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Single terrylenediimide molecules diluted in a 20-nm-thick polyvinylbutyral polymer film were localized and observed by scanning confocal fluorescence microscopy. A modular and compact confocal microscope and the high optical stability of the molecules allowed a repeated imaging and observation over >5 h at room temperature. Most of the molecules showed several “on–off–on” transitions (blinking) on a time scale from seconds to hours, before permanent bleaching occurred. We determined that >1.5 × 10^7 fluorescence photons are emitted from the most-stable molecules before the final bleaching step occurs. Despite the “on–off–on” transitions, however, the overall change in fluorescence intensity, either integrated over each image of a time series or summed for several individual molecules, resembled an exponential-like decay, familiar from measurements of many-molecule ensembles. We also observed the polarization of the fluorescence from single molecules during excitation with circular polarized light. From these measurements, possible rotations of the molecular dipoles were studied. Over a span of 5 h, the polarization angle in most cases did not change by >15°. This may explain the slow and small intensity changes but excludes molecular rotation as a reason for the blinking behavior.

Introduction

The microscopic observation of fluorescence from single chromophores adsorbed on surfaces or in thin films or membranes and in fluids at room temperature and low temperatures gives direct insight into several processes that are not accessible in ensemble measurements, where fluorescence is detected from many molecules simultaneously. For example, bleaching of a single molecule can appear in one step after several fluorescence cycles, the number of which may vary statistically between different molecules, whereas a fluorescence bleaching curve of a molecule ensemble decays continuously and nearly exponentially in time. Variations from a single-exponential bleaching curve may be related to the random orientation within the ensemble or to inhomogeneities in the individual local environment of each molecule. The investigation of single-molecule fluorescence is expected to uncover such inhomogeneities on a molecular scale. Apart from permanent bleaching, new effects have already been observed on the single-molecule level, such as large, photoinduced spectral shifts and, possibly related to this, a sudden reappearance of fluorescence some time after an instantaneous drop of the fluorescence intensity. One fascinating application of microscopic single-molecule fluorescence detection is the observation and localization of chemical reactions on the single-molecule level.

To obtain insights that are statistically relevant, one must collect data from many molecules. Conventional scanning and nonscanning microscopy, combined with detectors with high photon counting efficiencies such as cooled avalanche photodiodes and cooled charge-coupled device (CCD) cameras, are simple and reliable techniques for these investigations. Scanning near-field optical microscopy (SNOM), first used for imaging single dye molecules on surfaces, offers resolution superior to that of conventional far-field microscopy. Furthermore, because of the smaller excitation volume, SNOM should, in principle, decrease the background level against which the single molecule fluorescence occurs. However, at present production of coated optical-fiber-tip probes with apertures suited for single-molecule detection is difficult, making SNOM more cumbersome to use in this case than conventional microscopes. With a combined confocal and near-field optical microscope, we compared the confocal mode with the near-field mode. The resolution in the confocal mode is only about a factor of five worse than that of the SNOM mode of 50–150 nm but offers other important advantages: (1) better suppression (by 10-fold) of the excitation light because of the epi-illumination scheme;
(2) the possibility of higher excitation intensities than with the fiber tips; (3) three-dimensional imaging capability; and (4) ease of use. Therefore, for investigations of samples with diluted molecules, we found the confocal mode to be much better suited than SNOM.

Here we report on fluorescence bleaching experiments with terylene-dimide (TDI) molecules\(^{18}\) embedded in a polyvinylbutyral (PVB) polymer film. These dye molecules, excited by a continuous HeNe laser line at 633 nm, allow quantitative analysis of single-molecule fluorescence because of their high photostability and high fluorescence quantum yield. The fluorescence intensity of single molecules measured in the confocal microscope depends on the molecular orientation and, for individual molecules, differs between zero and some maximum value. We compared the influence of the orientation of the molecules’ dipole moments onto the local intensities in the images with the images obtained by computer simulations. Measurements of the fluorescence polarization were performed to investigate whether molecular rotation took place, which could be a reason for slow intensity changes or blinking.

Experimental Section

The sample is scanned line by line in the focus of an on-axis confocal microscope by using an 80-\(\mu\)m-range three-dimensional sample scanner to acquire the fluorescence intensity profiles (Figure 1). A similar instrument has been used for measurements in a low-temperature cryostat\(^{17}\) and is described in more detail elsewhere.\(^{17}\) These microscopes can also be used in a near-field optical mode to excite single molecules near an aperture of a coated fiber tip, if a subwavelength resolution is required. All measurements discussed here were done in the confocal mode. Our modular microscope is based on optomechanical cubes (partec) mounted onto an optical bench (Figure 1). The cubes are well suited for the setup of a modular microscope because they ensure that no light from the outside enters the optical path, as is obviously crucial for single-molecule experiments, where minute light powers have to be detected. Lenses, filters, diaphragms, and detectors can easily be mounted and switched in and out of position.

