

UC Irvine

UC Irvine Previously Published Works

Title

CAPRYDAA, an anthracene dye analog to LAURDAN: a comparative study using cuvette and microscopy

Permalink

<https://escholarship.org/uc/item/69x536s3>

Journal

Journal of Materials Chemistry B, 8(1)

ISSN

2050-750X

Authors

Castro-Castillo, Vicente
Gajardo, Javier
Sandoval-Altamirano, Catalina
[et al.](#)

Publication Date

2020-01-07

DOI

10.1039/c9tb01738k

Peer reviewed



HHS Public Access

Author manuscript

J Mater Chem B. Author manuscript; available in PMC 2021 January 07.

Published in final edited form as:

J Mater Chem B. 2020 January 07; 8(1): 88–99. doi:10.1039/c9tb01738k.

CAPRYDAA, an anthracene dye analog to LAURDAN: a comparative study using cuvette and microscopy

Vicente Castro-Castillo¹, Javier Gajardo¹, Catalina Sandoval-Altamirano¹, Enrico Gratton³, Susana Sanchez², Leonel Malacrida^{3,4}, German Gunther¹

¹Universidad de Chile, Facultad de Ciencias Químicas y Farmacéuticas, Departamento de Química Orgánica y Físicoquímica, Casilla 233, Santiago 1, Chile.

²Universidad de Concepción, Facultad de Química, Departamento de Polímeros, Concepción, Chile.

³Laboratory for Fluorescence Dynamics, Biomedical Engineering Department, University of California at Irvine, Irvine, CA, USA.

⁴Departamento de Fisiopatología, Unidad de Microscopia Avanzada y Bifotónica, Hospital de Clínicas, Facultad de Medicina, Universidad de la República, Montevideo, Uruguay.

Abstract

We synthesized an anthracene derivative with solvatochromic properties to be used as a molecular probe for membrane dynamics and supramolecular organization. A ten carbon atom acyl chain and a dimethylamino substitution were introduced at positions 2 and 6 of the anthracene ring, respectively. This derivative, (2-nonanoyl-6-(dimethylamino) anthracene (termed CAPRYDAA), is a molecular probe designed to mimic the well-known membrane probe LAURDAN's location and response in the lipid membranes. Due to the larger distance between the electron donor and acceptor groups, its absorption and emission bands are red-shifted according to the polarity of the media. The photophysical behavior of CAPRYDAA was measured in homogeneous media, synthetic bilayers and cells, both in a cuvette and in a fluorescence microscope, using one and two-photon excitation. Our results show a comparable physicochemical behavior of CAPRYDAA with LAURDAN, but with the advantage of using visible light (488 nm) as an excitation source. CAPRYDAA was also excitable by two-photon laser sources, making it easy to combine CAPRYDAA with either blue or red emission probes. In GUVs or cells, CAPRYDAA can discriminate the lipid phases and liquid-liquid phase heterogeneity. This new membrane probe shows the bathochromic properties of the PRODAN-based probes designed by Weber, overcoming the need for UV or two-photon excitation and facilitating the studies on the membrane properties using regular confocal microscopes.

Correspondence to: Leonel Malacrida (lmalacrida@hc.edu.uy), German Gunther (ggunther@ciq.uchile.cl).

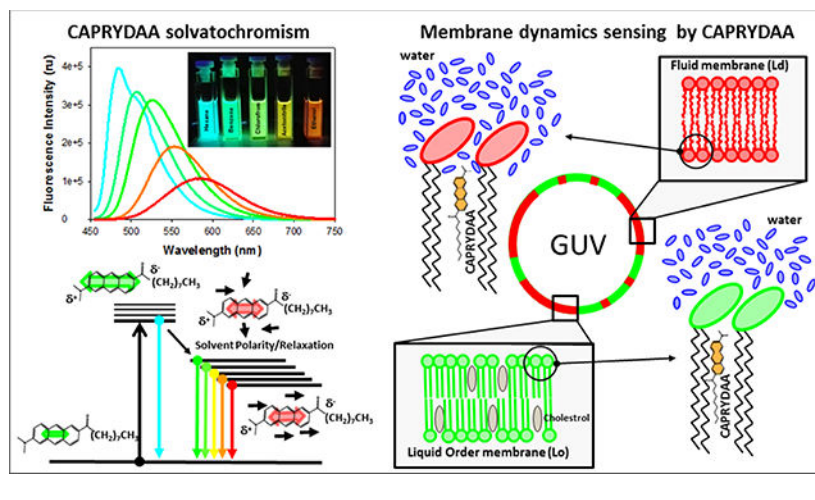
Author contribution

JG and VCC synthesized the probe; GG, LM and SS designed the experiments; CSA, GG, SS and LM performed the experiments; CSA, SS, GG and LM analyzed the data; SS, GG and LM wrote the manuscript; EG, SS, GG and LM revised and discussed the manuscript.

Conflict of interest

The authors declare no competing interests.

Graphical Abstract



1. INTRODUCTION

Fluorescent probes are essential tools in a wide range of biological research. In particular, environment-sensitive hydrophobic fluorescent probes are widely used in the study of molecular organization in the membranes.^{1–3} According to Weber's definition, an environmental probe must contain, in its molecular structure, at least two different substituent groups or organic functionalities, both attached to an aromatic ring, with one of them having an electron acceptor character and the other with an electron donor character.⁴ Generally, the magnitude of the spectral changes shown by the probe is directly related to the spatial separation between both substituent groups as well as to the basic molecular entity. The structure of the series of polarity sensitive fluorescent probes, designed and synthesized by Weber *et al.*⁴ in the late seventies, which undergo intramolecular photoinduced charge transfer, contains a naphthalene aromatic moiety substituted with a dimethylamino group in position 2 (donor group) and an alkyl carbonyl group in position 6 (acceptor group). PRODAN and LAURDAN, among others, belong to this family of probes. LAURDAN has been widely used, both in cuvettes and in microscopes, for the study of the synthetic membranes and live cells.^{5–17} These studies include the characterization of lipid phases in the membranes, determination of partition coefficients,¹⁸ observation of shape changes during the phase transition,¹⁹ observation of lipid-protein interactions,²⁰ and discrimination of cholesterol dynamics in living cells.²¹ The complex photophysics and solvatochromism properties of this family of compounds involve the contribution of solvent dielectric relaxation and intramolecular excited-state reactions. The separation of these two contributions allows a detailed characterization of the processes.²² In fact, the analysis of the spectral data using phasor analysis allows this distinction, making phasor analysis a very useful tool to describe the complex systems.^{16, 17, 23} At this point, it is important to acknowledge that several researchers have developed other analogs of the LAURDAN/PRODAN family for imaging lipid organization in cell membranes and model membrane systems. Among them, the fluorine-based push-pull probes^{24, 25} and push-pull pyrene derivatives²⁶ show a large shift in the emission as a function of the lipid order.²⁷

The LAURDAN's emission is mainly in the wavelength range between 400 and 530 nm, with maximum excitation at 370 nm. The possibility to shift absorption/emission to longer wavelengths is a major goal since such a red-shift will have advantages for biological imaging, including facilitating light penetration into tissues, reducing auto-fluorescence, and opening a different wavelength window for combination with other biological fluorescent markers. An option to achieve this purpose is to increase the number of aromatic rings in order to enlarge the distance between substituents involved in the intramolecular charge transfer. This methodology has been used to fabricate the anthracene derivative dyes. The excitation wavelengths of these derivatives are red-shifted (near-infrared in two-photon excitation),²⁸ and they show high quantum yields. As an example of this strategy, we can mention the PRODAN (2-propionyl-6-dimethylaminonaphthalene) homologs and Anthradan (6-Propionyl-2-(dihexylamino)anthracene), whose excitation and emission wavelengths shift around 100 nm and 80 nm, respectively. Further, the bathochromic displacement of the absorption bands was accomplished when a symmetric diaryl substitution on the positions 9 and 10 of the anthracene moiety was performed. These substituted probes (analogous of Acedan (2-acetyl-6-(dimethylamino) naphthalene) have been used for two-photon microscopy imaging.²⁹

In this report, we synthesized an anthracene derivative, with an acyl chain of ten carbon atoms and a dimethylamino substitution at positions 2 and 6 of the anthracene ring, respectively. The logic behind this synthetic design was to obtain a molecule with a similar membrane location and response to the environment as that of the LAURDAN probe, but with a bathochromic shift in the absorbance and emission bands.

We report here the synthesis, spectroscopic characterization, and applications of (2-nonanoyl-6-(dimethylamino) anthracene, CAPRYDAA. The photophysical behavior was measured in several media (homogeneous, synthetic bilayers, and cells), with the measurements being carried out both in cuvettes and under the microscope, using one- and two-photon excitation to establish its applicability under different experimental techniques and conditions as well as to compare with the LAURDAN probe. Our results show a comparable behavior of CAPRYDAA with LAURDAN, but with the advantage of using visible light (488 nm) for excitation. Moreover, CAPRYDAA was fully excitable with two-photon laser sources, due to which CAPRYDAA can easily combine with either blue or red emission probes. Further, by using CAPRYDAA, it was possible to discriminate lipid phases in GUVs as well as liquid phase heterogeneity in living cells. Therefore, here, we present a novel probe that extends the possibilities for studying the membrane dynamics using the fluorescent probes for *in vitro* as well as *in vivo* experiments and combining the opportunity to work in cuvettes and with imaging. The major advantage of CAPRYDAA is the possibility of working with single-photon excitation, thereby avoiding the need for 2-photon or UV illumination.

