Cyclic molecules can exhibit fundamentally different electronic dynamics compared with their linear counterparts due to the lack of chain end effects, high rotational symmetry, and the presence of ring strain.1,2 The photophysical properties of cyclic molecules have been extensively investigated for a large number of systems ranging from small aromatic rings3,4 to large molecular complexes.5 A recurrent key concept is exciton delocalization, which is known to lead to strong fluorescence suppression in cyclic molecular structures such as benzene.6 The high symmetry results in a dipole-forbidden lowest exciton (\(k = 0\)) state; however, emission is weakly enabled by Herzberg–Teller intensity borrowing from allowed higher energy transitions through the involvement of vibrations with suitable symmetry.8 Electronic delocalization also plays a major role in the efficient energy transfer between adjacent antennae in natural light-harvesting complex 2 (LH2) based on chlorophyll units,9,10 where the pigments are held loosely by protein scaffolds in circular structures.11 Such designs of nature have inspired many attempts to create synthetic nanoring analogues, for instance, cycloparaphenylene12,13 cyclic oligothiophenes14, nanorings synthesized from phenylcarbazole units linked with phenylene-ethynylene-butadiyne groups,15 and an extensive range of porphyrin arrays.16–19

While investigating electronic properties on structures with such high symmetry, an interesting question inevitably arises, namely, how symmetry distortions influence the photophysical properties. When defects are introduced into cyclic molecular structures, the lowest dipole-forbidden exciton state may be mixed with higher lying states and some absorption into the lowest state becomes possible.20,21 As the distortion becomes more and more prominent, deviations from high rotational symmetry dominate and the fluorescence is no longer suppressed.20 Symmetry breaking also plays an important role in structures found in natural systems, such as LH1 complexes containing a reaction center (RC) at their center, which can be divided into at least two distinct classes.9 The first class is monomeric, consisting of only one RC surrounded by one cyclic LH1 complex, as found in Rhodospirillum rubrum and Rhodospirillum palustris.22 However, while LH1 in Rsp. rubrum is believed to exhibit a highly symmetric ring-like structure,22 LH1 in Rsp. palustris displays a gap, providing a portal through which ubiquinone can transfer electrons across the boundaries of LH1, which is vital for photosynthetic processes.23,24 The second class of systems is dimeric, consisting of two circular RC-LH1 complexes as found in Rhodobacter sphaeroides.25 Such examples illustrate the urgent need for an understanding of how symmetry lowering in ring-like chromophore assemblies affects the electronic dipole transitions that control the absorption of light.

Previous studies have examined both experimentally and theoretically the effects of conformational and dynamic disorder on cyclic systems;26,27,29,30 however, investigations into the process of symmetry breaking based on ring-like compounds with geometrical defects require an excellent control over their structures. Fully \(\pi\)-conjugated porphyrin nanorings bound to wheel-like templates32,33 provide an ideal model system here: These nanorings have acquired a high level of rigidity through the butadiyne bridges and the template inside the rings, their initial excited states are fully delocalized over the entire ring.1

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and symmetry defects of varying extent can be introduced through modifications in synthesis. 

In this study we examine five synthetic porphyrin nanorings with structures mimicking those of the natural RC-LH1 complexes previously described. We investigate fully conjugated nanorings with and without defects, a nanoring featuring a gap due to broken conjugation, and systems consisting of two fused nanorings (see Figure 1). We reveal the effect of symmetry distortions on the electronic transitions in these systems, showing that as deviations from rotational symmetry increase, oscillator strength is transferred to the lowest \((k = 0)\) state. Concomitantly, the degeneracy of the dipole-allowed first excited \((k = \pm 1)\) state is shown to be lifted, leading to an ultrafast polarization switching effect in the emission from strongly symmetry-broken nanorings.

The synthesis and characterization of the nanoring complexes is described in detail in the Supporting Information and in previous literature reports.\(^{33-36}\) Figure 1 shows the structures and simplified graphical representations of the five investigated nanorings consisting of zinc porphyrin units joined by butadiyne bridges. The six-legged \((T6)\) templates inside all complexes bind to the porphyrin units, thus maintaining a rigid ring-like shape for the nanorings, which is vital because it suppresses competing symmetry weakening from ring distortions.

