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Solution and Interface Aggregation States of *Crotalus atrox* Venom Phospholipase A₂ by Two-Photon Excitation Fluorescence Correlation Spectroscopy†

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ABSTRACT: The dimeric *Crotalus atrox* venom PLA₂ is part of the secreted phospholipase A₂ (PLA₂) enzyme family that interacts at the lipid—solution interface to hydrolyze the sn-2 acyl ester bond of phospholipids. We have employed fluorescence correlation spectroscopy (FCS) to study the monomer—dimer equilibrium of the *C. atrox* venom PLA₂ in solution, in the presence of urea, and in the presence of monomeric and micellar n-dodecylphosphocholine (C12-PN), a phosphatidylcholine analogue. Dilution experiments show that PLA₂ is an extremely tight dimer, *K₅* ≤ 0.01 nM, in solution. Urea was introduced to weaken the subunit’s association, and an estimate for the PLA₂ dimer dissociation constant in buffer was obtained by linear extrapolation. The derived dissociation constant was at least several orders of magnitude greater than that suggested from the dilution experiments, indicating a complex interaction between urea and the PLA₂ dimer. FCS data indicate that the PLA₂ dimer begins to dissociate at 10 mM C12-PN in 10 mM Ca²⁺ and at 5 mM C12-PN in 1 mM EDTA. The PLA₂ tryptophan fluorescence displayed spectral shifts and intensity changes upon interacting with C12-PN. On the basis of the FCS and tryptophan fluorescence results, we postulate an intermediate state where the two monomers are in loose interaction within a protein—lipid micelle. As the concentration of C12-PN was increased, complete dissociation of the dimer was observed, inferred from the doubling of the particle number, and the average diffusion constant decreased to approximately 60 μm²/s, consistent with PLA₂ associated with a C12-PN micelle. The presence of Ca²⁺ makes the micelle intermediate more stable, retarding the separation of the monomers in the micellar suspension. Our data clearly indicate that PLA₂, though a strong dimer in the absence of lipids, is dissociated by micellar C12-PN and supports the monomer hypothesis for PLA₂ action.

Secreted phospholipase A₂’s (PLA₂) constitute a family of calcium-dependent enzymes which hydrolyze the sn-2 acyl ester bond of phospholipids. They may be isolated from the pancreas and snake or insect venoms, and some forms have now been cloned. Their primary structures are similar with 50–70% sequence homology, and their monomer molecular masses range from approximately 13 to 15 kDa. The secreted PLA₂’s are soluble enzymes that interact at the solution—lipid interface of micellar and bilayer phospholipids to hydrolyze phospholipid into lysophospholipid and free fatty acid.

The interactions of PLA₂ with lipids have been extensively studied over the last 30 years. The focus of these studies has been on the effect of lipid organization on enzymatic activity (1–3). The PLA₂ specific activity against monomeric phospholipids is poor but is greatly enhanced, from 10- to 100-fold, when the lipid is organized in micellar form (4). Under the appropriate conditions, PLA₂ interfacial activation in phospholipid bilayers can be observed as a delay period before the onset of active phospholipid hydrolysis (5, 6). Researchers exploring the nature of the lag phase events have implicated several factors which seem to be critical in this process, including the fluctuation of lipid microdomains (7–9), the extent of membrane curvature (10, 11), and the overall charge of the lipid membrane. Early in PLA₂ research, it was noted that some PLA₂s were dimeric in solution, which lead several groups to propose that interfacial activation of monomeric forms involves dimer formation (12–17). In fact, calcium, an essential cofactor for activity, has been proposed to favor protein dimerization for some PLA₂s (18–20). Considering the similarity in sequence and catalytic mechanism within this group of enzymes, it would appear that the monomer contains all of the necessary structure for full enzymatic activity. However, the existence of tight dimer PLA₂s such as the PLA₂ from *Crotalus atrox* venom begs the question as to the role that the dimer plays in PLA₂ function. Whether or not the dimer is the active form remains an unanswered question with sporadic support appearing in the literature (5, 13). Thus, our interest in the monomer—dimer equilibrium of the *C. atrox* PLA₂ stems from specific questions about the influence of the dimer on function, i.e.,

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1 Abbreviations: PLA₂, phospholipase A₂; FCS, fluorescence correlation spectroscopy; CMC, critical micelle concentration; C12-PN, n-dodecylphosphocholine; Fluo-PLA₂, fluorescein-labeled PLA₂; Laurdan, 6-dodecanoyl-2-(dimethylamino)naphthalene; C14-PN, n-tetradecylphosphocholine; C16-PN, n-hexadecylphosphocholine; EDTA, (ethylenedinitrilo)tetraacetic acid; GL, 3-D Gaussian—Lorentzian model.
the original active dimer (and half-the-sites) hypotheses presented in earlier works (15) or involvement with the lag/burst phase kinetics (5), but also from general questions on protein–protein interactions and protein interactions with a membrane interface.