2. EXPERIMENTAL

2.1. Materials

All the reagents and solvents were purchased from Sigma-Aldrich (St. Louis, MO, USA) or Merck (Darmstadt, Germany) and used without further purification.

Diioleoylphosphatidylcholine (DOPC), dipalmitoylphosphatidylcholine (DPPC), and cholesterol (Chol) were obtained from Avanti Polar Lipids (Alabaster, AL, USA) and used as received. Melting points were determined on a Reichert Galen III hot plate with a DUAL JTEK Dig–Sense thermocouple thermometer and were uncorrected. The ^1H and ^{13}C NMR spectra were recorded on Bruker Avance spectrometers at 200 or 400 MHz and 50 or 100 MHz, respectively, using CDCl_3 or $\text{DMSO}-d_6$ as solvents unless indicated otherwise. The chemical shifts were reported in terms of δ (ppm), downfield from TMS. The coupling constants (J) were given in Hz. The analytical TLC was performed on Merck silica gel 60 F254 foil-backed plates. The solvents used as eluent were specified in each case. The chromatographic purification of all the products was carried out on the silica gel columns, and the solvents used were specified in each case.

2.2 Methods

Synthesis of 2-nonanoyl-6-(dimethylamino) anthracene, CAPRYDAA

Figure 1 and scheme 1 shows the molecular structure and the synthetic path of CAPRYDAA, respectively. More details are given in the supplementary material.

Briefly, the synthesis of **6** was carried out by using a method reported in the literature.¹ Compound **2** was synthesized in good yield by the Sandmeyer reaction, using *tert*-butyl nitrite in dry acetonitrile. The treatment of **2** with CsF in DMSO afforded 2-bromo-6-fluoro-9,10-anthraquinone, which was treated, without purification, with aqueous dimethylamine to afford 2-bromo-6-dimethylamino-9,10-anthraquinone (**3**). The subsequent sodium borohydride reduction followed by the dehydration of **3** and further subjection to a Rosenmund-von Braun reaction yielded **5**. Finally, **6** was prepared by the nucleophilic addition of octylmagnesium bromide to the cyano compound followed by the hydrolysis of the imine intermediate (more details and NMR spectra are given in the Supplementary Section).

Cuvette spectroscopy measurements—The UV-Vis spectra were recorded on an Agilent 8453 Diode-Array spectrophotometer (Agilent, USA) in the range of 250–700 nm. The fluorescence emission spectra were obtained by using an ISS PC1 spectrofluorometer (ISS, Champaign IL) or a FluoroMax 4CP (Horiba Jobin Yvon, Japan) at room temperature. Further, the fluorescence lifetime measurements were carried out by using the time-correlated single-photon counting (TCSPC) method on a PicoQuant FluoTime 200 fluorescence lifetime spectrometer (PicoQuant, Germany) equipped with a multichannel scaler, PicoQuant's Timeharp 250 (Pico Quant, Germany). LEDs or lasers of appropriate wavelengths were employed as excitation sources. The fluorescence quantum yields were measured in a FluoroMax 4CP (Horiba Jobin Yvon, Japan) spectrofluorometer equipped with the Quanta-Phi integrating sphere.

Partition constant measurements—The water/dodecane partition for CAPRYDAA was measured by HPLC. The solution was allowed to equilibrate for 48 h and before the measurement, the aqueous and organic phases were centrifuged at 3000 rpm for 10 min (Hermle Z200A) to split the phases properly. The partition constant was obtained from the ratios of the areas of the chromatographic peaks obtained in a Waters 600 HPLC system.

The HPLC was equipped with a Waters 996 PDA pump, the column used had a 1.15 μm pore, 100 \times 4.6 mm Merck Chromolith RP-18 (Merck KGaA, Darmstadt, Germany), and the mobile phase and chromatographic conditions were: isocratic acetonitrile having a flow rate of 1 mL/min with subsequent detection at 315 nm.

One- and two-photon lifetime and hyperspectral microscopy—The samples were incubated with either 1 μM of CAPRYDAA or LAURDAN for 30 min at 37 $^{\circ}\text{C}$. The fluorescence lifetime imaging (FLIM) and the hyperspectral microscopy were acquired using a Zeiss LSM880 laser scanning microscope (Carl Zeiss, Dublin, CA-USA). The instrument was equipped with a 20 MHz pulsed supercontinuum white laser (A2, SC390, Fianium Inc, USA) for 488 nm excitation and a Ti:Sapphire laser (Spectra-Physics, Newport-CA-USA) with 80 fs pulses and 80 MHz repetition rate for two-photon excitation. The calibration of the lifetime measurements was performed using a 100 nM solution of Coumarin 6, having a reference lifetime of 2.5 ns (www.iss.com). For FLIM data acquisition, an ISS A320 FastFLIM box (ISS, Champaign, USA) was used to collect the lifetime decay data. The images were acquired using a 40 \times water immersion objective 1.2 N.A. (Carl Zeiss). CAPRYDAA was excited using either a 488 nm or 900 nm excitation sources. LAURDAN was excited using 780 nm excitation source. For the FLIM measurements, two high-efficiency GaAsP Hybrid Detectors (HPM-100-40, Becker & Hickl GmbH) were used, and before the detector, a filter cube composed of two bandpass filters (440/60nm, channel-1, and 500/60nm, channel-2) and a 470 nm dichroic mirror from Semrock was employed. We defined channel-1 and channel-2 as blue and green filters, respectively. For the hyperspectral imaging, the lambda mode configuration of the Zeiss LSM8800 was used. A 32-channel configuration was used with a bandwidth of 9.7 nm, and the total range acquired was between 410 nm and 728 nm. The image acquisitions in both the configurations (FLIM and hyperspectral) were performed with a pixel frame size of 256 \times 256 and a pixel dwell time of 16.38 μs /pixel. The hyperspectral and FLIM data were processed using the SimFCS software developed at the Laboratory for Fluorescence Dynamics (www.lfd.uci.edu).

GUVs preparation—Giant unilamellar vesicles (GUVs) were prepared by following the electroformation protocol described by Pott *et al.*³⁰ Briefly, dioleoylphosphatidylcholine (DOPC) or dipalmitoylphosphatidylcholine:cholesterol (1:1 molar, DPPC:Chol) was prepared at a final concentration of 0.3 mM in chloroform. CAPRYDAA or LAURDAN at 0.5% molar was added to the lipid mixture in chloroform. Then, 4 μL of the organic mixture was applied to each platinum wire and dried under vacuum overnight. The electroformation chamber was filled with 400 μL of 200 mM sucrose at 50 $^{\circ}\text{C}$. To keep the temperature constant, we used a water circulating bath. For the GUVs growth, we applied a sinusoidal potential of 2 volts and 10 Hz for 1.5 hours. To detach the GUVs from the wire, the frequency of the sinusoidal potential was reduced to 1 Hz for 10 min, and then, the function generator and circulating bath were turned off. For the measurements, 50 mL of the GUVs dispersion were transferred to an 8-well bottom-glass imaging chamber, containing 300 μL of a 200 mM glucose solution. The imaging chamber was previously coated with 1 mg/mL of the bovine serum albumin (BSA) solution (to avoid vesicle burst on settling), and before transferring the GUVs, each well was washed 10 times with 200 mM glucose solution.

Cell culture—The NIH-3T3 fibroblasts derived from mouse (ATCC® CRL-1658™) were grown at 37 °C, with 5% CO₂ in Dulbecco's modified Eagle's medium (Thermo Fisher Scientific Inc. Huntington Beach, CA-USA), supplemented with 10% fetal bovine serum, and 5 mL of Pen-Strep. The cells (2 × 10⁴ cells/dish) were plated 24 h before the experiments onto fibronectin-coated 35 mm glass-bottom dishes (MatTek Corporation, MA-USA). On the day of the experiments, the cells were incubated for 30–60 min with a final solution of 3 μM CAPRYDAA or LAURDAN. Before imaging, the dyes were removed by replacing it with a fresh medium. The concentration of LAURDAN was determined using a molar extinction coefficient of 20,000 M⁻¹ cm⁻¹ at 364 nm in methanol. During the imaging, on the Zeiss LSM880 microscope, the cells were kept at 37 °C with 5% CO₂ using the Zeiss incubator.