The most highly symmetric compound, \(c-P6\cdot T6\), has a closed ring structure with six porphyrin units,\(^{37}\) comparable to the \(D_{6h}\) symmetry of benzene. For ring-like systems with 6-fold symmetry, such as benzene, the energy-level structure can easily be visualized through the simple Hückel molecular orbital method for \(\pi\)-conjugated molecules. This approach postulates two nondegenerate \((k = 0, k = 3)\) and two pairwise degenerate \((k = \pm 1, k = \pm 2)\) exciton states; however, simple analysis of the transition dipole moments to these states reveals that only the two degenerate \(k = \pm 1\) states of orthogonal polarization can carry significant oscillator strength. In particular, the transitions to the lowest \((k = 0)\) state can be visualized by the simple Kasha model\(^{38}\) as dipole contributions summing to zero in a hexagonal arrangement. Therefore, once the molecule has relaxed into its lowest state, emission is expected to be strongly suppressed, as we have shown to be the case for \(c-P6\cdot T6\)\(^{2,50}\).

Figure 1 also displays four other compounds whose symmetry has been reduced with respect to \(c-P6\cdot T6\). Here \(c-P6\cdot T6\) has a very similar structure, but with the minor modification that two opposite porphyrins have additional alkyne bridges with TIPS (triisopropylsilyl) end groups attached (see SI), leading to reduced symmetry. Stronger deviations are introduced by \(l-P6\cdot T6\), which comprises a linear six-porphyrin chain bound to the \(T6\) template and therefore features a “gap” in the \(\pi\)-conjugated backbone. The two spiro fused ring structures \(s-P11\cdot (T6)\)\(_2\) and \(s_r\cdot P11\cdot (T6)\)\(_2\) comprise 11 porphyrin units each and consist of two interlinked six-porphyrin nanorings with perpendicular template planes, as shown in Figure 1, with the only difference being the ethyl side chains attached to the shared porphyrin unit in the center of \(s_r\cdot P11\cdot (T6)\)\(_2\) (see SI).\(^{36}\) For all measurements reported, \(l-P6\cdot T6\) was dissolved in toluene, while all other samples were prepared in toluene/1% pyridine solutions to avoid aggregation.

We proceed to investigate the impact of symmetry on the electronic properties of ring-like molecules by investigating the changes in absorption and emission of these five porphyrin nanoring systems, ranging from almost perfect 6-fold symmetry, through modifications in structure, to ring structures with broken \(\pi\)-conjugation. As a first comprehensive insight into the energy level structure in these ring systems, Figure 2 shows the extinction coefficient spectra of the five compounds in the region of the \(Q\)-band. Extinction coefficient spectra over a wider range of photon energies and emission spectra can be found in the SI. In contrast with zinc porphyrin monomers that display degenerate \(S_0-S_1\) transitions at \(\sim 550\) nm \((Q\)-band\),\(^{39}\)
the Q\textsubscript{x}-band of the ring structures is split into Q\textsubscript{y} and Q\textsubscript{z} bands with polarizations parallel and perpendicular to the molecular backbone, respectively, and the Q\textsubscript{x} band is significantly red-shifted to \sim 800 nm because of the increased conjugation length.\textsuperscript{28}

For the most highly symmetric molecule, c-P6\textsubscript{T6}, the extinction coefficient spectrum shows three distinctive strong peaks with approximately equal energy difference. We have previously shown that excitations in this system are fully delocalized over the entire ring.\textsuperscript{2,37} Therefore, the simple Hückel model should hold as previously described, and only the lowest allowed \( k = \pm 1 \) exciton state carries significant oscillator strength. As we have previously discussed,\textsuperscript{37} the multiple peaks most likely arise from vibrational contributions, with the approximate 75 meV energy spacing between the three main peaks tentatively supporting an interpretation in terms of Franck–Condon vibronic progression within an adiabatic Born–Oppenheimer framework. As the symmetry is lowered for the other nanorings systems, two effects can be clearly observed in the extinction coefficient spectra. First, new features appear at the low-energy end (1.2 to 1.4 eV) of the spectra, which grow in intensity (red arrows in Figure 2) as symmetry distortions become more severe. We propose that such deviations from 6-fold symmetry may lead to oscillator strength being transferred to the lowest \( k = 0 \) exciton state. For the ring with broken \( \pi \)-conjugation (I-P6\textsubscript{T6}) this low-energy peak rises to considerable strength (Figure 2e), suggesting a strongly allowed transition to the \( k = 0 \) state becomes possible. This phenomenon has analogies to observations in the natural RC-LH1 complexes from \textit{Rps. palustris}, which features a gap in the LH1 ring\textsuperscript{24} that enhances absorption contributions for the \( k = 0 \) state. Concomitant with these changes, the distortions from 6-fold symmetry appear to also affect the peak intensity ratio of the three strong peaks visible in all extinction spectra. Compared with c-P6\textsubscript{T6}, the first peak in the series, as indicated with green arrows in Figure 2, gradually rises with respect to the second peak as symmetry constraints are lifted. This effect may derive from higher energy vibronic progressions of the emerging \( k = 0 \) state overlapping with the \( k = 1 \) features and is therefore most likely of similar origin.

