkoxy chains (e.g. 12) resulted in a slight bathochromic shift, with the spectral shape remaining similar otherwise. The spectra of the acyclic 15 differed somewhat from the macrocycles, which was expected due to the major structural difference. Its absorption spectrum was blue-shifted relative to the others, possibly due to reduced conjugation resulting from the absence of the second diyne linker. A much larger Stokes shift was also observed, which could indicate reduced rigidity, once again as a result of having only a single linker. Fluorescence quantum yields of the compounds were measured against quinine sulfate in 0.1N H₂SO₄ (Table 1). The three macrocycles in chloroform solution showed fairly high quantum yields between 0.40 and 0.50, whereas the singly-bridged 15 had a lower value of 0.35.

**Table 1.** Photophysical properties of 6a, 6b, 12, and 15.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Absorption max (nm)</th>
<th>Emission max (nm)</th>
<th>Quantum yield, Φ_F</th>
<th>Extinction coefficient (M⁻¹ cm⁻¹)</th>
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</thead>
<tbody>
<tr>
<td>6a</td>
<td>443</td>
<td>456</td>
<td>0.45</td>
<td>90141 (at 443 nm)</td>
</tr>
<tr>
<td>6b</td>
<td>443</td>
<td>456</td>
<td>0.43</td>
<td>63569 (at 443 nm)</td>
</tr>
<tr>
<td>12</td>
<td>448</td>
<td>461</td>
<td>0.47</td>
<td>79113 (at 448 nm)</td>
</tr>
<tr>
<td>15</td>
<td>395</td>
<td>440</td>
<td>0.35</td>
<td>38206 (at 395 nm)</td>
</tr>
</tbody>
</table>
FIGURE 1. Normalized absorbance (solid lines) and emission (dotted lines) spectra of 6a, 6b, 12 and 15 in chloroform.

To test for the presence of J-aggregates, we investigated the thin film photophysics of the macrocycles. As 6a was synthesized in the largest quantity, films of this compound were studied in greatest detail. The initial films were produced by spin-coating a fairly concentrated (5 mg/mL) toluene solution of 6a on to glass or quartz cover-slips (18 x 18 mm). Fortuitously, the first few films showed promising UV-Vis absorption features consistent with J-aggregates (Figure 2).

![Absorption Spectra](image)

FIGURE 2. Absorption spectra of 6a, solution (blue line) vs. film (red line), normalized to the absorbance at 340 nm.

Compared with the solution spectrum, the 6a film spectrum shows an aggregate absorption peak at 467 nm (red-shifted by (23±1) nm from the solution). Even more notable is the high intensity and narrow linewidth of this peak (J-band), which dominates all other spectral features. This is in stark contrast to the solution spectrum, in which the peak at 443 nm shows much lower intensity than those between 300 nm and 360 nm (absorptions due to pendant p-alkoxyphenyl moieties). Normalizing the solution and film absorbances at 340 nm, the enhancement in the peak intensity (at 467 nm) relative to the other spectral features becomes evident (Figure 2). The bathochromic shift and the strong intensity
of the aggregate peak, are photophysical characteristics of J-aggregates.\textsuperscript{23} From the emission spectra of the 6a films we find the Stokes shift to be only 4 nm (Figure 3), versus 13 nm in solution phase. Such minimal Stokes shift is also consistent with the existence of J-aggregates.\textsuperscript{24} It is notable that the fluorescence band is a mirror image of the low energy edge of the J-band absorption.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{spectra.png}
\caption{Normalized absorption (blue line) and fluorescence (pink line) spectra of 6a (film).}
\end{figure}