Early studies on the C. atrox venom PLA2 quickly identified this enzyme as a strong dimer, similar to another dimeric PLA2 isolated from Crotalus adamanteus venom (21–23). The secreted PLA2 from C. atrox is a dimer when free in solution (20) with a reported $K_d$ in the nanomolar range (24). Using chromatographic profiles, Myatt et al. also studied the effect of Ca$^{2+}$ on the $K_d$ and they report a 5-fold decrease of the dissociation constant when the cofactor is present. Hydrolysis activity studies on the strong dimer PLA2S indicated that the active unit was likely a dimer with a dissociation constant of 0.6 nM in the presence of lipid substrate and Ca$^{2+}$ cofactor (21). Other researchers have questioned the general conclusions and began examining monomeric PLA2S and searching for ways to identify the active unit in the presence of substrate (18, 19, 25, 26).

A common experimental observation has been reported for PLA2S when they interact with lipid interfaces: changes in tryptophan emission (5, 6, 27, 28). In studies on micellar lipid interactions with PLA2S, at lipid concentrations below the CMC (critical micellar concentration), small decreases in the tryptophan emission may occur depending on the enzyme source and pH (29). At lipid concentrations above their CMC there is normally an increase in tryptophan emission accompanied by a blue shift of the emission maximum (29). This increase in tryptophan fluorescence has been attributed to direct lipid–protein interactions, conformational changes in the enzyme/lipid complex, and, the most controversial one, postulated changes in the aggregation state of the enzyme. Direct biophysical studies to determine the aggregation state of this enzyme when interacting with lipids in any of its conformations (micelles, bilayers, etc.) have been rare. Hydrolysis data by Jain and co-workers suggest that C. atrox venom PLA2 is a dimer in solution but dissociates at the lipid interface, but no physical evidence has been forthcoming. In contrast, a recent report by Ferreira et al. on immobilized PLA2 from C. atrox venom identified this enzyme as an active dimer (13).

Few techniques can be applied onto a membrane or micelle surface. One of the most commonly used fluorescence techniques in studying oligomer dissociation is fluorescence polarization (30–32). The ratio of the rotational correlation time of the dimer to the monomer is 2, due to the volume change, which allows studying monomer/dimer equilibrium in solution for small proteins. However, in the PLA2 case in which the dimer dissociates upon interaction with lipids, there is an additional change in mass due to the number of lipids that associate with the protein. Thus, fluorescence polarization cannot be used to study dimer dissociation under this condition.

An alternative fluorescence technique that can provide information about monomer/dimer equilibrium and, at the same time, determine the mass of the protein/lipid aggregates is fluorescence correlation spectroscopy (FCS). This technique allows one to count the number of fluorophore-labeled proteins in a particle aggregate using the fluctuation of the fluorescence intensity. The fluctuation amplitude is inversely proportional to the number of particles in the excitation volume for a single molecular species. In our case, a decrease in the fluctuation amplitude, $G(0)$, by a factor of 2 will indicate complete dissociation of the PLA2 (33). An additional advantage of the FCS techniques is the very high sensitivity permitting its use at very low protein concentrations (subnanomolar range), which is crucial for the present study.