Data Analysis and interpretation

Fluorescence Lifetime Imaging and phasor analysis (FLIM phasor plot): The FLIM phasor plot was obtained using equations (1) and (2), as described previously.^{22, 31} Briefly, the fluorescence decay, $I(t)$, was acquired at each pixel of an image, and then, the G and S coordinates were calculated. The G and S coordinate for each pixel was plotted in a polar plot (referred to as phasor plot or phasor space) as follows:

$$\text{x-coordinate} = G_{(t)} = \frac{\int_0^T I(t) \cos(\omega t) dt}{\int_0^T I(t) dt} \quad (1)$$

$$\text{y-coordinate} = S_{(t)} = \frac{\int_0^T I(t) \sin(\omega t) dt}{\int_0^T I(t) dt} \quad (2)$$

where, ω is the angular modulation frequency (equal to $2\pi f$, and f is the repetition frequency of the laser) and T is the period of the laser frequency. Phasors follow the rules of vector addition, orthogonality, and reciprocity, i.e., pixels that contain a linear combination of two independent fluorescent species will appear on the line joining the two independent emissions in the phasor plot. A detailed description and discussion can be found elsewhere.³²

Hyperspectral Imaging and phasor analysis (Spectral phasor plot): The fluorescence spectrum, at each pixel, in a spectral image was transformed into phasor coordinates, ($G(\lambda)$) and ($S(\lambda)$), as described in equations 3 and 4, respectively.¹⁶ The $G(\lambda)$ and $S(\lambda)$ coordinates are the real and imaginary components of the Fourier transformation. $I(\lambda)$ represents the intensity at every wavelength (channel in our case), n is the number of the harmonic, and λ_i the initial wavelength. The x and y coordinates were plotted in the spectral phasor plot, as previously shown by Malacrida *et al.*¹⁶

$$\text{x coordinate} = G_{(\lambda)} = \frac{\int_{\lambda_{min}}^{\lambda_{max}} I(\lambda) \cos\left(\frac{2\pi n(\lambda - \lambda_i)}{\lambda_{max} - \lambda_{min}}\right) d\lambda}{\int_{\lambda_{min}}^{\lambda_{max}} I(\lambda) d\lambda} \quad (3)$$

$$y \text{ coordinate} = S(\lambda) = \frac{\int_{\lambda_{min}}^{\lambda_{max}} I(\lambda) \sin\left(\frac{2\pi n(\lambda - \lambda_i)}{\lambda_{max} - \lambda_{min}}\right) d\lambda}{\int_{\lambda_{min}}^{\lambda_{max}} I(\lambda) d\lambda} \quad (4)$$

The position for every pixel in the spectral phasor plot can be defined by the phase angle and the modulus (M), given the coordinates G and S (for both spectral or lifetime), in the following way:

$$\theta = \arctan (S/G) \quad (5)$$

$$M = \sqrt{S_2 + G_2} \quad (6)$$

The angular position in the spectral phasor plot relates to the center of mass of the emission spectrum, and the modulus depends on the spectrum's full width at half maximum (FWHM). For instance, if the spectrum becomes broad, the phasor cluster should move closer to the center (modulation is decreasing). On the other hand, for the red-shifted spectra, the position of the phasor will move counter-clockwise towards the increasing angles from position (1,0), i.e., the phase angle will increase. The spectral phasors share all the vector properties with the lifetime phasors. Moreover, an in-depth description of the properties of the spectral phasor plots is given in the reference.¹⁶

3. RESULTS and DISCUSSION

Cuvette spectroscopic characterization

The absorption and emission spectra of CAPRYDAA and LAURDAN in solvents with increasing polarity were compared. The absorption spectrum of CAPRYDAA (Fig. 2A) was found to undergo a shift of around 100 nm towards the red region as compared with the LAURDAN probe (Fig. 2D). In homogeneous media, the absorption spectra and molar absorptivity at maxima absorption wavelengths, for both CAPRYDAA and LAURDAN, showed a fair dependency on the solvent polarity. For the LAURDAN probe, the molar absorptivity, at the absorption maxima at a longer wavelength (around 20,000 M⁻¹cm⁻¹), was twice to that of CAPRYDAA's molar absorptivity, which was determined to be around 9,000 M⁻¹ cm⁻¹ (Fig. 2A and 2D).

Further, clear bathochromic shifts were observed for CAPRYDAA (Figs 2B and 2C) and LAURDAN (Figs. 2E and 2F). To compare the spectral shift (Stokes shift) for CAPRYDAA and LAURDAN, the Lippert-Mataga equation was used:^{33, 34}

$$\nu_A - \nu_F = \frac{2\Delta f (\mu_E - \mu_G)^2}{hc a^3}, \Delta f = \frac{\epsilon - 1}{2\epsilon + 1} - \frac{n^2 - 1}{2n^2 - 1}$$

In this equation, μ_E and μ_G corresponds to the dipole moments of the probe in the excited and ground state, respectively; a is a radius of an assumed spherical Onsager cavity, n^2 is the

square of the refractive index, and ϵ is the dielectric permittivity of the solvent. Figure 3 shows the experimental data and fitting to the Lippert-Mataga equation. The trend observed for both probes was the same (similar slopes were observed in Fig. 3). Assuming a larger volume for the cavity of CAPRYDAA, these data indicate that the charge transfer dipole of CAPRYDAA must be smaller than that of LAURDAN.

Additionally, it must be stated that both absorption and emission properties as well as their dependency on the solvent for CAPRYDAA were found to be similar to that reported for Anthradan by Lu *et al.*²⁸

Several CAPRYDAA photophysical parameters related to the first singlet excited state in homogeneous media were measured (Table 1). The singlet state emission decay of CAPRYDAA, in homogeneous media with different polarities, can be fitted to the monoexponential decays, with lifetime values (τ) between 5 and 8 ns (Table 1). Further, the lifetime values for LAURDAN, i.e., between 4 and 8 ns, were found to be similar to the values reported in the literature.³⁵ However, the fluorescence quantum yield (Φ_F) for CAPRYDAA decreased as the polarity of the solvent medium increased (Table 1).

Additionally, the effect of temperature in the CAPRYDAA emission from DPPC SUVs is shown in the supplementary material (Fig. S9).

Microscope intensity measurements

Two-photon excitation characterization—Generally, the measurements of the LAURDAN probe in a confocal microscope require two-photon excitation in order to avoid severe photobleaching that occurs with one-photon excitation. Thus, the first comparison for the newly synthesized CAPRYDAA and LAURDAN was the two-photon cross-sectional excitation spectra (Fig.4). For CAPRYDAA, the optimal excitation by two-photon excitation was determined to be around 900 nm, while the LAURDAN excitation maximum was found around 780 nm.

Fluorescence Lifetime characterization (FLIM-phasor analysis)—The second comparison was the dye response to the physical order of the lipid bilayer. The CAPRYDAA partition in water was estimated by the partition constant measurements in a water/dodecane mixture. $A_{LogP} > 4$ was experimentally obtained, and hence, it can be considered that there is no partition of CAPRYDAA in the solvent. Further, the fluorescence lifetime measurements in MLVs (multilamellar vesicles) of lipids in different phases were used to evaluate the performance of CAPRYDAA (Fig. 5). Three different sets of MLVs were prepared for this comparison, and the measurements were carried out at 37 °C: MLVs of DPPC (gel phase), DOPC (fluid phase), and DOPC:Cholesterol (1:1, molar) were used as a liquid order model. The vesicles were labeled with either CAPRYDAA or LAURDAN, and the lifetime measurements were performed by collecting the emission in two different channels: blue (channel-1) and green (channel-2) wavelength range of the dyes. Using this strategy, it was possible to isolate the polarity (blue region) from the dipolar relaxation effect of the dye in the excited state (green region).²² Additionally, the fluorescence lifetime measurements were carried out in a microscope, using two-photon excitation at 780 nm for

LAURDAN, and 900 nm for CAPRYDAA. For CAPRYDAA, one-photon excitation was also used since no significant photobleaching was observed.

Channel-1 (Fig. 5) gives information about the polarity of the media surrounding the dye molecule, and therefore the physical state of the membrane. As expected, the L_d phase showed a shorter lifetime (indicated by the red cursor in Fig. 5 A, B, and C, corresponding to channel-1). This result was consistent for both the dyes using one and/or 2-photon excitation. However, for L_β and L_o phases, the FLIM-phasor analysis showed some differences between CAPRYDAA and LAURDAN. The CAPRYDAA probe showed differences in the lifetime values when one or two-photon excitation was used (Fig. 5A and 5B). When one photon excitation was used, the lifetime values of CAPRYDAA were different, namely being shorter in the L_β phase than in the L_o phase (Fig. 5A); however, the two-photon excitation cannot differentiate between these two phases (Fig.5B, channel-1). On the other hand, for LAURDAN, the two-photon excitation led to a longer lifetime, which is presented by the L_β phase (Fig. 5C).

The lifetime data acquired in channel-2 (Fig 5A, 5B, and 5C; the bottom panel) gives information regarding the dipolar relaxation, and in the phasor representations shown in Figure 5, this phenomenon is illustrated by the position of the phasors outside the universal circle. Generally, a position outside the universal circle is associated with a reaction in the excited state, and in this case, it is due to the dipolar relaxation of the water molecules around the probe. The solvent dipolar relaxation takes time, and this duration is responsible for the increase in the phase angle, and hence, the position outside the phasor plot is observed. For CAPRYDAA, the dipolar relaxation was smaller for one-photon as compared with the two-photon excitation. In Table 2, the lifetime for CAPRYDAA and LAURDAN using phase and demodulation analysis is shown. The difference between the tau phase (τ^P) and tau modulation (τ^M) values in channel-1 is indicative of a complex decay. On the other hand, the larger values for the tau phase over the tau modulation in channel-2 were found to be related to the occurrence of a reaction in the excited state. For instance, delays in the phase were present when the dipolar relaxation of the solvent molecules occurred around the probe in the excited state.

Spectral shift comparison—The spectral shift of the probes in different lipid environments was analyzed using the spectral phasor approach^{16, 36} (Figure 6). The emission spectrum of CAPRYDAA was found to be similar, whether one-photon or two-photon excitation was used, as shown in the spectral phasor plots (Fig.6A and 6B). The spectral phasors for CAPRYDAA in the L_d phase showed the largest phase and smallest modulation, as evidenced by the more red-shifted spectra and higher emission heterogeneity (broader spectrum) (Fig.6A and 6B). Further, the blue-shifted spectra were found for CAPRYDAA in the membranes in the L_o phase. Finally, the gel phase (L_β) had similar modulation as L_o but with an increased phase angle, indicative of a blue-shifted spectrum. For LAURDAN, a larger spectral shift was obtained, but the most blue-shifted spectrum was identified for DPPC (gel phase, see Fig. 6C). The liquid order membrane showed a slightly increased phase angle, but with a significant decrease in modulation (Fig. 6C).

