Solvation Dynamics of DCM in Lipid

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Solvation dynamics of 4-(dicyanomethylene)-2-methyl-6(ρ-dimethylaminostyryl)-4H-pyran (DCM) in dimyristoylphosphatidylcholine (DMPC) vesicles in water is studied by using the picosecond time-resolved Stokes shift technique. In unilamellar DMPC vesicles, DCM exhibits wavelength-dependent fluorescence decays with a fast decay at the blue end and a slow decay preceded by a distinct growth at the red end of the emission spectrum. The solvation dynamics of DCM in DMPC vesicles is found to be biexponential with two components 230 ps (40%) and 1.6 ns (60%) and is attributed to the relaxation of the water molecules inside the water pool of DMPC vesicles.

1. Introduction

One of the longstanding goals of chemistry and biology is to elucidate the dynamics of various processes occurring in complex biological assemblies. Most processes in biological systems occur in a confined region having dimensions of a few nanometers. Confinement in a small space and the severely altered local properties markedly modify dynamics of many chemical processes. As a result, chemistry in biological and confined environments is often very different from that in bulk water. The behavior of water in such a confined environment plays a fundamental role in many natural processes. In recent years several groups have studied solvation dynamics and dielectric relaxation of water in many organized and constrained media such as cyclodextrin, proteins, microemulsions, micelles, water surface, and lipid vesicles. Of these, the study of lipid vesicles is most relevant to understanding the behavior of water in a biological cell. In a vesicle, an aqueous volume is surrounded by a bilayer of surfactants and is dispersed in bulk water. In a recent paper, we have demonstrated that for coumarin 480 (C-480) in dimyristoylphosphatidylcholine (DMPC) lipid the solvation dynamics exhibits a component of 600 ps. This is substantially slower than the subpicosecond (0.3 ps) solvation dynamics exhibited in bulk water. In the present work, we report on the ultrafast dynamics of the laser dye 4-(dicyanomethylene)-2-methyl-6(ρ-dimethylaminostyryl)-4H-pyran (DCM, I) in DMPC vesicles. Recent femtosecond studies reveal that solvation dynamics of DCM in methanol exhibits an inertial component on the 100 fs time scale and another of a few picoseconds. The structure of DCM suggests the possibility of various ultrafast processes, such as isomerization about the olefinic double bond, twisted intramolecular charge transfer (TICT), apart from the ordinary solvation dynamics. Because of the TICT process, the excited-state dipole moment of DCM (23.6 D) is very much higher than that in the ground state. It is obviously of interest to find out how these ultrafast processes are affected in organized assemblies. The TICT process is usually slowed in various organized assemblies, and thus it is important to know whether the TICT process is retarded sufficiently to exhibit dual emission. In a recent work, we studied DCM in AOT microemulsions. The microemulsions are basically nanometer sized water droplets surrounded by a surfactant layer and dispersed in bulk n-heptane. We observed that DCM does not exhibit dual emission in AOT microemulsions and exhibits only the TICT emission band. Furthermore, it is observed that in AOT microemulsions DCM displays slow solvation dynamics with an average solvation time of 1.23 ns. In the present work, we report on the photophysical processes of DCM in DMPC lipids.

2. Experimental Section

DCM (laser grade, Exciton) and DMPC (Sigma) were used as received. Absorption and emission spectra were recorded by a JASCO 7850 and a Perkin-Elmer 44B instruments, respectively. For lifetime measurements, the sample was excited at 300 nm with the second harmonic of a cavity dumped rhodamine 6G dye laser (Coherent 702-1) pumped by a cw mode locked Nd:YAG laser (Coherent Antares 76s). The emission was detected at magic angle polarization, using a Hamamatsu MCP photomultiplier (2809U). The full width at half-maximum of the instrument response at 300 nm is ≈80 ps. Fluorescence decays were deconvoluted by using global lifetime analysis software (PTI).

The lipid was prepared following the methanol injection method. Two milligrams of DMPC was dissolved in 50 μL of a solution of DCM in methanol. Using a microliter syringe, the whole solution was rapidly injected into tris buffer of pH 7.4. In this way the methanol is diluted almost instantaneously in water and the phospholipid molecules are dispersed evenly throughout the medium. The mixture is kept for 1 h at 30 °C. The concentration of the final solution of DMPC was 1 mM.

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and DCM concentration was $1 \times 10^{-5}$ M so that the probe-to-
lipid ratio was 1:100. This procedure yields a high proportion
of small unilamellar vesicles of diameter ca. 50 nm.\textsuperscript{10c,d} Emission
quantum yields were obtained by using the reported quantum
yield (0.44) of DCM in methanol.\textsuperscript{14a} All steady-state and time-
resolved measurements were done at 26 °C, i.e., above the gel
transition temperature of DMPC (23 °C).\textsuperscript{10}

3. Results

3.1. Steady-State Spectra. In the DMPC lipid, DCM exhibits
an absorption spectrum with two peaks at 445 and 470 nm
(Figure 1a) and a strong emission band at 615 nm with
$\phi_i = 0.24$ (Figure 1b). It is readily seen that DCM does not show
dual emission in DMPC. The emission maximum of DCM
exhibits a red shift with increase in solvent polarity, from 530
nm in $n$-heptane to 620 nm in methanol. DCM is insoluble in
water and hence, the steady-state spectral properties of DCM
in lipid could not be compared with those in water. However,
the observed emission maximum of DCM in lipid is similar to
that in methanol.