We also find that it is possible to spin-coat films of 6a that did not display a J-band. These less strongly absorbing films could be produced when: the spin rate (of the spin-coating process) was high (e.g. 4000 rpm), a lower concentration (< 2 mg/mL) of 6a in solvent was used, or when a more volatile solvent (e.g. THF) was employed. The use of these parameters provided for less-than-favorable conditions for aggregate formation. But when these ‘non-aggregated’ (i.e. monomeric) films were then subjected to conditions conducive to aggregate formation, the typical J-aggregate spectral features were found to emerge with time. This was achieved by vapor-annealing the films in a solvent chamber saturated with toluene vapor for 45 minutes, and then retrieving them for spectral (UV-Vis) re-acquisition. It can be seen (Figure 4) that the vapor-annealing, which should result in more ordered thin films\textsuperscript{25}, precipitates the appearance of the highly intense J-band, confirming that molecular organization
was indeed important in producing the desired J-aggregate photophysics. When the volatile THF is used as the spin-casting solvent (particularly with low 6a concentration) the resulting films lacked J-aggregate features. However, J-aggregate features can be recovered when these films are placed in a solvent chamber containing THF vapor (Figure 4).

**FIGURE 4.** Absorption spectra of 6a (film), before and after annealing under toluene (PhMe) and THF vapor, normalized to the absorbance at 340 nm.

Similar photophysical experiments were also performed on films of the two longer-chained macrocyclic analogs 6b and 12. In both cases, the films are cast from THF solutions of the macrocycles, and subsequently annealed under THF vapor for 45 minutes. UV-Vis data are acquired before and after the annealing process, and the spectra of 6b and 12 are shown in Figure 5. The spectra of the pre-annealed films did not show J-bands, but these appeared in both cases upon annealing. Therefore, the results obtained with 6b and 12 were analogous to those of 6a, suggesting that the doubling of chain length of the peripheral alkyl/alkoxy groups had little effect on the photophysics, be it in solution or in the film-state. Similar J-aggregate photophysics could not be observed with the non-cyclic 15, implying that the aggregate formation may require approximate molecular planarity (steric hindrance in the non-cyclic 15 produces a larger deviation from planarity, since the two dibenzanthracene sub-units are less
It is likely that J-aggregation of these polycyclic aromatics in the solid-state relies on π-π stacking interactions that could be disrupted if the non-planarity became too pronounced.

**FIGURE 5.** Absorption spectra of 6b (film) and 12 (film), before and after annealing under THF vapor, normalized to the absorbance at 340 nm.

Additional experiments examining the photophysics of 6a as a function of concentration were also undertaken. A series of films were spin-coated using 6a solutions (polyisobutylene matrix/chlorobenzene as solvent) of varying concentrations, and their photoluminescence spectra, excitation spectra, and fluorescence lifetimes were measured. Chlorobenzene was chosen as it provided for optimal co-miscibility of 6a, polyisobutylene, and solvent. When a film containing a very low concentration of 6a (i.e. 0.0005 mg in a 1 mL solution of 40 mg/mL polyisobutylene (PIB) in chlorobenzene) was used, its emission peak was at 455 nm (Figure 6), identical to that observed in solution spectra. As the amount of 6a used in the spin-coating process was increased to 0.002 mg/mL of PIB/chlorobenzene, aggregate peaks began to emerge at 470 nm with a shoulder at 500 nm, while the 455 nm “monomer” peak diminished. A further increase in 6a concentration to 0.005 mg/mL resulted in further reduction in the 455 nm peak, so that at 0.02 mg/mL the monomer peak can no longer be observed, at which point the emission spectrum begins to resemble those obtained with neat films discussed above (pure 6a, no PIB...
From the excitation spectra (Figure 7), no J-band at 470 nm could be observed at the lowest concentration of 0.0005 mg/mL, but as the concentration was increased ten-fold, a peak at 470 nm appeared, becoming more pronounced with increasing 6a concentrations. Fluorescence lifetimes (at 470 nm) for a series of solutions and films of different concentrations were also measured. Solution lifetimes were found to be between (1.7 ± 0.1) ns regardless of concentration. In the film state, it can be observed from Table 2 that the lifetimes generally decrease as the concentration of 6a was increased from 0.0005 to 2.000 mg/mL (only a small incidence of scatter is observed in the trends). In particular, with a concentration of 0.0005 mg/mL, a lifetime of $\tau_m = 1.3$ ns was obtained, comparable to what was observed in chlorobenzene solutions, while at higher concentrations, lifetimes of about $\tau_J = 0.24$ ns are observed. Higher doping concentration also leads to a measurable increase in thin film photoluminescence (PL) quantum yield (QY) from $\Phi_m = 43 \pm 6$ % for the monomeric film to $\Phi_J = 92 \pm 8$ % for the aggregate films. To determine PL QY, we compared the PL counts from the 6a film to a thin film standard of known QY, accounting for relative differences in absorption strength of the films. The standard was a thin film of thickness 75 nm of the small molecule tris(8-hydroxyquinolinato)aluminum (Alq$_3$). The Alq$_3$ film was prepared by thermally evaporating recrystallized Alq$_3$ in ultra high vacuum (growth pressure below $10^{-6}$ Torr) onto a quartz substrate that was carefully solvent cleaned and oxygen plasma treated to remove trace impurities. The published QY for Alq$_3$ in thin film is $32 \pm 2\%$. We erred on the side of caution and used a value of QY = 30% for our calculations. To make a fair comparison of QY, for each film in consideration, the percentage of absorbed optical excitation was determined from optical transmission measurements. Measured PL counts were then normalized to the percent absorption values, on a film by film basis. For 6a in monomeric form, we calculated the PL QY to be $43 \pm 6 \%$, which is similar to the QY for 6a in solution, and for 6a at high doping concentrations, we determined the QY to be $92 \pm 8 \%$.