Spatial resolution in model systems—To test the spatial resolution of CAPRYDAA under hyperspectral imaging, i.e., its ability to discriminate between the membrane phases, we prepared giant unilamellar vesicles (GUVs) containing L_o , L_d , or coexistence of the L_o/L_d phases (Fig. 7). Using the cursor selection on the spectral phasor plot (Fig. 7A and 7B), we were able to segment the liquid order from the liquid disorder membrane either in the single lipid phases or in the coexistence of L_o/L_d phases.

Spatial resolution in live cells—After the GUV experiments, we moved to *in vivo* cell imaging. We used NIH-3T3 cell, and the CAPRYDAA fluorescence was acquired by using the lifetime and hyperspectral imaging (Fig. 8 and 9, respectively). To evaluate the dye response in different subcellular environments, we used segmentation by intensity and hand drawn masking. The lipid droplets were assigned to the high-intensity pixels due to the accumulation of the dye in the cell interiors and significant increment in the quantum yield (Fig. 8A and 8B). The spectral shift was color coded using the three-cursor selection as described previously by Ranjit *et al.*³⁷ (Fig. 8E and 8H). Then, the interior plus plasma membrane or the plasma membranes, isolated by the hand-drawn masks, were analyzed (Fig. 8C, 8G, 8D, and 8H). The histogram of the spectral shift was plotted as a fraction of the green cursor, representing the fraction of the L_o membrane (Fig. 8J). Using this approach, we can quantitatively evaluate the differences within the cellular membranes, since the lipid droplets correspond to the bluer environment. At this point, it is important to mention the biochemical and structural differences between the membrane and the lipid droplets. Further, concerning the solvent polarity, the dye present in the membrane senses the polarity at the phospholipid/water interphase, but in the lipid droplet, the dye will report from the homogenous apolar environment of the core. This capability to sense the two environments has also been demonstrated in other dyes, like push-pull fluorenes²⁵ and pyrenes.²⁷ CAPRYDAA efficiently discriminates between the plasma membrane and the intracellular membranes. In Figure S10 in the supplementary material, we present the “classical” generalized polarization⁵ ($GP = \text{Ch}_1 - \text{Ch}_2 / \text{Ch}_1 + \text{Ch}_2$) from the hyperspectral images using the two bands centered at 536 nm (channel-1) and 614 nm (channel-2).

Using the two FLIM channels, we can independently study the polarity and the dipolar relaxation of CAPRYDAA (Fig. 9). Additionally, the “classical” FLIM image generated by the tau phase histogram is shown in Figure S11 of the supplementary material. In the blue channel, we can see a trajectory associated with different fractions of L_o and L_d membranes in the cell (Fig. 9A, 9B, and 9C). The dipolar relaxation of water around the dye at the membrane interphase was assessed using the green FLIM channel (Fig. 9D, 9E, and 9F). An increased dipolar relaxation was found at the intracellular interior, while the plasma membrane presented lower dipolar relaxation when compared with the intracellular membranes (Figure 9F).

4. FINAL REMARKS

Biological membranes are a supramolecular assembly, classically associated with the subcellular compartmentalization; however, membranes are sensitive to the thermodynamic considerations, and thus, their physical order and dynamics can be modified. An important methodology to study the membrane dynamics and organization is the combination of

fluorescent dyes with microscopy. In this study, environment-sensitive probes were designed with the final goal of understanding the macromolecular fluctuations and conformation. LAURDAN has been widely described as a probe located at the membrane interphase and due to the charge transfer upon excitation, the increase in the dipole moment was used to test the environment. Weber has originally described the fundamentals between the polarity and the dipolar relaxation. The polarity was modified due to the apparent dielectric constant at the membrane interphase, while the dipolar relaxation was found to be related to the re-orientation of the solvent dipoles around the dye during the excited-state lifetime. LAURDAN is perhaps one of the most widely used membrane dyes in the literature, and up until now, it is the probe with the best-understood photophysics behavior. An important limitation of the use of LAURDAN is the need for two-photon excitation with microscopy in addition to the restriction to the blue spectral emission. Even upon using one-photon excitation of 405 nm, it is important to consider that such red excitation selects a subpopulation of LAURDAN's ground state, and thus the UV illumination may be detrimental to live cells. Therefore, novel probes with more red-shifted excitation spectra and capabilities for either one- or two-photon excitation will prove to be very useful in overcoming some of these drawbacks.

CAPRYDAA was designed based on the LAURDAN logic and considering the availability of new spectroscopy and image analysis tools. The rationale for the design of CAPRYDAA was related to the spectral-shifted absorption of anthracene with respect to naphthalene, but keeping the electron donor and acceptor groups as a key point in the photophysics of the dye.

This new probe demonstrated spectral shifts similar to LAURDAN with increasing solvent polarity but was found to undergo red-shift, making it easier to excite by the single-photon methods. This point must be emphasized, considering the limitation on the use of 405 nm for the LAURDAN excitation. Nowadays, some variants of LAURDAN, like C-LAURDAN or M-LAURDAN, claim better photostability than LAURDAN under UV excitation. Nevertheless, the major limitation of this kind of excitation is the deleterious effect on cells and the preferential ground state selection due to the red-edge excitation. CAPRYDAA was also excitable by two-photon excitation at 900 nm, which facilitates the combination with other fluorophores showing blue or red excitation spectra. This characteristic can be very useful when exploiting the linear combination properties of the phasor plots since the dye can be used in combination with the red fluorescent proteins to track particular subcellular structures.

CAPRYDAA was also sensitive to the dipolar relaxation; this property of LAURDAN is unique and is related to the delay in the emission due to the reorganization of the water molecules around the dye in the excited state. Water has a rotational time of the order of the probe lifetime, and for instance, any change in the number of water molecules or their dynamics can be sensed as a dipolar relaxation. The physical state of the membrane is responsible for the final number of water molecules, and thus, the dipolar relaxation can also be related to the membrane dynamics and organization. Interestingly, CAPRYDAA can be studied either by spectral or lifetime shifts, showing a tight correlation between both. The use of the phasor plots (for lifetime or spectra) enhances the bathochromic properties of

CAPRYDAA in the synthetic lipid vesicles. In GUVs, it was possible to identify the liquid/liquid phase separation and the characteristic domain formation in a ternary lipid mixture, which is the representative of lipid rafts. Finally, when CAPRYDAA was tested *in vivo* with live cells, it was possible to isolate the organization of the subcellular membrane by the spectral phasor analysis and image segmentation. This spectral shift is very useful in addressing the membrane dynamics in the *in vivo* experiments, as well as the dynamic range showed by the phasor plot can be applied to different kinds of experiments. With the FLIM data, a map of the polarity or the dipolar relaxation in the cell can be generated, which is valuable to understand the supramolecular heterogeneity of live cells. These two properties (polarity and dipolar relaxation) provide complementary information, in which the first one was related to the apparent dielectric constant at the membrane interphase and the second one to the relaxation of water molecules around the probe during its excited-state lifetime. It is interesting to discuss the difference in the dipolar relaxation observed by LAURDAN and CAPRYDAA, namely, it seems that LAURDAN has a larger dipolar relaxation shift than CAPRYDAA. However, it is important to consider the difference of the laser sources (one or two-photon), and another important point is the actual position of each dye in the membrane. It is known that LAURDAN is located right behind the carbonyl group at the membrane interphase. On the other hand, it looks as if CAPRYDAA was positioned a little deeper in the bilayer, which can also be explained by the smaller phase-shift observed in the green channel. An important aspect of the probe to highlight is its lack of distribution to the water molecules during the water/dodecane experiment, which besides its low emission from the water, is fundamental for studying the membranes avoiding an emission component from the bulk phase. This characteristic is shared with LAURDAN and is an important advantage of the probe over other soluble LAURDAN versions, which always present a water component.

Finally, CAPRYDAA shows excellent labeling in the *in vivo* experiments, where a hundred percent of the cells were strongly stained after 30 minutes of incubation, with the staining being homogeneous from cell to cell and inside the cell (data not shown). This point is an important property of CAPRYDAA to highlight, considering there is no preferential affinity for different regions in the cell that improves their uses for full-cell experiments. Besides, considering one of the main applications for CAPRYDAA in the *in cellulo* and *in vivo* experiments, easy labeling is one of the fundamental use.

To summarize, we introduced a new molecular probe for membrane dynamics and supramolecular organization with an excellent performance, which was comparable with the “gold standard” in the field, i.e., LAURDAN.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgment

We would like to dedicate this paper to the memory of Dr. Vicente Castro-Castillo who passed away during the revision stage of this manuscript. Vicente was fundamental to the development of CAPRYDAA and he was a wonderful scientist and person. He will be greatly missed by all who knew him. The authors would like to thanks to MilkaStacic for the cells preparation and Prof. David Jameson for his valuable suggestion during the manuscript

preparation. This work was partially supported by grants Grants NIH P41-GM103540 and NIH P50-GM076516. LM is supported for the Universidad de la República-Uruguay as a full time professor.

