Fluorescence Lifetimes and Emission Patterns Probe the 3D Orientation of the Emitting Chromophore in a Multichromophoric System

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The optoelectronic properties of dyes embedded in thin polymer films are currently under intense study due to the use of such materials in nanotechnology1 and nano-electronics.2 The fluorescence lifetime of a single fluorescent molecule (SM) embedded in a thin film can fluctuate as a direct manifestation of the SM nanoenvironment (e.g., as a result of polymer dynamics).3–5 Furthermore, it has been shown that the fluorescence lifetime strongly depends on the position and orientation of the SM with respect to the boundaries of the film, an effect known as the influence of the electromagnetic boundary conditions (EBC).6–13 This effect is especially important in very thin polymer films (<30 nm). Therefore, several groups have devoted efforts to determine the 3D orientation of a SM in thin polymer films.14–17 Techniques such as annular illumination (probing the orientation of the absorption transition dipole)15,16 and wide-field defocusing (probing the emission transition dipole moment) have been proposed and exploited.17

In this Communication, we report on the fluorescence behavior of a single first generation multichromophoric dendrimer with four perylene imide chromophores at the rim (G1R4) embedded in a thin polymer film (Chart 1).18,19 A model developed to describe energy transfer in this system13 invokes energy hopping between the different chromophores. It was suggested that, at any moment in time, the emission of a single dendrimer originates from the chromophore that has the lowest energy (fluorescent trapping site). Since the emission patterns give direct evidence for the hopping model between the tetrahedral oriented chromophores in individual G1R4 molecules, the EBC effect for a SM of G1R4 should result in discrete changes in the fluorescence lifetime. By using a scanning confocal optical microscope with polarization-sensitive FIFO mode detection, the intensity, the degree of polarization P, and the fluorescence lifetime of a G1R4 SM were simultaneously determined.6

Figure 2 shows the intensity recorded in the two orthogonal polarization channels (a), the degree of polarization (b), and the fluorescence lifetime (c) trajectories of a single G1R4. Each emissive level clearly corresponds to a different polarization value. The four different emissive levels are thus related to four different emitting chromophores. The lifetime trajectory shows an increase of the fluorescence lifetime from 4.8 ns for the highest emissive level to approximately 7 ns for the other three lower emissive levels. Such an anticorrelated behavior between the lifetime and the intensity can be attributed to the EBC effect. Indeed, the emission of a chromophore oriented more or less in the plane of the sample becomes feasible. Böhmer et al.17 provided exact wave-optical calculations of these defocused images. Upon comparing the experimental data with calculated results, we obtained the emission dipole orientation.

The patterns shown in Figure 1 were observed sequentially during the indicated period. One sees that the emission pattern of a single G1R4 molecule changes as function of time. The patterns provide direct evidence that different chromophores of the G1R4 SM emit in time, as different dipole orientations are observed successively. In this series we observe only three different patterns, probably one chromophore bleached during alignment. On the basis of the simulated patterns (Figure 1b),6,21 following the sequence shown in Figure 1a, we attribute the first and second patterns to a chromophore with an emission dipole oriented in-plane with angles of 40° and 70°, respectively, with respect to the x-axis (θ = 90°, φ = 40° and θ = 90°, φ = 70°). The third pattern indicates an out-of-plane oriented chromophore (θ = 0°, φ = 0°). Note that the error for the out-of-plane orientation θ is ±15°, whereas a very high accuracy in the xy plane can be obtained.21

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An important parameter related to the hopping model lifetimes obtained for parallel and perpendicular oriented chromophores in a 25 nm film is from 4 to 12 ns, in good agreement with the calculated lifetime fluctuations. The largest observed lifetime fluctuation in a single molecule was recorded in G1R4. A further lifetime fluctuation study, clearly confirming this assumption. Both imaging and lifetime measurements confirm hopping to a fluorescent trap in G1R4.

For the investigated molecules (158), 40% show EBC-type lifetimes. The absence of jumps in the simultaneously recorded emission spectra ensures furthermore that the lifetime jumps are not connected to other processes in the matrix. As mentioned earlier, the structure of a G1R4 SM in a polymer film resembles a tetrahedron. The four chromophores of the system can thus be placed in such a way that three of them are similarly oriented more perpendicular, relative to the interface, while in comparison the fourth chromophore is oriented more parallel. Figure 2d shows a simulation of the lifetime change of G1R4 for such an orientation, clearly confirming this assumption. Both imaging and lifetime measurements confirm hopping to a fluorescent trap in G1R4.

For the investigated molecules (158), 40% show EBC-type lifetimes. The largest observed lifetime fluctuation in a 10 nm film is from 4 to 12 ns, in good agreement with the calculated lifetimes obtained for parallel and perpendicular oriented chromophores. An important parameter related to the hopping model presented for G1R4 is the relative inhomogeneous character of the environment acting on the SM. In such a situation, each chromophore feels a slightly different environment. In contrast, for a more homogeneous local environment, the energies of the four chromophores are nearly identical. In the latter case, each individual chromophore has now a probability to emit. Due to the finite integration time, the detected signal then results in an averaging of the fluorescence of the different chromophores. An example is given in Figure 3. For the series of emission patterns depicted in Figure 3a, the 3D orientations of three patterns could be assigned on the basis of simulated patterns (Figure 3b) as θ = 90°, φ = 0° for the first pattern, θ = 90°, φ = 45° for the second pattern, and θ = 90°, φ = 135° for the fourth pattern.

A movie of the emission patterns series shown in part a is available. The switching in time between distinct patterns related to chromophores acting as fluorescent traps and the average pattern (even at 0.5 s time resolution) indicates that the homogeneity of the polymer surrounding this particular molecule fluctuates as a result of polymer dynamics. A more detailed study of the influence of local polymer inhomogeneity will be published elsewhere.

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Supporting Information Available: Details on the setups and techniques used, spectra of the SM depicted in Figure 2, and calculations. This material is available free of charge via the Internet at http://pubs.acs.org.

References

(6) See Supporting Information.