Light from a 20 mW HeNe laser with \(\lambda = 633\) nm is coupled into the entry of a single-mode fiber. The 4-\(\mu\)m-diameter core on the end of the fiber is magnified into one of the intermediate-image planes of the microscope objective, after being attenuated by neutral filters and band-pass-filtered by an excitation filter (633NB3.0, part of Omega “dye set XF46”). By inserting a linear or circular polarizer, one can select linear or nearly circular polarized excitation light. After passing the excitation pinhole, the laser light is reflected into the objective by a dichroic mirror (Omega 645DRLP02) with a corner wavelength of 645 nm. An objective with a numerical aperture of 0.8 (partec 50 \(\times\) 0.8) focuses the laser light into the focal plane on-axis to give an excitation spot with a diameter of 0.8–1.2 \(\mu\)m at one-half the intensity maximum, as can be deduced from the spots originating from single molecules in the images (to be shown later). The fluorescence of the sample is collected by the same objective and passes the dichroic mirror and an interference band-pass filter (Omega 670DF40), which blocks the excitation light reflected from the sample and objective lenses. A pinhole of 200 \(\mu\)m diameter in the intermediate-image plane of the emission light path suppresses background light originating from locations more than \(\sim 5\) \(\mu\)m from the excitation spot. By using of smaller pinholes, the resolution as well as the background suppression could be increased further. However, because of different wavelengths of the fluorescence emitted and the use of objectives that show chromatic aberrations, the loss of fluorescence signal becomes too high when smaller pinholes are used. The detection pinhole is demagnified onto the 0.1 \(\times\) 0.1 mm\(^2\) sensitive area of an avalanche photodiode (EG&G SPMC200), which detects fluorescence photons with a counting efficiency of 40%, while the dark count rate is < 4 s\(^{-1}\). The number of photons counted in 5 ms is registered for each image pixel and displayed in a grayscale image. The total detection efficiency
of the microscope is estimated to be $1\text{–}2\%$. To determine the fluorescence polarization, we used a scheme with two avalanche photodiode detectors (Figure 1), as was also used in by Ruiter et al.$^5$ A polarization beam-splitting cube splits the two linear polarization components of the fluorescence onto the photodiodes. The photon signals of both detectors are counted simultaneously by two independent counters. In contrast to the previous study,$^5$ we used circular polarized excitation light to excite all in-plane molecular absorption dipoles, rotating or not, with the same strength.

Sample Preparation

A drop of a solution of TDI$^{18}$ molecules (Figure 2), $0.8 \times 10^{-3}$ mol/L, and PVB, 2 mg/mL, in CHCl$_3$ is spin-coated onto a cleaned quartz cover glass.$^{12}$ This yields homogeneous polymer films 20–30 nm thick and a calculated average of 12 ± 6 molecules/µm$^2$. In the fluorescence images, however, only a fraction less than 10% of this number appears as a spot, as will become clear from the images. By taking into account the photobleaching, working with this concentration makes it convenient to compare and analyze a statistically relevant number of molecules simultaneously in one experiment. The thickness of the films was measured by atomic force microscopy and scanning shear force microscopy by evaluating the depth of the holes that are occasionally present in the films. The dye molecules are well suited for excitation with an inexpensive HeNe laser or a semiconductor laser diode, given their high absorptivity between 590 and 680 nm (Figure 2).

Results

Number of Fluorescing Molecules: Simulation vs Experiment. One of the aims of our numerical simulation was to get a qualitative impression of the fluorescence images that can be obtained from distributions of single molecules. The orientation of the molecular dipole moments plays an important double role concerning the fluorescence signal reaching the detector: first, the excitation rate and, second, the fluorescence detection efficiency strongly depend on the molecular orientation relative to the optical axis.$^{20}$ Under our experimental conditions, excitation and (spontaneous) fluorescence emission are incoherent processes and can be described independently of each other. For the simulation, we used a semiclassical treatment, i.e., describing the electromagnetic field in terms of polarized electromagnetic waves. The excitation light was approximated by a linear polarized plane wave with an Airy-shaped intensity distribution. The width of this distribution corresponds to the size of the laser spot, the lower limit being set by diffraction at the aperture of the objective. Fluorescence excitation and emission as well as photon detection were described as independent stochastic processes; i.e., the quantum statistical probabilities for excitation and emission were used.