3.2. Time-Resolved Studies. The fluorescence decay of DCM
in DMPC exhibits a marked wavelength dependence. At the
blue end a fast decay is observed, while at the red end the decay
is preceded by a growth (Figure 2). Such a wavelength depend-
ence is typical of molecules undergoing solvation dynamics.\textsuperscript{17}
From the parameters of best fit for the emission decays and
using the steady-state emission spectra, time-resolved emission
spectra (TRES, Figure 3) have been constructed following the
procedure described by Fleming and Maroncelli.\textsuperscript{17c} The solva-
tion dynamics is described by the response function $C(t)$ which is
defined as

\begin{equation}
C(t) = \frac{\nu(t) - \nu(\infty)}{\nu(0) - \nu(\infty)}
\end{equation}

The decay of $C(t)$ is shown in Figure 4. The decay parameters
of $C(t)$ are summarized in Table 1. It is readily seen that the
total Stokes shift ($\nu(0) - \nu(\infty)$) is 1250 cm$^{-1}$ and the decay of
$C(t)$ is biexponential with two components 230 ps (40%) and
1600 ps (60%) leading to an average solvation time, $<t_\alpha>$
($=a_1f_1 + a_2f_2$) of 1050 ps.

4. Discussion

As noted earlier, DCM does not exhibit dual emission in the
DMPC lipid. The observed emission at 615 nm is very close to
that of the TICT band of DCM in polar solvents.\textsuperscript{13,14} This
indicates that the TICT process of DCM remains ultrafast in
the lipid.

The time-dependence Stokes shift (TDSS) indicates that DCM
exhibits a $\Delta\nu = 1250$ cm$^{-1}$ in the DMPC lipid. It may be noted
that recent femtosecond studies indicate that DCM exhibits a
$\Delta\nu = 3800$ cm$^{-1}$ in methanol and 2400 cm$^{-1}$ in ethylene
glycol.\textsuperscript{13b,c} Though the total Stokes shift depends on solvent, in
our apparatus with limited time resolution ($\sim$80 ps) one cannot
eliminate the possibility of missing a large part of solvation
which occurs in the femtosecond time scale.\textsuperscript{13,14} Despite this
limitation it is evident that the solvation dynamics of DCM in
DMPC exhibits a component of about 1 ns, which is substan-
tially slower than those obtained in the femtosecond studies.\textsuperscript{13,14}

In a lipid vesicle, there are three possible locations of the probe,
bulk water, inner water pool, and bilayer. Since DCM is
insoluble in water the possibility of DCM in bulk water is ruled
out. In a hydrocarbon the emission quantum yield of DCM is
extremely low (0.01) and the lifetime is very short (<50 ps).\textsuperscript{16}
Further, in $n$-heptane the emission maximum of DCM is very
much blue-shifted to 530 nm and the decay does not exhibit a
wavelength dependence.\textsuperscript{15} Thus the DCM molecules staying in
the "dry" hydrocarbon-like bilayer is not expected to contribute to
the observed solvation dynamics. Thus the solvation dynamics
appears to be due exclusively to the DCM molecules in the
inner water pool. The observed solvation time of 1 ns is very
similar to that of DCM obtained in the water pool of the
microemulsion\textsuperscript{16} and is very close to the 600-ps component
reported for DMPC using C-480 as a probe.\textsuperscript{9} This indicates
solvent relaxation time of the water molecules confined by the
lipid bilayer is 1 ns. This is substantially slower than the
solvation dynamics in ordinary water. Kaatze and co-workers reported that lipid vesicles exhibit two prominent dielectric relaxation time (τD) in 10 and 0.1 ns time scales. They assigned the 10-ns relaxation to the solvent relaxation time. Since in the continuum model solvation time, τS = (ε∞-ε0)τD, if one assumes ε∞ of the inner water pool same as that of water, i.e., 5, and ε0 is of the inner water pool same as that of methanol (≈ 40), one immediately calculates a solvation time of (5/40) × 10 ns, i.e., 1.25 ns, which is remarkably close to the observed solvation time of 1 ns. The role of methanol appears to be minor because the overall concentration of methanol added is about 2%, most of which should evaporate during preparation of the lipid. Further, the solvation dynamics of DCM and other probes in methanol occurs with an inertial component in the 100-fs time scale and another in the <10-ps time scale. Thus the observed nanosecond dynamics cannot be because of traces of methanol. Again, the polarity of the inner pool for lipids prepared by the methanol injection method is similar to that of sonicated unilamellar vesicles.

5. Conclusion

The present work shows that DCM does not exhibit dual emission in the DMPC lipid. This suggests that the TICT process of DCM remains ultrafast in lipids. The solvation dynamics indicate that in the inner water pool the water molecules relax on a 1-ns time scale. The solvation dynamics studied is consistent with the previous dielectric relaxation studies in lipids. Since the water molecules inside the water pool of vesicles resemble those in biological cells, this study demonstrates that the dynamics of biological water molecules is substantially slower than that of ordinary water molecules.

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References and Notes