The emergence of a red-shifted narrower linewidth optical transition at higher 6a concentrations, the corresponding reduction in lifetime, and increase in quantum yield of aggregates as compared to
monomers are indicative of J-aggregate formation. In J-aggregates, strong coupling between the monomer transition dipoles produces a new cooperative molecular state. The coupling results in a new optical transition called the J-band, when the interaction strength exceeds the monomeric dephasing processes. The interaction between monomeric transition dipoles lowers the overall energy of the cooperative state; hence the J-band absorption/fluorescence is red-shifted relative to that of the monomer. In the J-aggregate state, multiple molecules coherently couple, the number being denoted by $N_c$, and the J-aggregate exciton delocalizes over all of them. Coherent coupling amongst the $N_c$ molecules leads to the acceleration of the radiative rate of the J-band states by a factor of $N_c$ relative to the monomer, which translates into shorter excited state lifetime and higher PL QY. The radiative rate enhancement is typically referred to as a superradiance phenomenon since it is caused by coherent exciton coupling, though in J-aggregates the mechanism for the coupling is near-field Coulombic interactions while in classic superradiant systems, the origin is interference effects in the spontaneous light emission process. Since the radiative rate of a J-aggregate increases relative to that of the monomer by a factor of $N_c$, from a comparison of lifetimes ($\tau_j$ vs. $\tau_m$) and quantum yields ($\Phi_j$ vs. $\Phi_m$), $N_c$ can be determined using the equation:

$$N_c = \frac{\tau_m \Phi_j}{\tau_j \Phi_m}$$

The data obtained suggest that $N_c$ is on the order of 12 for our 6a J-aggregate films. Coherent coupling also leads to a narrower total linewidth for the J-aggregate optical transition relative to the monomer, because the delocalized exciton averages out site-to-site variations, and suppresses the inhomogeneous broadening. The linewidths of the monomer optical transition and the J-band are dominated by inhomogeneous broadening. Nevertheless, the width of the J-band relative to the monomer spectrum does characterize the coherence of the system. The linewidth of the J-band is narrower than the monomer optical transition because in the J-aggregate state, the exciton is delocalized over the $N_c$ molecules that are coherently coupled, which tends to average out site-to-site inhomogeneities in the exciton energy. This motional narrowing is manifest in the smaller linewidth for J-aggregate absorption.
and emission spectra. This coherent coupling also results in the accelerated radiative process in the J-aggregate state, which translates into the higher observed QY and shorter exciton lifetime for the J-aggregate compared to the monomer.

**FIGURE 6.** Photoluminescence intensity vs. concentration of 6a in thin films (PL scaled by subtracting the background and scaling by integrated intensity at all wavelengths).
FIGURE 7. Excitation vs. concentration of 6a in thin films (PLE scaled by subtracting background and scaling by integrated intensity at all wavelengths).