To ascertain whether a particular lowering of the nanoring symmetry indeed leads to the emergence of a dipole-allowed lowest \( (k = 0) \) state, we further probed the emissive properties of these molecules, following excitation into the Q\textsubscript{y}-band. While for a rotationally symmetric molecule, transitions from the \( k = 0 \) exciton state are dipole forbidden within the Franck–Condon approximation, weak emission is possible via Herzberg–Teller intensity borrowing through joint electronic-vibrational transitions that match the symmetry of the higher-lying allowed \( (k = 1) \) transition.\textsuperscript{37} A weakening of symmetry constraints, however, is expected to relieve the strict selection rules on the \( k = 0 \) state, leading to additional Franck–Condon emission.

To investigate such effects, we therefore measured the radiative emission rate of all molecules, which reflects the oscillator strength in the \( k = 0 \) state into which the system rapidly relaxes following excitation. For this purpose, the fluorescence lifetime was measured using time-correlated single-photon counting (TCSPC) and the quantum yield was obtained by comparing the fluorescence spectrum integral against that of a reference standard (full details of measurement are provided in SI). From these independent measurements, the radiative \( (\Gamma_{\text{r}}) \) and nonradiative \( (\Gamma_{\text{nr}}) \) rate contributions were evaluated, as listed in Table 1. It can be seen that the

![Extinction coefficient spectrum](image)

**Figure 2.** Extinction coefficient spectrum in the region of the Q\textsubscript{y}-band for (a) c-P6\textsubscript{T6}, (b) c-P6\textsubscript{y}T6, (c) s\textsubscript{Et}P11(T6)\textsubscript{y}, (d) s-P11(T6)\textsubscript{y} in toluene/1% pyridine, and (e) I-P6\textsubscript{T6} in toluene. Arrows indicate the trend in peak intensity ratios for each compound in comparison with c-P6\textsubscript{T6}.

<table>
<thead>
<tr>
<th>molecule</th>
<th>( \tau ) (ps)</th>
<th>( \Phi_{\text{F}} )</th>
<th>( \Gamma_{\text{r}} ) (ns\textsuperscript{-1})</th>
<th>( \Gamma_{\text{nr}} ) (ns\textsuperscript{-1})</th>
</tr>
</thead>
<tbody>
<tr>
<td>c-P6\textsubscript{T6}</td>
<td>627</td>
<td>0.7%</td>
<td>0.010</td>
<td>1.60</td>
</tr>
<tr>
<td>c-P6\textsubscript{y}T6</td>
<td>682</td>
<td>1.3%</td>
<td>0.019</td>
<td>1.45</td>
</tr>
<tr>
<td>s\textsubscript{Et}P11(T6)\textsubscript{y}</td>
<td>271</td>
<td>0.5%</td>
<td>0.020</td>
<td>3.73</td>
</tr>
<tr>
<td>s-P11(T6)\textsubscript{y}</td>
<td>343</td>
<td>0.8%</td>
<td>0.024</td>
<td>2.90</td>
</tr>
<tr>
<td>I-P6\textsubscript{T6}</td>
<td>364</td>
<td>2.3%</td>
<td>0.064</td>
<td>2.69</td>
</tr>
</tbody>
</table>