In this work, we have determined an upper limit for the $K_d$ by dilution of the enzyme through at least 3 orders of magnitude in concentration, in the presence and absence of Ca$^{2+}$. We also studied the effect of urea on the dissociation of the enzyme. In the presence of urea the enzyme dissociates without being denatured (as reported by the CD signal at 220 nm). We also studied the interaction of PLA2 with lipids to answer specific questions as to the state of the enzyme (monomeric vs dimeric) below and above the CMC of the C12-PN lipid analogue. Since we were independently following the dissociation of the enzyme using FCS, we were also able to determine whether there is correlation between the changes in tryptophan emission with enzyme dissociation. We measured the number of enzyme particles when lipid monomers are present (below the CMC) and when micelles are already formed (above the CMC). The comparison of dissociation curves of the number of particles and of tryptophan emission at different lipid concentrations in the presence and absence of Ca$^{2+}$ allowed us to address the above questions and to propose a structural model for PLA2/lipid interactions. Most importantly, our data indicate that the PLA2 dimer dissociates into monomers given sufficient micellar lipid and suggest that under assay conditions and in the presence of substrate the monomer form is the active enzyme form.

Finally, this study shows that extending traditional fluorescence measurements to FCS can be a promising approach to study the interaction of protein in its natural biological environment (interacting with lipids). Here, FCS provides a direct measure of the dissociation process, while conventional fluorescence probes environmental factors affecting the chromophore. The combination of both techniques provides complementary information that allows us to formulate a more detailed model.

**MATERIALS AND METHODS**

*Sample Preparation.* The PLA2 from C. atrox venom (Miami Serpentarium, Punta Gorda, FL) was purified in our laboratory using standard methods (34). Fluorescein conjugates of PLA2 were prepared with fluorescein succinidyl ester (Molecular Probes, Eugene, OR) using standard methodology. Laurdan was purchased form Molecular Probes (Molecular Probes, Eugene, OR). The labeled fluorescein/enzyme ratio was calculated using an extinction coefficient for fluorescein of $\epsilon_{499} = 70,000$ M$^{-1}$ cm$^{-1}$ (35), and the protein concentration was determined using either Bio-Rad Protein Assay (Bio-Rad, CA) for the labeled enzyme or $\epsilon_{280} = 25,000$ M$^{-1}$ cm$^{-1}$ (34) for the enzyme without labeling. Enzymatic activity was measured in a pH-stat using a mixed micelle assay at pH = 8.0 (36). The labeling ratio 2 fluoresceins per protein and the enzyme conjugates retained >90% of the activity of unlabeled PLA2. The phospholipid analogue 1,2-dodecylphosphocholine was from Avanti Polar Lipids (Alabaster, AL). Urea and general chemicals were from Fisher Scientific (Fair Lawn, NY).


FCS Experiments. The instrumentation for two-photon fluctuation experiments is similar to that described by Berland et al. (37) with the following modifications: the experiments were carried out using a Zeiss Axiovert 135 TV microscope (Thornwood, NY) with a 63X Plan Apochromat-oil immersion objective (NA = 1.4). A mode-locked Ti: sapphire laser (Mira 900; Coherent, Palo Alto, CA) pumped by an intracavity doubled Nd:YVO4 vanadate laser (Verdi; Coherent Inc., Santa Clara, CA) was used as a two-photon excitation source. For all measurements, an excitation wavelength in the range from 770 to 780 nm was used, while the average power at the sample ranged from 7 to 2 mW. Photon counts were detected using an Avalanche photodiode detector (APD) (Model SPCM-AQ-151; EG&G). The output of the APD unit was directly connected to the data acquisition card. The photon counts were sampled at either 20 or 100 kHz. The recorded photon counts were later analyzed with programs written in PV-WAVE version 6.10 (Visual Numerics). Samples were mounted in a sample holder fabricated out of Delring (Illini Plastic, IL). A drilled hole in the center of the chamber was covered by a 1.5 mm standard microscope cover glass and used as the window for the microscope objective. There were no additional treatments applied to the objective. There were no additional treatments applied to the chamber. The chamber was covered by a 1.5 mm standard microscope cover glass and used as the window for the microscope objective. There were no additional treatments applied to the objective. There were no additional treatments applied to the objective.

(A) Dilution Experiments. In the dilution experiments, the sample holder was placed in the microscope, coupled with immersion oil, and kept there until the last sample was measured. Three sets of samples were measured: fluorescein, fluorescein–PLA2 (Fluo-PLA2) in 1 mM EDTA, and Fluo-PLA2 plus 10 mM CaCl2 in 50 mM Tris, pH 8.0. A fluorescein standard was used to calibrate the equipment. Each series of measurements started by the buffer alone, then the Fluo-PLA2 was diluted from a concentrated stock to the initial concentration of the set, and all the following dilutions were done in the same cuvette. Adsorption of molecules to the container walls is a concern when performing titration experiments at low concentrations. To ensure that adsorption did not influence our results, we performed the experiments in both directions: starting from either low or high PLA2 concentrations. The results were independent of the protocol, indicating that PLA2 adsorption did not adversely affect our studies.