Table 2. Fluorescence lifetimes of 6a (solutions and films) at different concentrations.

<table>
<thead>
<tr>
<th>Concentration (mg/mL)</th>
<th>Lifetimes (bimodal) (ns)</th>
<th>State</th>
</tr>
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<tr>
<td>0.002</td>
<td>1.7 (100%)</td>
<td>Solution (PhCl)</td>
</tr>
<tr>
<td>0.02</td>
<td>1.7 (100%)</td>
<td>Solution (PhCl)</td>
</tr>
<tr>
<td>0.2</td>
<td>1.8 (100%)</td>
<td>Solution (PhCl)</td>
</tr>
<tr>
<td>0.0005</td>
<td>1.3 (99.9%) 4.8 (0.1%)</td>
<td>Film</td>
</tr>
<tr>
<td>0.002</td>
<td>0.2 (96.7%) 1.3 (3.3%)</td>
<td>Film</td>
</tr>
<tr>
<td>0.005</td>
<td>0.3 (96.6%) 1.2 (3.4%)</td>
<td>Film</td>
</tr>
<tr>
<td>0.01</td>
<td>0.3 (97.4%) 1.0 (2.6%)</td>
<td>Film</td>
</tr>
<tr>
<td>0.02</td>
<td>0.2 (98.1%) 1.1 (1.9%)</td>
<td>Film</td>
</tr>
<tr>
<td>0.05</td>
<td>0.6 (84.5%) 1.5 (15.5%)</td>
<td>Film</td>
</tr>
<tr>
<td>0.10</td>
<td>0.4 (82.7%) 1.4 (17.3%)</td>
<td>Film</td>
</tr>
<tr>
<td>0.20</td>
<td>0.4 (94.9%) 1.4 (5.1%)</td>
<td>Film</td>
</tr>
<tr>
<td>2.00</td>
<td>0.2 (97.4%) 0.7 (2.6%)</td>
<td>Film</td>
</tr>
</tbody>
</table>

Molecular modelling. In order to better visualize the equilibrium geometry of the macrocycles 6a, 6b, and 12, molecular calculations were performed at the semi-empirical PM3 level, using a model
compound (Figure 8) with deliberately shortened alkyl sidechains to enable more rapid completion of the calculation. As can be seen in Figure 8, the macrocycle is composed of two 1,3-butadiyne-linked planar dibenz[\(a,j\)]anthracene sub-units that are slightly staggered relative to each other as a result of steric crowding in the middle of the molecule. Despite this structural distortion, the core of the macrocycle retains some overall planarity, which would still allow for intermolecular \(\pi-\pi\) stacking interactions. By comparison, the acyclic analog shows greater non-planarity (Figure 8g), since the two non-restrained aromatic sub-units have more freedom to minimize steric repulsions. As a result, \(\pi-\pi\) stacking interactions in acyclic 15 may be weakened.
FIGURE 8. PM3-calculated models (a) top-down view of geometry-optimized macroryclic structure, (b) molecular electrostatic potential map, (c) optimized structure tilted to emphasize steric crowding, (d) edge-on view of optimized structure, (e) frontier HOMO, and (f) frontier LUMO, (g) edge-on view of the acyclic model structure, (h) top-down view of the acyclic structure.

CONCLUSION. In summary, three dibenz[a,j]anthracene-based macrocycles have been synthesized and spectroscopically characterized. The conjugated macrocycles display pronounced photophysical properties in the solid-state, such as the intense red-shifted absorbances, narrow linewidths, and small Stokes shifts, indicating J-aggregate formation. These new compounds may have the potential to be utilized in various optoelectronic devices (e.g. lasers, photovoltaics, and polaritonic devices\textsuperscript{16a,36}).

ACKNOWLEDGMENT. This work was supported by the National Science Foundation and the Army Research Office’s IED Stand-Off Detection Research Program (W911NF-07-1-0654), and the U.S. Army through the Institute for Soldier Nanotechnologies (DAAD-19-02-0002).

SUPPORTING INFORMATION AVAILABLE: Full experimental details pertaining to the synthesis of all new compounds described herein. This information is available free of charge via the Internet at http://pubs.acs.org.

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