Bibliography

1. Bouvrais H, Pott T, Bagatolli LA, Ipsen JH and Meleard P, *Bba-Biomembranes*, 2010, 1798, 1333–1337. [PubMed: 20398624]
2. Sezgin E, Levental I, Mayor S and Eggeling C, *Nat Rev Mol Cell Bio*, 2017, 18, 361–374. [PubMed: 28356571]
3. Sklar LA, Hudson BS and Simoni RD, *Biochemistry-U S*, 1977, 16, 819–828.
4. Weber G and Farris FJ, *Biochem*, 1979, 18, 3075–3078. [PubMed: 465454]
5. Parasassi T, De Stasio G, Ravagnan G, Rusch R, M and Gratton E, *Biophys. J*, 1991, 60, 179–189. [PubMed: 1883937]
6. Parasassi T and Gratton E, *J Fluoresc*, 1995, 5, 59–69. [PubMed: 24226612]
7. Bagatolli LA, Maggio B, Aguilar F, Sotomayor CP and Fidelio GD, *BBA-Biomembranes*, 1997, 1325, 80–90. [PubMed: 9106485]
8. Bagatolli LA, Gratton E and Fidelio GD, *Biophys. J*, 1998, 75, 331–341. [PubMed: 9649390]
9. E. Lissi A., Abuin E, B., Rubio M, A., Ceron A and Langmuir, 2000, 16, 178–181.
10. Harris F, S. Smith M, J. Bell K and D, *J. Biol. Chem*, 2001, 276, 22722–22731. [PubMed: 11294853]
11. Viard M, Gally J, Vincent M and Paternostre M, *Biophys. J*, 2001, 80, 347–359. [PubMed: 11159407]
12. Sanchez SA, Bagatolli LA, Gratton E and Hazlett TL, *Biophys. J*, 2002, 82, 2232–2243. [PubMed: 11916878]
13. Tricerri M, J. Toledo A, S. Sanchez D, T. Hazlett A, E. Gratton L, Jonas A, Garda H and A, *J. Lipid Res*, 2005, 46, 669–678. [PubMed: 15654128]
14. Bagatolli LA, *BBA-Biomembranes*, 2006, 1758, 1541–1556. [PubMed: 16854370]
15. De Vequi-Suplicy C, C. Benatti C, M. Lamy R and T, *J Fluoresc*, 2006, 16, 431–439. [PubMed: 16791507]
16. Malacrida L, Gratton E, Jameson DM, *Methods Appl Fluores*, 2015, 3.
17. Malacrida L, Astrada S, Briva A, Bollati-Fogolin M, Gratton E, Bagatolli L and A, *BBA-Biomembranes*, 2016, 1858, 2625–2635. [PubMed: 27480804]
18. Strohmeier A, Forst G, Tauber P and Schubert R, *Biophys. J*, 2016, 111, 1714–1723. [PubMed: 27760358]
19. Bagatolli LA and Gratton E, *Biophys. J*, 1999, 77, 2090–2101. [PubMed: 10512829]
20. Sanchez SA, Tricerri MA, Ossato G and Gratton E, *Bba-Biomembranes*, 2010, 1798, 1399–1408. [PubMed: 20347719]
21. Rodriguez-Agudo D, Malacrida L, Kakiyama G, Sparrer T, Fortes C, Maceyka M, Subler MA, Windle JJ, Gratton E, Pandak WM and Gil G, *J. Lipid Res*, 2019, 60, 1087–1098. [PubMed: 31015253]
22. Malacrida L, Jameson DM and Gratton E, *Sci Rep-Uk*, 2017, 7, 9215.
23. Malacrida L and Gratton E, *Free Rad. Biol. Med*, 2018, 128, 144–156. [PubMed: 29885356]
24. Kucherak O, P. Didier A, Mely Y, Klymchenko A and S, *J Phys Chem Lett*, 2010, 1, 616–620.
25. Shaya J, Collot M, Benailly F, Mahmoud N, Mely Y, Michel B, A. Klymchenko Y, S and Burger A, *Acs Chem Biol*, 2017, 12, 3022–3030. [PubMed: 29053920]
26. Niko Y, Kawauchi S and Konishi G, *Chem. Eur. J*, 2013, 19, 9760–9765. [PubMed: 23744761]
27. Niko Y, Didier P, Mely Y, Konishi G, Klymchenko A and S, *Sci Rep-Uk*, 2016, 6.
28. Lu Z, S. Lord K, H. Wang J, Moerner W, R. Twieg E and J, *J Org Chem*, 2006, 71, 9651–9657. [PubMed: 17168582]
29. Moon H, Jun Y, D. Kim W, Ryu H, T. Wang G, Kim K, Y. Huh H, Jung J, Ahn K and H, *Chem-Asian J*, 2016, 11, 2518–2523. [PubMed: 27535006]
30. Pott T, Bouvrais H and Meleard P, *Chem. Phys. Lipids*, 2008, 154, 115–119. [PubMed: 18405664]

31. Jameson DM, Gratton E and Hall RD, *Appl Spectrosc Rev*, 1984, 20, 55–106.
32. Ranjit S, Malacrida L, Jameson DM and Grafton E, *Nat Protoc*, 2018, 13.
33. Lippert E, *Naturforsch A Z Astrophys Phys. Physikalisch Chem*, 1955, 10, 541–545.
34. Mataga N, Kaifu Y and Koizumi M, *Bull Chem. Soc Jpn*, 1956, 29, 465–470.
35. Ma Y, Benda A, Kwiatek J, Owen DM and Gaus K, *Biophys. J*, 2018, 115, 1498–1508. [PubMed: 30269886]
36. Sena F, Sotelo-Silveira M, Astrada S, Botella M, L. Malacrida A and Borsani O, *Plant Physiol. Biochem*, 2017, 119, 224–231. [PubMed: 28910707]
37. Ranjit S, Dvornikov A, Dobrinskikh E, Wang XX, Luo YH, Levi M and Gratton E, *Biomed Opt Express*, 2017, 8, 3143–3154. [PubMed: 28717559]

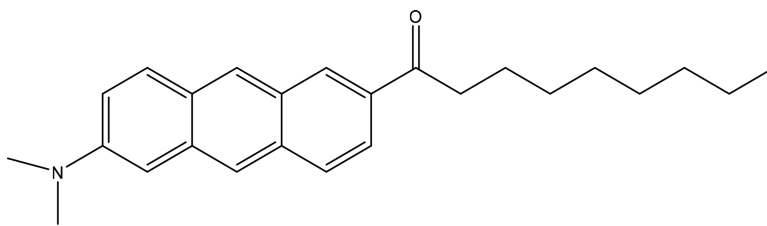


Figure 1.
The molecular structure of CAPRYDAA.

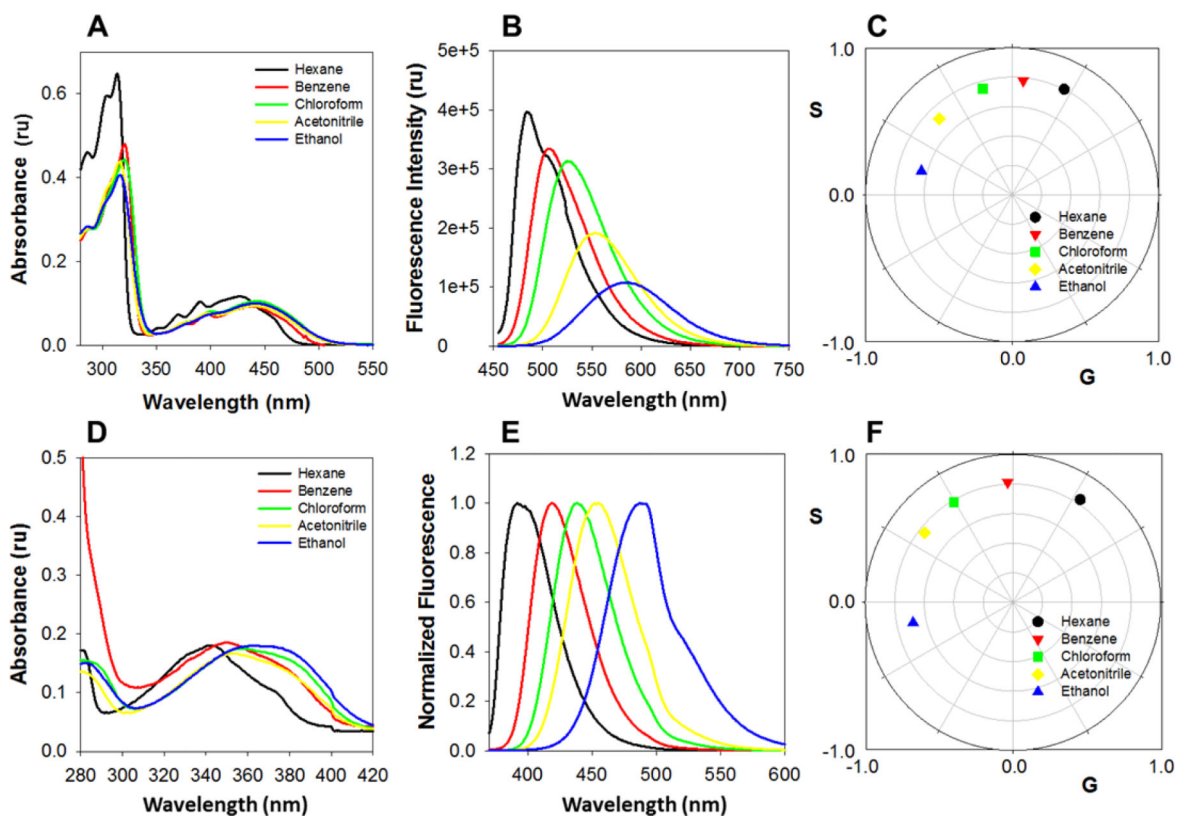
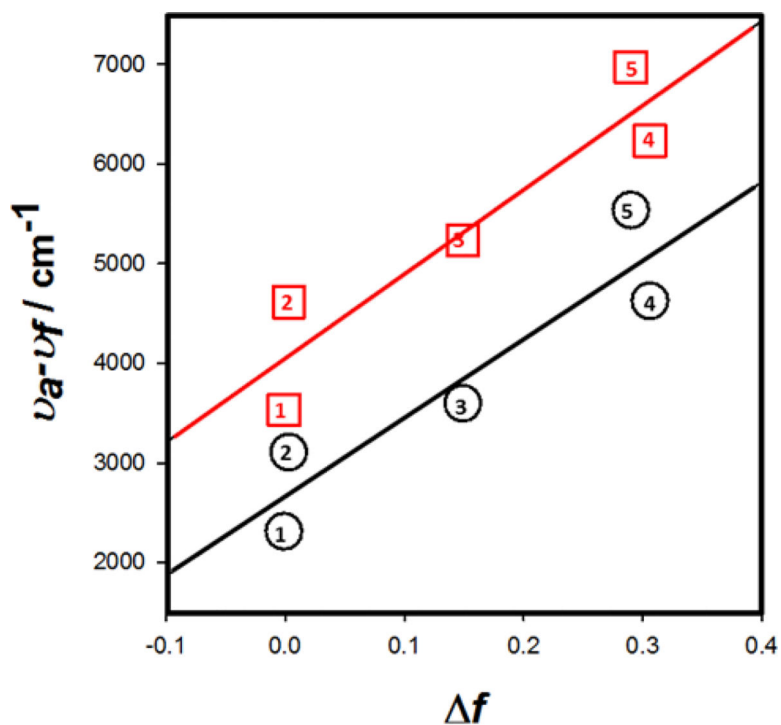