Results of the simulation of 2000 molecules on an area of $40 \times 40$ µm$^2$ on the optical axis and an acquisition rate of 200 pixels/s, Figure 3c shows a typical image obtained at room temperature. The scan direction is from bottom to top for each image. All spots in the Figure 3c have a width of 0.8–1.2 µm and are an image of the intensity distribution of the laser spot. As already mentioned, different intensities of the spots can result from different molecular orientations with respect to the excitation polarization and from differences in excitation or emission yields. The counting rate in the brightest spots reaches 300/5 ms, which, taking the detection efficiency of 1–2% into account, corresponds to (3–
6) \times 10^6 \text{ fluorescence photons/s. This is 10-fold less than the}
\text{saturation rate of TDI of } R_\infty = k_{21}(1 + k_{isc}/k_{21})(2 + k_{isc}/k_{T}) =
82 \times 10^6 \text{ s, calculated from the transition probabilities given}
\text{earlier}.^{18,19} \text{ The corresponding binary image of these data}
(Figure 3d) shows only 57 \pm 10 bright spots that exceed one-
half of the maximum intensity in the image (above the
background). However, the concentration of dye molecules in
the PVB film is 12 \pm 6 molecules/\mu m^2, and the total number
of molecules on the imaged area is 10 \times 10^3 \text{ to } 30 \times 10^3.
This ratio of bright spots to the total number of molecules, 0.3\% \pm 0.2\%,
is to be compared with the ratio of 5\% between bright molecules
and all molecules in the simulated images in Figure 3b, where
a completely random distribution of the dipole moments was
assumed.

With the low-temperature setup, similar images were also
acquired at \text{T} = 202 \text{ and 96 K (results not shown). Down to}
\text{temperatures as low as 96 K, we observed no temperature}
dependencies, e.g., an increased fluorescence yield. Differences
in the counting rates are attributed to slightly different illumina-
tion adjustments. From the comparison of the experimental with
the simulated data, and the fact that no inhomogeneity of the
samples was found on a scale up to a few hundred micrometers,
one can be sure that each spot represents no more than a few
molecules. In fact, there is a strong evidence that each spot
represents the light from exactly one molecule, as will be shown
now.

\textbf{Blinking and Bleaching in Fluorescence Images.} Figure
4a shows a series of 20 scans of the same area, taken over 5 h.
In these images, most of the bright spots instantaneously
disappear during the first scans, which is characteristic for single
molecules bleaching in one step through transition to a
nonfluorescing state, in contrast to the continuous exponential-
like bleaching curve of the fluorescence of an ensemble of many
molecules. Two other effects can be seen in the images, or
even better, when the images are combined in sequence to make
a movie: First, some spots reappear after having disappeared
for some time, or show dark stripes, demonstrating a discrete
“on–off–on” switching process. This blinking is a character-
istic effect in the fluorescence light of single molecules. Second,
the intensity of some spots, in comparison with the others,
changes slowly between the images. During the 5-h experiment,
we observed no diffusion out of a range of 200 nm. Plotting
the intensity of each of the 20 images, integrated over the whole
image against the image number or time, yielded a nearly
exponentially decaying curve for bleaching, as would be
expected for a molecule ensemble (Figure 4b). A slight
variation of the experimental data from an exact exponential
decay is to be expected because of the limited number of
molecules contributing to the total intensity and the possible
inhomogeneities in the various surrounding molecules. Another
effect not considered here are differences in the excitation rates
for the randomly oriented molecules, for which the bleaching
decay of the ensemble would be expected to be a sum of
decaying exponentials rather than a single exponential.

The high background intensity in the bleaching curve,
compared with the fairly good signal-to-background ratio (in
Figure 4a) of 5:1 (related to the counts in single pixels), is a consequence of the small overall number of fluorescing dye molecules in the imaged volume of \(20 \, \text{nm} \times 40 \, \mu\text{m} \times 40 \, \mu\text{m}\), compared with the number of PVB molecules, which contribute to the background. The ratio of TDI molecules to PVB molecules is on the order of \(10^3:10^{10}\). In experiments with shear force and atomic force microscopes, holes were sometimes detected in the PVB film. The black spot in the background in Figure 4a is interpreted to indicate such a hole in the polymer film. This is a hint that impurities in the PVB film or in the PVB itself contribute to the fluorescence signal more strongly than does the nonideal excitation light suppression by the filters. In the series of the 20 images, some molecules remain fluorescing during the experiment. Interestingly, some of these “most-stable” molecules (arrows) are comparatively bright, emitting, in total, many more fluorescence photons than other dim but fast-bleaching molecules.