(B) Dissociation with Urea. The samples, at different concentrations of urea, were prepared and incubated over-night. Protein concentration was kept constant at 0.1 μM. The sample holder was also kept in place until the last sample was measured. In this case after each sample was measured, the cuvette was washed with buffer several times until the fluorescence reached the background counts of the buffer. In this type of experiment we also used fluorescein at the same urea concentrations as a control.

(C) Lipid–PLA2 Interactions. Independent samples with different n-dodecylphosphocholine (C12-PN) concentrations in 50 mM Tris, pH 8 (with or without Ca2+), were prepared, and the fluorescently labeled enzyme was added. Protein concentration was kept constant at 0.14 μM. The chamber was also washed exhaustively with buffer before samples.

PLA2 Emission Spectra. Tryptophan emission spectra were recorded between 310 and 450 nm, with excitation at 295 nm on a Fluoromax-2 (JOBIN YUON-SPEX, Instruments S.A., Inc., Edison, NJ) connected to a water bath at 25 °C. Protein concentration was 0.14 μM, unless otherwise stated. Unlabeled protein was used to avoid any quenching of the signal due to energy transfer between the tryptophan and fluorescein. For each emission spectrum, two parameters were calculated: the center of mass and the total area of the emission band.

CD Spectra. Circular dichroism spectra were taken on a JASCO J-720 spectropolarimeter. Spectra were obtained for PLA2 under varying concentrations of urea at a protein concentration of 14 μM in Tris buffer, pH 8. Samples were incubated with denaturant overnight before the spectra were collected. Each spectrum was obtained at 25 °C in a 0.1 cm optical path cell. Wavelengths were scanned between 250 and 190 nm at 50 nm/min with a band-pass of 0.5 nm.

FCS Data Analysis. Experimental autocorrelation functions were fitted with an equation assuming a Gaussian–Lorentzian intensity profile, as described in a previous work which contains the explicit formulas for the point spread function and the definition of the beam waist used (37). The beam waist of the Gaussian–Lorentzian function depends on the instrument setup and must be calibrated each time the system is aligned. For this purpose a substance with a known concentration and diffusion coefficient (D) was used to calibrate the excitation volume. In this work, fluorescein (in 50 mM Tris, pH 8.0), with a reported diffusion constant of 330 μm²/s (33), was used as the standard. The recovered beam waist value of 0.37 μm was used to perform a global analysis with each set of data (dilution, urea, etc.). For a single species, the fluctuation amplitude, G(0), is related to the number of particles by

$$G(0) = \gamma/N$$  \hspace{1cm} (1)

where $\bar{N}$ is the average number of molecules inside the excitation volume and $\gamma$ is a geometric factor that is only determined by the shape of the point spread function, determined by the mathematical model used and the determined width parameters. The geometric factor for the Gaussian–Lorentzian model is 0.0762 (38). For a mixture of species, the G(0) value is related to the number of molecules of the individual species by (33)

$$G(0) = \sum_{m=1}^{M} \frac{(F_{m})^{2}}{G_{m}(0)}$$  \hspace{1cm} (2)

where $\langle F_{m} \rangle$ and $\langle F_{T} \rangle$ are the average fluorescence intensities of the overall mixture and the individual species, respectively, and $M$ is the total number of species.

The G(0) value was corrected for the buffer contribution by extending eq 2 for two species as shown in the equation:

$$G_{\text{measured}}(0) = \left( \frac{\langle F_{\text{sample}} \rangle}{\langle F_{T} \rangle} \right)^{2} G_{\text{sample}}(0) + \left( \frac{\langle F_{\text{buffer}} \rangle}{\langle F_{T} \rangle} \right)^{2} G_{\text{buffer}}(0)$$  \hspace{1cm} (3)

where the fluorescence intensity of the buffer, $F_{\text{buffer}}$, and the G(0) value of the buffer, $G_{\text{buffer}}(0)$, can be determined independently by performing FCS experiment on the buffer. $G_{\text{measured}}(0)$ is extrapolated by the experimental autocorrelation function, and $F_{\text{sample}}$ is the fluorescence intensity of the sample of interest and is equal to $\langle F_{T} \rangle - \langle F_{\text{buffer}} \rangle$. For our
experiments, the $G_{\text{buffer}}(0)$ is 9.5 and $F_{\text{buffer}}$ is 240 cps (counts per second), respectively.