*Details of measurements are given in the Supporting Information.*

Increasingly severe deviations from ring-like symmetry introduced along the group of molecules lead to a gradual increase in their radiative emission rates. The strongest enhancements are seen for I-P6\textsubscript{T6}, for which \( \Gamma_{\text{r}} \) increases more than 6-fold with respect to c-P6\textsubscript{T6} as the \( \pi \)-conjugation is broken, resulting in efficient emission that suggests strong Franck–Condon allowed transitions. The spiro-fused rings show weaker increases, with the radiative rate more than doubling in comparison with the 6-fold symmetric c-P6\textsubscript{T6}. The weakest effects are seen for c-P6\textsubscript{y}T6, which may be partly because a \( C_2 \)-type symmetry reduction alone is not expected to lead to oscillator strength being transferred to the \( k = 0 \) state.\textsuperscript{3,40}

In contrast and as expected, no correlation between nonradiative decay rates and molecular symmetry is observed. The fluorescence lifetime of s\textsubscript{Et}P11(T6)\textsubscript{y} is particularly short, which is caused by stronger nonradiative decay channels \( (\Gamma_{\text{nr}}) \) Table 1) possibly introduced by the presence of the ethyl side chains attached to the shared porphyrin unit.
We proceed by establishing a quantitative correlation between such reduced-symmetry enhancements in radiative rates and the changes in the absorption features previously discussed. Figure 3 shows the correlation between (a) the absorption intensity of the $k = 0$ state and (b) the intensity ratio of the first two main peaks, plotted against the radiative emission rate for each of the five compounds. To capture the contributions to the absorption into the emerging $k = 0$ state (indicated by red arrows in Figure 2), we integrated the extinction coefficient spectrum over the low-energy wing up to a full width at half-maximum below the first main peak and given in relation to the integral over the full spectrum of the Qs band. (b) Intensity ratio of the first two peaks in the main ($k = \pm 1$) series of the extinction coefficient spectrum. Each data point corresponds to a ring complex with the colors depicted matching that of the icons above the graphs representing the structure. Dashed lines are guides to the eye, highlighting the strong correlation between the parameters, but do not imply a particular functional dependence.

<table>
<thead>
<tr>
<th>Radiative rate [ns$^{-1}$]</th>
<th>Mean peak ratio</th>
<th>k=0 abs.integral</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.02</td>
<td>0.1</td>
<td>0.3</td>
</tr>
<tr>
<td>0.04</td>
<td>0.2</td>
<td>0.1</td>
</tr>
<tr>
<td>0.06</td>
<td>0.01</td>
<td>0.0</td>
</tr>
</tbody>
</table>

Figure 3. Analysis of the extinction coefficient spectra for the nanoring complexes c-P6-T6 (red), c-P6-T6 (blue), s-P11-T6 (yellow), s-P11-T6 (magenta), and l-P6-T6 (green) in relation to their radiative emission rates. (a) Spectrally integrated extinction coefficient of the emerging low-energy ($k = 0$) peak, determined by integration of the spectra shown in Figure 2 up to a full width at half-maximum below the first main peak and given in relation to the integral over the full spectrum of the Qs band. (b) Intensity ratio of the first two peaks in the main ($k = \pm 1$) series of the extinction coefficient spectrum. Each data point corresponds to a ring complex with the colors depicted matching that of the icons above the graphs representing the structure. Dashed lines are guides to the eye, highlighting the strong correlation between the parameters, but do not imply a particular functional dependence.

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lowered as a result of the clear break in conjugation. Figure 4b illustrates the mechanism that explains our experimental observations. Highly rotationally symmetric systems such as c-P6-T6 absorb light through the degenerate $k = \pm 1$ states that are orthogonally polarized. Following an ultrafast relaxation to the $k = 0$ state, Franck–Condon forbidden emission from this state takes place via weak Herzberg–Teller coupling. Because there is no preferred excitation polarization direction nor a net emission dipole moment in the nanoring plane, the polarization memory is completely lost in this 2D plane, resulting in $\gamma = 0.1$; however, for I-P6-T6 the lowered symmetry lifts the degeneracy of the $k = \pm 1$ states and results in the creation of new higher-lying states (labeled as $k = 1, 2...$ in Figure 4b) with transition dipoles of well-defined orientation with respect to the symmetry plane. These changes are accompanied by a strong recovery of Franck–Condon allowed transitions from the lowest excited state, as previously discussed and examined in natural light-harvesting complexes. As indicated in Figure 4b, this scenario creates a well-defined relation between the orientation of the dipole moment associated with the lowest ($k = 0$) state and those of the nondegenerate higher-lying states, with some combinations being parallel and others orthogonal.