Figure 2: Spectroscopic characterization of CAPRYDAA and LAURDAN.

A) The absorption spectra of CAPRYDAA in solvents with increasing polarity (hexane, benzene, chloroform, acetonitrile, and ethanol). B) The emission spectra for CAPRYDAA in increasing solvent polarity. The excitation wavelength used was 450 nm. C) The spectral phasor transformation for the emission spectra of CAPRYDAA shown in B. D) The absorption spectra of LAURDAN in solvent with increasing polarity (hexane, benzene, chloroform, acetonitrile, and ethanol). E) The normalized emission fluorescence spectra for LAURDAN in increasing solvent polarity. The excitation wavelength used was 360 nm. F) The spectral phasor transformation for the LAURDAN emission spectra shown in E.



Fitting parameters for Lippert-Mataga plot

Equation $y = a + b \cdot x$

Probe	b	a	R ²
CAPRYDAA	7910.5	2654.1	0.813
LAURDAN	8464.6	4053.2	0.828

1-Hexane, 2-Benzene, 3-Chloroform, 4-ACN, 5-EtOH

Figure 3: The Lippert-Mataga plot for CAPRYDAA (red squares) and LAURDAN (black circles) as a function of solvent polarity.

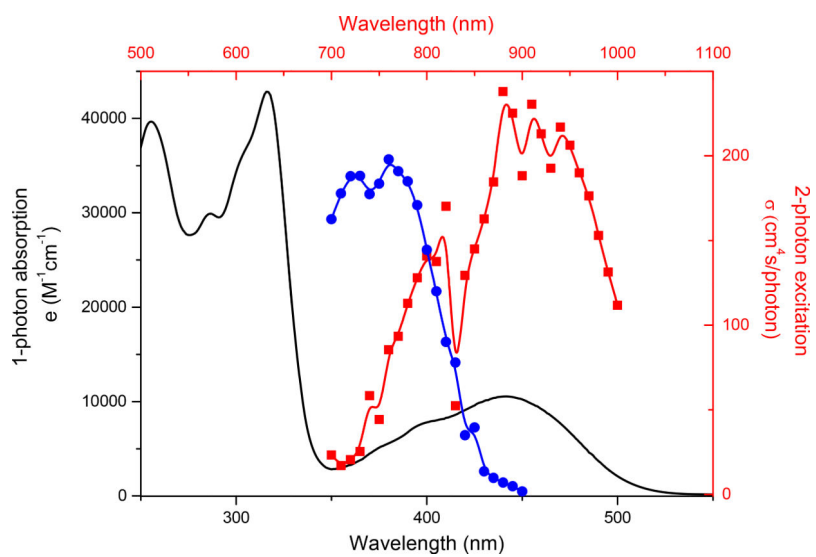


Figure 4. Two-photon excitation spectra for CAPRYDAA and LAURDAN: The black line shows the CAPRYDAA absorption spectrum in DMSO. The red line and square boxes show the two-photon excitation spectra for CAPRYDAA in DMSO. The LAURDAN two-photon excitation spectrum in DMSO is shown by the blue trace and circles.

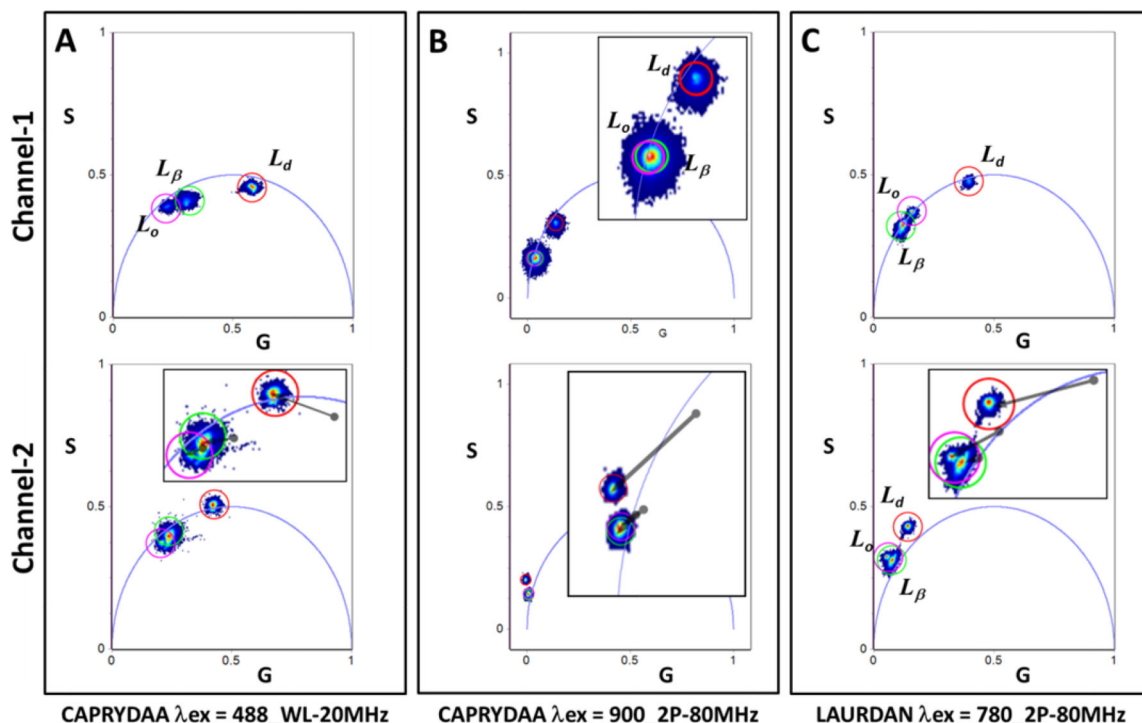


Figure 5: CAPRYDAA and LAURDAN lifetime comparison in multilamellar vesicles by FLIM-Phasor plot.

A) The FLIM-phasor plot for CAPRYDAA in MLVs with L_o , L_d , and L_β phases using 20 MHz pulsed supercontinuum laser (488 nm excitation) and collecting the emission decay in two channels: channel-1 (polarity) and channel-2 (dipolar relaxation). The arrows in the inset illustrated the shift of the phasor position from channel-1 to channel-2. B) The phasor plot distribution for CAPRYDAA in MLVs with L_o , L_d , L_β phases using 900 nm excitation at 80 MHz. C) The phasor plot distribution for LAURDAN in MLVs with L_o , L_d , and L_β phases using 780 nm excitation at 80 MHz. L_o was DPPC:Cholesterol 1/1 molar, L_d was DOPC, and L_β was DPPC.