We obtained the extrapolated $G(0)$ and the diffusion coefficient, for each sample in a series, by fitting the experimental autocorrelation function to eq 3. For a mixture of species, the recovered diffusion constant reflects the average diffusion constant of the mixture. The $G(0)_{\text{sample}}$ when multiple species are present, such as the dimeric and monomeric forms of PLA$_2$, gives an apparent number of particles weighted by the $G(0)_{s}$ and fractional contributions of each species. Once the protein is fully dissociated, only the monomeric species is present and the fluctuation amplitude characterizes again the true number of particles.

The standard deviations for $D$ and $G(0)$ were obtained by collecting a large single data stream and dividing it into 10 separate sets consisting of 9000–13 000 kB each. The resolved values and their associated errors were subsequently calculated from the independent fits to the 10 data sets. However, experimental uncertainties due to different instrument setup, optical alignment, and sample preparation from day to day, gave larger contributions to the point-to-point variations than the statistical error (Figure 3).

Dimer—Monomer Equilibrium. According to classical chemical thermodynamics, the equilibrium of dimer and monomers admits a single dissociation constant $K_d$ related to $\alpha$, the degree of dissociation of the dimer, by the expression:

$$K_d = \frac{4\alpha^2 C}{1 - \alpha}$$  

(4)

where $C$ is the total protein concentration in terms of the protein dimer. In logarithmic form:

$$\log_{4C} K_d = \log \left[ \frac{\alpha^2}{(1 - \alpha)} \right]$$  

(5)

The plot of degree of dissociation $\alpha$ against $\log C$ has a characteristic span of 2.86 decimal logarithmic unit between $\alpha = 0.1$ and $\alpha = 0.9$ (39, 40).

**RESULTS**

$K_d$ Estimation by Dilution. To determine the solution dissociation constant for the C. atrox PLA$_2$ dimer, we have measured the number of fluorescent particles in the excitation volume using the fluctuation amplitude in the FCS experiment (eq 1) as we dilute the enzyme solution. Figure 1 shows typical autocorrelation curves obtained in this study. They were measured for fluorescein (used for instrument calibration) and Flu-PLA$_2$. The diffusion coefficient obtained for the protein was 75 $\mu$m$^2$/s. The $G(0)$ was normalized in this figure but will depend on the protein concentration at each point of the dilution process. In Figure 2, the number of particles multiplied by the dilution factor is plotted versus concentration. In this plot, a straight line indicates that the number of molecules changes in proportion with the dilution. If there is dissociation, a deviation from the straight line will be observed. For the control experiment (dilution of fluorescein in 50 mM Tris, pH 8.0, 1 mM EDTA), we observed a straight line down to $1.5 \times 10^{-10}$ M (open squares). This is the expected result since fluorescein does not self-aggregate nor stick to the surfaces of our sample holder, so the number of particles changes exclusively due to the dilution. When the Flu-PLA$_2$ was diluted, we observed a straight line in the analysis plot (Figure 2). The addition of 10 mM calcium did not change the scenario: the number of molecules decreased proportionally with the dilution, indicating that the PLA$_2$ dimer remained intact. Using eq 4, we calculated that the maximum value of $K_d$ compatible with these data would be in the 0.01 nM range. Assuming a $K_d$ value, a simulated curve can be drawn that represents the hypothetical dissociation pattern in a given concentration range. In Figure 2, two simulations are shown for dissociation constants of 36 (a) and 4.3 nM (b).