**Photostability of TDI Molecules.** Figure 4c is an evaluation of \(N = 29\) molecules from which the total number of counts obtained in the 20 images in Figure 5a are classified into a
histogram. For some molecules, >300,000 counts were registered; taking the detection efficiency of 1–2% into account, this corresponds to at least $1.5 \times 10^7$ fluorescence photons at a fluorescence rate as great as $6 \times 10^6$ s for an excitation intensity of $I = 300 \mu W/\mu m^2$. Taking into account the scanning process, where only a fraction $[1 \mu m^2/(40 \mu m \times 40 \mu m) = 1/1600]$ of the image area is illuminated at any given time, the average excitation intensity is $I = 1/1600 \times 300 \mu W/\mu m^2 = 0.19 \mu W/\mu m^2$. The lifetime of the ensemble, determined as the 1/e decay time in Figure 4b, is 60 min. The lifetime for single molecules (during continuous illumination), determined by the photon number and the rate from above, is 2.5 s. The product of lifetime and intensity should be constant. Indeed, for single molecules it is similar to that of the ensemble, since $2.5 \times 300 \mu W/\mu m^2 \approx 60 \times 0.19 \mu W/\mu m^2$.

From the histogram can be deduced a broad distribution of this total fluorescence yield. For some molecules, which bleached during the experiment, <1/8 of the counts of the bright and stable molecules were detected. This may simply be an effect of fundamental statistics: If the ensemble shows an exponential decay, then even if the molecules and their surroundings are identical (as in a homogeneous ensemble), most of the molecules must be less stable and bleach at the beginning of the experiment, while only few survive to the end. However, strong inhomogeneities between molecules or in their local environments can also be responsible for these widely differing total photon yields. Another reason for the different number of counts before bleaching is the dependence of the detection efficiency on the orientation of the molecules, which was not taken into account in the calculation of the minimum number of emitted photons above.

Blinking and Bleaching in Timeline Scans. To get a more quantitative insight into this effect, we repeatedly scanned intensity profiles $C(x, t)$ above one line on the sample (Figure 5a). This view allowed us to follow the fluorescence intensity of several single molecules over a long period quasi-simultaneously. The scan profiles were 40 μm long. The acquisition time for each line was 2.6 s. In total, 768 lines were collected in 2000 s. Molecules that are fluorescing constantly appear as a continuous line (E), whereas other molecules switch “off” and “on,” on a time scale of seconds (D) to many minutes (A). Figure 5b shows intensity vs time curves $C(x, t)$ extracted from Figure 5a by summing the intensity over a range of $\Delta x = 8$ pixels at different positions A–F, again showing the “inhomogeneity” in the behavior of the molecules. Jump heights of different intensities can be explained by different molecular orientations with respect to the optical axis. For some time, traces A and D show a second or third intensity level, perhaps

Figure 6. (a, b) Linear polarization components of fluorescence light in single-molecule images with circular polarized excitation. Both images were taken simultaneously with two independent photon counters. On–off transitions occurring during the scan (arrows) appear simultaneously in (a) and (b). (c) Polarization distribution calculated from (a) and (b), with the low-intensity regions of the image blanked out to better visualize the positions of the molecules. (d, e) Temporal dependence of the fluorescence polarization for single molecules [circles A–F in (c)], as deduced from a series of 20 images: (d) blinking, and (e) nonblinking molecules. For all of the molecules, the polarization does not change by >15–200° during the 5 h of observation, which excludes a molecular rotation as the reason for the on–off–on transitions.
from two or three molecules contributing to the fluorescence. Summing the intensities in Figure 5a over the scan lines yields Figure 5c. Although only a few molecules contribute to the fluorescence, the intensity decay curve compares well with the exponential decay curve (smooth curve) that would be roughly expected for a molecule ensemble.