In particular, the conjugation break in I-P6-T6 causes the lowest ($k = 0$) state wave function to have a lobe (probability density maximum) at the ring side opposite to the break, leading to a net dipole moment in the $x$ direction (here we take the $y$ direction to lie within the mirror-symmetry plane). Such deviation from a fully delocalized (i.e., no nodes) wave function in the $k = 0$ state of the highly symmetric c-P6-T6 thus transfers oscillator strength to the ground state of I-P6-T6, in accordance with the observed recovery in radiative rate. The higher lying $k = 1$ state of I-P6-T6 has some analogy to the degenerate $k = 1$ state of the symmetric c-P6-T6 as its nodes fall onto the symmetry plane and it is hence less perturbed by the conjugation gap than the orthogonal $k = \pm 1$ state. Therefore, the observed polarization switch in I-P6-T6 can be understood in terms of an initial excitation into nondegenerate higher-lying states, such as the $k = 1$ state with polarization along the $y$ direction, followed by relaxation into the $k = 0$ state that is associated with a dipole moment in the orthogonal $x$ direction. Strictly speaking, orthogonal absorbing and emitting transition dipole moments should yield $\gamma = \pm 0.2$, whereas collinear dipole moments result in $\gamma = 0.4$ for randomly oriented molecules in solution; however, contributions from higher lying ($k \geq 1$) states and their vibrational progressions are broadly overlapping (clearly separate features are not discernible in the absorption of I-P6-T6; see Figure 2). Hence, in reality, anisotropy values between $0.1$ and $0.2$ are observed for the range over which the excitation wavelength is varied (Figure 4a); however, the strong observed variation of polarization anisotropy from negative to positive values as the excitation energy is tuned through the higher-lying excitation band provides clear evidence that severe symmetry lowering in such systems leads to a lifting of the degeneracy of the ring-like $k = \pm 1$ states, creating a series of states with well-defined transition dipole moments.

In conclusion, $\pi$-conjugated six-porphyrin nanorings with various distortions to the cyclic structures have been investigated to explore the influence of symmetry lowering on their photophysical properties. We have demonstrated that as symmetry distortions increase in severity along the series of structures, oscillator strength is transferred to the transition into the lowest ($k = 0$) state, which is Franck–Condon forbidden in the fully 6-fold rotationally symmetric rings. The associated increases in oscillator strength were shown to correlate with gradually emerging $k = 0$ absorption features at the lower energy end of the absorption spectra and changes in intensity ratio of higher-lying overlapping absorption peaks. Thus, symmetry plays a vital role in influencing and correlating absorptive and emissive properties of ring-like molecules, which we have been able to capture quantitatively. Furthermore, we have revealed through time-resolved photoluminescence anisotropy probes that strong symmetry breaking, induced, for example, through full breaks in conjugation, lifts the degeneracy of the $k = \pm 1$ states found in highly symmetric ring assemblies, instead creating a series of states with well-defined transition dipole moments. We note that these findings have direct relevance to ring-like complexes found in natural light-harvesting systems, which have a rich diversity of geometrical structures. Here chromophores can be arranged in circular, elliptically deformed or array arrangements, exhibit a gap, or form dimeric structures; however, comparable investigations of the impact of structural symmetries on photophysical properties are challenging in such biological systems, as the chromophores are held loosely together and interact significantly with their environment. Our study therefore highlights the outstanding potential of using synthetic biomimetic porphyrin nanorings as model systems to investigate the effect of symmetry and geometric arrangement on the fundamental nature of the light-harvesting states.
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■ ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpcllett.5b02617.

Synthetic procedures and characterization of all compounds. Full extinction coefficient and emission spectra of all compounds. Experimental details of up-conversion technique, TCSPC, and quantum yield measurement. PL transients and anisotropy dynamics. (PDF)

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■ REFERENCES


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