![Figure 1: Normalized autocorrelation curve $G(\tau)$. Normalized autocorrelation for data taken at 100 kHz for fluorescein (filled circles) and fluorescein-labeled PLA$_2$ (open circles). Curved lines correspond to the best fit parameters (see Materials and Methods). $G(0)$ values have been normalized, and the instrument was calibrated with fluorescein using a diffusion coefficient of 300 $\mu$m$^2$/s. The dimer PLA$_2$ data were best fit to a diffusion coefficient of 75 $\mu$m$^2$/s.](Image 3)

![Figure 2: Changes of particle number with dilution. Samples of fluorescein-labeled C. atrox PLA$_2$ have been diluted in the absence (open circles) and in the presence of 10 mM Ca$^{2+}$ (filled circles) in 50 mM Tris, pH 8. Fluorescein in the same buffer was used as a control (open squares). The number of particles was calculated from the $G(0)$, and the number of particles multiplied by the dilution factor was plotted against protein concentration (in terms of monomer concentration). For reference, simulation curves are shown for dissociation constants of 36 (a) and 4.3 nM (b).](Image 4)
These $K_d$ values correspond to those reported in the literature for the dissociation constant in the absence and in the presence of Ca$^{2+}$ (24).

Urea was used as an external agent to dissociate the PLA$_2$ dimer. The number of fluorescent particles in the excitation volume at different concentrations of urea is shown in Figure 3. Again fluorescein was used as a control (filled squares), and the number of molecules did not change with increasing concentration of denaturant. In the case of the PLA$_2$ dimer in the presence (filled circles) and absence (open circles) of calcium, the number of molecules in the excitation volume changed from 4 to 8. However, the concentration ranges where these changes occur are different. When calcium is present, the changes occur between 1 and 3.5 M with a middle point around 2.0 M urea. In the absence of calcium, the middle point occurs at a lower concentration, 0.8 M urea.

For each concentration of urea an apparent $K_d$ was calculated using eq 4. We used the method proposed by Santoro and Bolen (41) to extrapolate the dissociation constants at high urea concentration to zero urea concentration. We obtained extrapolated $K_d$ values of 3.5 nM for the dissociation in the absence of calcium and 0.63 nM when calcium was present. Figure 3 also shows the CD signal at 222 nm for each urea concentration is also shown (open diamonds).

Table 1: Solution Dissociation Constants for C. atrox PLA$_2$

<table>
<thead>
<tr>
<th>Conditions</th>
<th>$K_d$ (nM)</th>
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</thead>
<tbody>
<tr>
<td>FCS dilution experiments</td>
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<tr>
<td>1 mM EDTA</td>
<td>$\leq$ 0.01</td>
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<tr>
<td>10 mM Ca$^{2+}$</td>
<td>$\leq$ 0.01</td>
</tr>
<tr>
<td>FCS urea dissociation$^a$</td>
<td>1 mM EDTA</td>
</tr>
<tr>
<td>10 mM Ca$^{2+}$</td>
<td>0.63</td>
</tr>
<tr>
<td>chromatography$^b$</td>
<td>1 mM EDTA</td>
</tr>
<tr>
<td>10 mM Ca$^{2+}$</td>
<td>4.3</td>
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</tbody>
</table>

$^a$ Dissociation constant obtained using FCS results and the linear extrapolation method. $^b$Data reported by Myatt et al. (24).

PLA$_2$–Lipid Interactions. We measured three parameters, (a) center of mass (squares), (b) area of the tryptophan emission peak (crosses), and (c) number of particles in the excitation volume (filled circles), to study the interaction of C. atrox PLA$_2$ with the lipid analogue n-dodecylphosphocholine (C12-PN) in the absence and presence of 10 mM Ca$^{2+}$ (Figure 4).

Ca$^{2+}$ (Figure 4). The number of PLA$_2$ particles in the excitation volume (filled circles) behaves similarly in the presence or absence of Ca$^{2+}$ and increases from 6 to 12 as a function of lipid analogue concentration. However, the concentration range where the changes occurred depended on the presence of Ca$^{2+}$. In the absence of calcium, the range spanned from 6 to 20 mM with a middle point at 8 mM. In the presence of 10 mM Ca$^{2+}$ the range becomes larger, from 2.5 to 20 mM, and the middle point moves up to 10.5 mM. At 20 mM lipids the PLA$_2$ dimers were completely dissociated in both cases.
Table 2: Diffusion Coefficients (μm²/s) for the Fluoro-PLA₂, C12-PN Micelles, and Fluoro-PLA₂/C12-PN Mixtures

<table>
<thead>
<tr>
<th>C12-PN (mM)</th>
<th>1 mM EDTA</th>
<th>10 mM calcium</th>
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<tbody>
<tr>
<td>0.0</td>
<td>75</td>
<td>75</td>
</tr>
<tr>
<td>0.04</td>
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<td>0.08</td>
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<tr>
<td>15</td>
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<td>60</td>
</tr>
<tr>
<td>20</td>
<td>60</td>
<td>55</td>
</tr>
<tr>
<td>C12-PN micelles</td>
<td>57</td>
<td></td>
</tr>
</tbody>
</table>

*Obtained from the fitting of the autocorrelation function as described in Materials and Methods. Errors in the diffusion constants were estimated to be no greater than 10%. 8 50 mM Tris, pH 8. C12-PN micelles (10 mM) containing Laurdan.