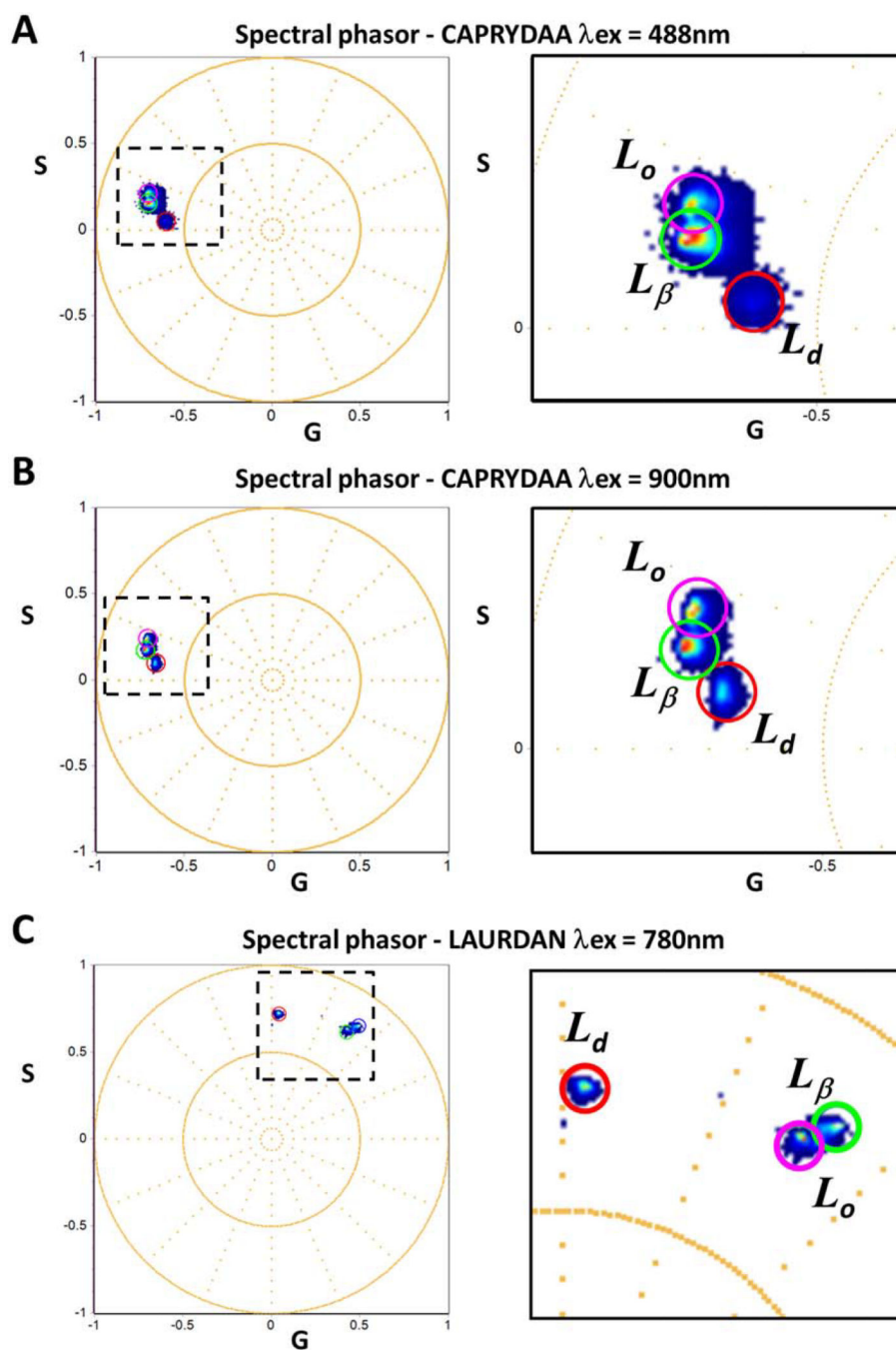


Figure 6: Spectral phasor characterization for single and two-photon excitation of CAPRYDAA versus LAURDAN in multilamellar vesicles. (A) Spectral phasor characterization of CAPRYDAA in MLVs with different lipid phases (L_b -DPPC green cursor; L_o -DPPC:cholesterol 1/1 molar pink cursor; L_d -DOPC red cursor) using single photon excitation (488 nm). (B) Spectral phasor characterization of CAPRYDAA in MLVs with different lipid phases using twophoton excitation (900 nm). (C) Spectral phasor of LAURDAN in MLVs with different

lipid phases using two-photon excitation (780 nm). All the experiments were done at 37 1C. At the right of each figure is an enlargement of the ROI represented as the dashed box.

Author Manuscript

Author Manuscript

Author Manuscript

Author Manuscript

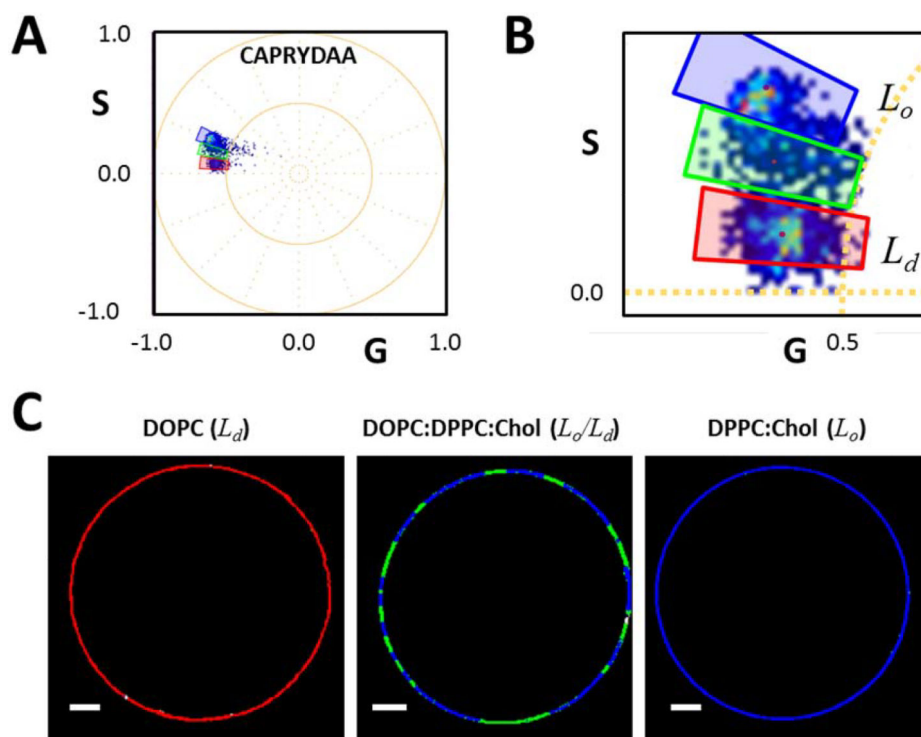


Figure 7: CAPRYDAA fluorescence in GUVs with different phases.

A) The spectral phasor plot for CAPRYDAA in L_α , L_α/L_β and L_β membranes. B) Zoom of A. C) The representative images of GUVs with L_β , L_α/L_β and L_α phases, respectively.

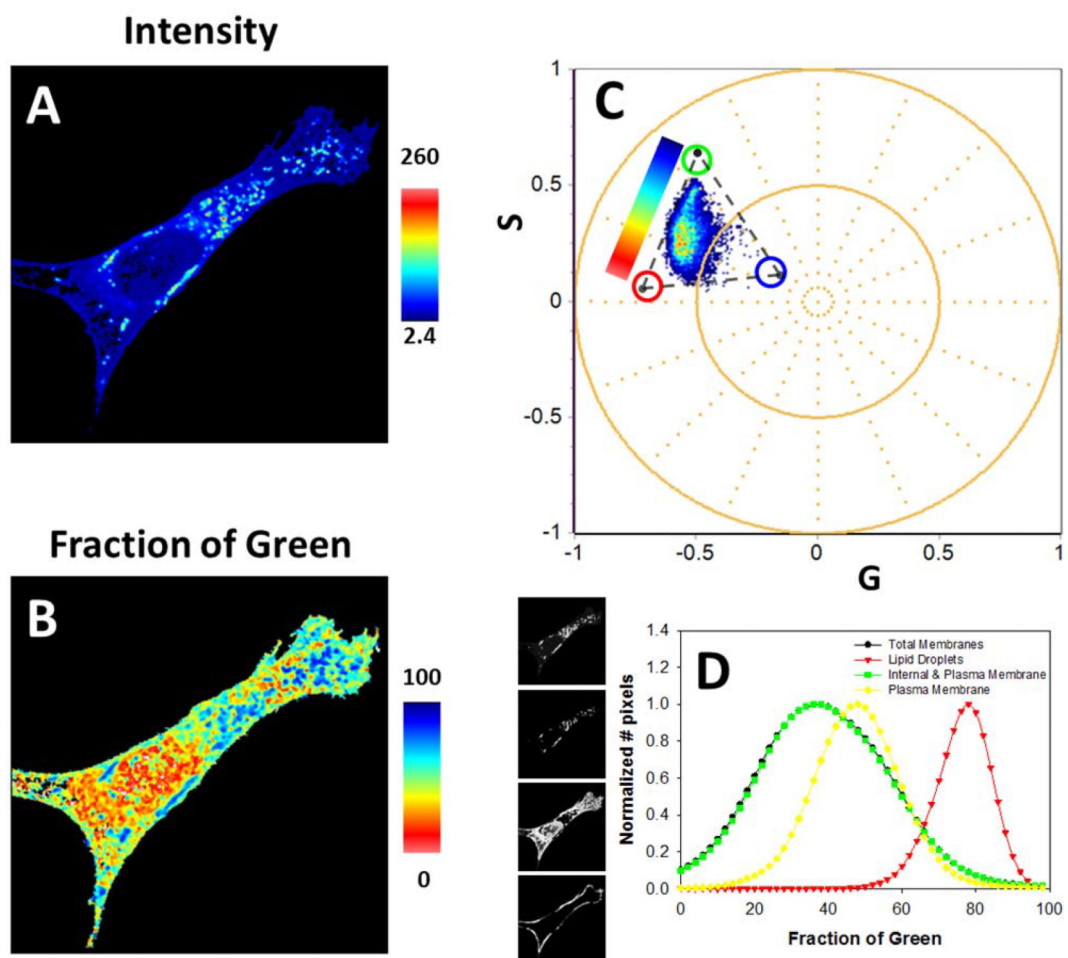


Figure 8:

Hyperspectral study of CAPRYDAA fluorescence: (A) intensity image for CAPRYDAA fluorescence using two-photon excitation. (B) Pseudo-color image generated by the spectral phasor analysis of the CAPRYDAA spectrum shift represented as a fraction of green cursor in the spectral phasor plot of figure (C). The color scale is representative of the membrane order, where blue means ordered membrane and red means fluid membrane. (C) The spectral phasor plot of CAPRYDAA fluorescence. The cloud of pixels was colored using the scale shown between the red and green cursors. The red and green cursors are associated with the fluid to ordered fraction, respectively. (D) The histogram of CAPRYDAA spectral shift in different cellular membranes, obtained by masking the subcellular regions (inset figures at the left). The fraction of green is related to the fraction of membrane order.