**Fluorescence Polarization.** In another experiment we studied the polarization of the fluorescence light from single molecules, when they were excited by circularly polarized light. The emission polarization for each spot directly corresponded to the molecular emission dipole component parallel to the object plane. Two images of both linear polarization components in the microscope were simultaneously acquired, as reported elsewhere. By calculating the inverse tangents from the relation of the intensities in the two detection channels for each pixel, we deduced an image of the polarization distribution. Figure 6 shows an image of the x- and y-polarized light and the corresponding polarization image, where in the polarization images the low intensities of the fluorescence are blanked out. By comparing Figures 6a and 6b, we can see that molecules simultaneously blink and bleach in both polarization channels: Spots disappear and reappear at the same scan position (arrows). Figures 6d and 6e show the time-dependence of the orientation of single molecules, which is extracted from a series of 20 polarization images. For all molecules, whether blinking (Figure 6d) or stably emitting (Figure 6e), the change of polarization over the 5 h is <20°; for most molecules, it is <15°. This very slow rotation of some molecules can be responsible for small and slow changes in intensity. However, since obviously the dye molecules make no strong rotation in the polymer film, then for our preparation, a rotation of the molecules as the origin for the blinking can be excluded on a time scale of hours. Consequently, another mechanism must be responsible for the long-time blinking.

**Discussion**

The observation of fluorescence intensity and polarization of single dye molecules over a period of seconds to several hours uncovers short- and long-time blinking, which cannot be detected in ensemble measurements. Our polarization measurements exclude fast rotation of the complete molecule as the reason for instantaneous intensity changes. However, very slow intensity changes, which are observed for very few molecules, can be explained by a slow molecular rotation, probably together with the surrounding polymer strands. The question remains, as to the origin of the instantaneous blinking behavior. Several models have been discussed in the literature. Investigating with a near-field microscope, attributed small translational shifts of rhodamine 6G in PVB to a molecular rotation. Xie and Dunn measured the molecular rotation of sulfurhexamol 101 (dispersed on glass) on the time scale of a minute but found no rotation. Studying Dif (diocodectyletrimethylindocarbocyanine) molecules in PMMA (poly(methyl methacrylate)), found a very slow molecular rotation on the time scale of 1 h; they attributed blinking to temporary transitions to dark states or spectral jumps. In this study, a sudden rotation was found to be unlikely. In single-molecule emission spectra, strong spectral shifts in the range of several nanometers were observed. Shifts in the excitation spectrum may explain one-step changes in the fluorescence intensity as well as slow changes. These spectral changes could be due to conformational changes of the molecules or of their surroundings. For the TDI molecules, spectral changes are expected for rotation of the substituents on the nitrogen atoms (Figure 2). For free molecules, a specific orientation of the substituents in relation to the main molecular frame to form the π-system, which is responsible for the optical transition, is energetically favorable. In the PVB film, we must assume that the ability of the endgroups to rotate is strongly influenced by the individual local surroundings formed by the polymer strands. Some substituents might never have the chance to be in a favorable orientation with respect to the π-system because of limited space and thus would prevent an excitation by the laser. This could help account for the discrepancy between the number of fluorescing molecules observed and the number of molecules expected to be on the surface. Apart from spectral jumps, another possible explanation of the temporary “off” states could be temporary reactions with surrounding molecules. For example, the temporary presence of O2 in the vicinity of the dye molecule could, by influencing the symmetry of the quantum mechanical levels of the molecule, reduce the excitation transition probability and consequently decrease the fluorescence intensity to a very small amount. After some time, which would vary stochastically between individual molecules because of fundamental statistics or different local surroundings, the oxygen could be released from the molecule and the fluorescence would reappear instantaneously. An experimental test of this model would be to study the dependence of the blinking behavior on the oxygen concentration. Recent results of such experiments show that the blinking is indeed strongly influenced by the surrounding oxygen concentration.

**Conclusions**

TDI molecules, diluted in a PVB polymer film, were investigated over a span of 5 h. Characteristic discrete long-time blinking and bleaching in the fluorescence demonstrates that the distribution of single TDI molecules can be imaged with our modular confocal microscope with high quality over a long time. No diffusion out of a range of 200 nm was observed during 5 h. Some TDI molecules delivered >1.5 × 105 fluorescence photons without bleaching. The fluorescence, summed for several molecules either in a series of 20 successively scanned images or in time-line scans, decayed in a nearly exponential way, as would be expected for a molecule ensemble. Polarization measurements of the fluorescence light with two parallel counters allowed us to follow molecular rotations by using circular polarized excitation light. For TDI in PVB, the polarization direction did not change by >20° over the 5 h, which excludes molecular rotation as the origin of the instantaneous blinking. The influence of oxygen on the blinking and bleaching will be topic of a forthcoming paper.

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