In addition to the particle number, the diffusion constant of the particle gives a measure of the extent of C12-PN association with PLAs₂. The diffusion constant of Fluo-PLA₂ as a function of C12-PN concentration is given in Table 2. At monomer C12-PN concentrations [below the CMC of 1.1 mM (42)] the particle diffusion constant was found to be approximately 72 μm²/s, similar to the value found for the PLAs₂ dimer, 75 μm²/s. As the concentration of C12-PN was increased above the CMC, both in the presence and in the absence of Ca²⁺, the diffusion constant progressively decreased, indicating an increase in particle size. At 20 mM C12-PN a diffusion constant of 60 μm²/s (55 μm²/s in the presence of Ca²⁺) was reached. The diffusion constant for micellar C12-PN was obtained through the collection of FCS data on Laurdan-containing micelles and was found to be 57 μm²/s (Table 2).

The area of the tryptophan emission spectra also changed when C12-PN was added. The final intensity for the two cases studied, with and without Ca²⁺, was the same, but the starting point was lower in the presence of Ca²⁺. The fluorescence increase in both cases followed a sigmoidal behavior. In the presence of EDTA, the spectral changes occurred in the lipid concentration range between 1 and 10 mM with a middle point at 4.5 mM. In contrast, when calcium was added, the range was larger, from 0.3 to 20 mM, and the middle point was near 3 mM C12-PN.

The center of mass of tryptophan emission spectra is another marker of lipid/protein interaction. A blue shift of tryptophan emission spectra was observed upon lipid titration in both cases, in the presence and absence of Ca²⁺. The plateau of the blue shift was observed around 20 mM in both cases with a value of 349.5 and 348.0 nm in the presence and absence of 10 mM Ca²⁺, respectively.

**DISCUSSION**

The primary focus of this work is on the dimer—monomer equilibrium of the strong dimer PLA₂ from C. atrox venom. To determine the dissociation constant, we used an approach based on the fluctuation correlation spectroscopy technique. This technique is capable of determining the number of fluorescent particles in the excitation volume (approximately 0.1 FL in our instrument) and the diffusion coefficient of the particles. In this way, we were able to determine the aggregation state of the protein (monomer vs dimer) in the presence of denaturant and in the presence of lipids. Our work constitutes a significant advance because we were able to follow directly the dissociation of dimer PLA₂ in solution, in the presence of lipid monomers and lipid micelles. The determination of the dissociation of the enzyme has historically been a difficult task since the PLA₂ subunit affinity is high and the protein needs to be diluted several orders of magnitude to accurately determine the dissociation constant (39, 43). At each point of the dilution, we must measure a property which can distinguish monomer from dimer. The first experimental problem generally encountered is the sensitivity of a technique for measuring the desired property. The tighter the dimer, the more challenging it is to find an adequately sensitive technique. Fluorescence spectroscopy is one of the more sensitive spectroscopic methodologies commonly applied to studies on binding equilibria. In fact, fluorescence methods can achieve single molecule detection levels. In some systems the intrinsic protein fluorescence can be used, but tryptophan fluorescence, the primary protein fluorophore, is not strong enough for measuring K_d values much below 10⁻⁸ M. For high sensitivity such as this work, extrinsic fluorophores must be used.