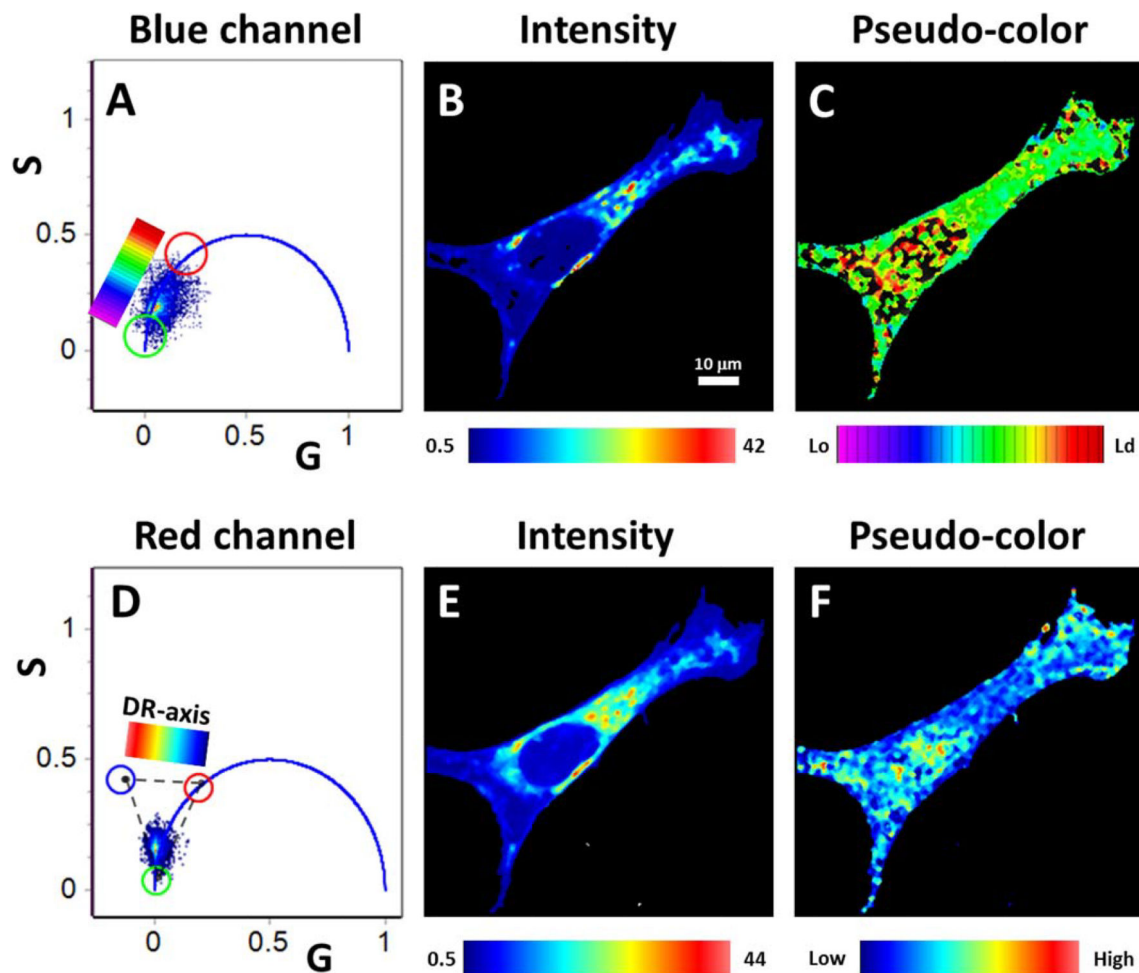
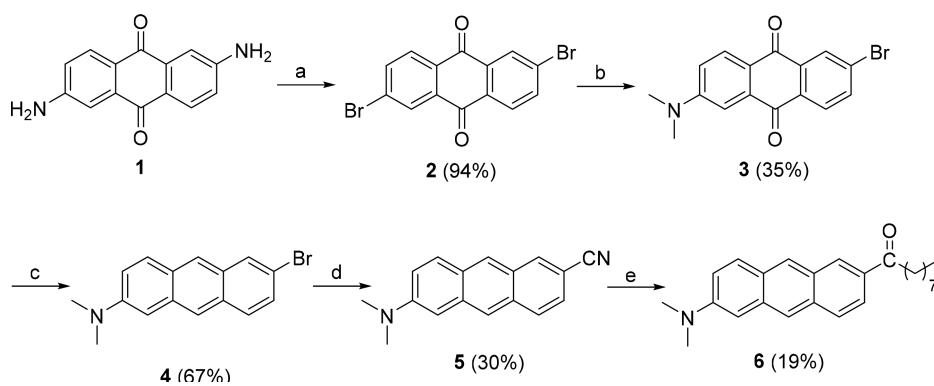


Figure 9: FLIM characterization of CAPRYDAA fluorescence in cells:

A, B, and D) The FLIM-phasor analysis of CAPRYDAA using a blue filter. The blue filter is used to characterize the polar part of the CAPRYDAA emission. Using the color scale shown in Figure A we generated Figure C. Red and green cursor represent the “pure” fluid and ordered phases, respectively. In between it is possible to see the cloud of pixels with different fraction of the two components. D, E, and F) The FLIM-phasor analysis of CAPRYDAA fluorescence using a red filter. The red filter is used to identify the occurrence of dipolar relaxation. The dipolar relaxation is a reaction in the excited-state and for instance should produce a delay in the phase, which pull out the universal circle the pixels that undergoes to dipolar relaxation, see figure D. To monitor the change in the dipolar relaxation, we use a three cursor analysis as previously described by Malacrida et al.²³ And the dipolar relaxation axis, the DR-axis was colored using the color scale shown in figure D. In the dipolar relaxation figure (F) red means higher dipolar relaxation. It is important to notice that almost all the pixels are outside the universal circle.

**Scheme 1.**

Reagents and conditions: (a) *tert*-butylnitrite, CuBr₂, CH₃CN, 90 °C, 24 h; (b) (1) CsF, DMSO, N₂, 170 °C, 5 h; (2) (CH₃)₂NH (ac) 40%, K₂CO₃, 70 °C, 17 h; (c) (1) NaBH₄, 2-propanol, reflux, 12 h; (2) HCl (ac) 37%; (d) CuCN, DMF, N₂, reflux 12 h; (e) CH₃(CH₂)₆CH₂MgBr, CuBr (cat.), THF, N₂, room temperature, 1 h.

Table 1:

Results summary for the photophysics properties of CAPRYDAA in solvents with different polarity.

Solvent	Φ_F	τ/ns	λ_{abs}/nm	λ_{em}/nm	$\epsilon/M^{-1}cm^{-1}$
Hexane	0.44	6.46	314 – 428	475	52589 (314) 9437 (428)
Benzene	0.51	7.84	320 – 438	507	50566 (320) 9816 (438)
CHCl ₃	0.44	7.68	321 – 443	527	42469 (321) 10113 (443)
ACN	0.29	7.56	316 – 441	554	46148 (316) 9924 (441)
EtOH	0.10 (MEOH)	5.29	316 – 443	587	42841 (316) 10552 (443)

Φ_F = Fluorescence Quantum Yield

τ = Fluorescence lifetime

λ_{abs} = absorbance wavelength

λ_{em} = emission fluorescence wavelength

ϵ = extinction coefficient (value and absorbance wavelength)

Table 2:

Fluorescence Lifetime of CAPRYDAA and LAURDAN in MLVs.

Lipid Mixture	Channel-1		Channel-2	
	τ^P	τ^M	τ^P	τ^M
CAPRYDAA 900nm				
DPPC	7.07±0.05	11.33±0.05	34.00±2.405	13.67±0.05
POPC	4.37±0.12	5.50±0.00	----*	9.50±0.08
DPPC:Chol (1:1)	9.60±0.05	11.73±0.17	29.37±4.84	13.33±0.12
LAURDAN 780nm				
DPPC	5.27±0.05	5.37±0.05	8.10±0.22	5.93±0.12
POPC	2.37±0.05	2.47±0.05	5.93±0.05	3.87±0.05
DPPC:Chol (1:1)	5.23±0.05	5.43±0.12	8.07±0.17	5.93±0.12

 τ^P tau phase τ^M Tau modulation

n = 3 for each sample

* phase larger than 90 degree.