A second consideration in studying binding equilibria is that the chosen method must adequately distinguish monomer from dimer. Fluorescence polarization is commonly used to study aggregation equilibria in proteins. Polarization measurements are based on the changes in particle hydrodynamic volume during protein dissociation (32, 43, 44). Although dependent on the particle shape, the ratio between dimer and monomer rotational correlation times should be approximately 2. This technique can be applied to solution studies, but it cannot be easily applied for our specific case in which the changes in the association state of PLAs₂ are accompanied by changes in the interaction of PLA₂ with organized lipids. For example, in our condition in which the PLAs₂ begins associating with lipids and simultaneously dissociating from dimer to monomer, there would be an increase in the hydrodynamic volume due to lipid binding (in the form of monomers, micelles, or vesicles) but a decrease in the volume due to the dissociation of the dimer. In this scenario we would observe an average polarization change, a distribution of rotational correlation times, which would be difficult to accurately interpret. In contrast, fluorescence correlation spectroscopy will not only give us the sensitivity, which we expect from fluorescence techniques, but also allows us to detect dissociation in terms of the number of molecules in the excitation volume. If a dimer dissociates into its monomers, the number of fluorescent molecules in the excitation volume will increase by a factor of 2 due to dissociation (dimer to monomer). The upper concentration in the FCS experiments is determined mainly by the fluctuation amplitude, G(0). The inverse relation between the number of molecules in the excitation volume (N) and G(0) indicates the importance of limiting the number of molecules inside the excitation volume to observe appreciable fluctuations. If the volume of excitation contains many molecules, the average intensity will fluctuate very little. The lower concentration limit in the fluorescence fluctuation spectroscopy experiments is imposed by the signal-to-noise ratio, which in our case is determined by the fluorescence.
contribution from the buffer. The fluorescence intensity of the buffer is relatively high when compared to water. Therefore, it is important for the more diluted samples that we take the buffer contribution into account (eq 2).

The effect of pH and Ca\(^{2+}\) on the binding constants has been previously studied using chromatographic profiles (24). By examining the width and position of the gel filtration profiles, the authors estimated that the \(K_d\) is approximately 36 nM and 4.3 nM in the absence and presence of Ca\(^{2+}\), respectively. From our dilution data, no dissociation is observed. The enzyme \(K_d\), according to our calculation, would be on the order of 0.01 nM. Plotting our data and the simulated curves using their \(K_d\) values (Figure 2) shows obvious disagreement.

The discrepancy between our data and the reported values might be explained by the nature of the PLA\(_2\). The phospholipases A\(_2\) are surface-active proteins. Techniques such as gel filtration rely on the assumption that the protein does not associate with the column matrix. Any interaction will drive the protein to longer retention times and lower apparent molecular weights. These effects are particularly troublesome at low protein concentrations. It is also true that these techniques are carried out under nonequilibrium conditions, which can also influence the results. In contrast, in FCS one collects data on a protein solution under equilibrium conditions. However, we must acknowledge that we have an extrinsic label attached to the PLA\(_2\) which may alter the monomer–dimer equilibrium. We believe our conjugate is well behaved since its activity appears normal against micellar substrate, and modifications would generally be thought to disrupt specific interactions, not enhance them.

The use of reagents such as urea to weaken the molecular interactions is a common tool in oligomer biochemistry. The problem associated with using these types of reagents is that it is possible to denature the protein subunits as well as perturb the oligomer equilibrium (32). The fact that the CD signal does not change in the range of urea concentrations used indicates that at least the secondary structure of the protein was unaffected. The stability of PLA\(_2\) in urea has been reported by others, and it is not a surprising result, since C. atrox venom PLA\(_2\) contains seven sulfur bridges per monomer which greatly stabilize the subunit structure (45).

We explored the possibility of obtaining indirectly the \(K_d\) values using the “linear extrapolation method” (41) from urea dissociation experiments. This methodology is normally used in denaturation studies to calculate the stabilization free energy of the folded state by extrapolation of the denaturant-unfolded state’s free energies to zero denaturant concentration. The assumption for the application of this methodology is the existence of two states, native (N) and unfolded (U) in equilibrium, with a constant \(K_{\text{denat}}\) and that urea does not directly participate in the equilibrium. In the case of a monomer–dimer equilibrium we have two states, monomer and dimer, with a single equilibrium constant, \(K_d\). We calculate the extrapolated \(K_d\) in the presence (\(K_d = 0.63\) nM) and in the absence of 10 mM Ca\(^{2+}\) (\(K_d = 3.5\) nM), which indicates that Ca\(^{2+}\) increases the dimer stability. However, the extrapolated \(K_d\) values are not compatible with the upper limit obtained from the direct dilution experiments, indicating that urea participates in a complex manner. An interesting work published by Weber’s group some years ago studied the pressure dissociation of the R17 bacteriophage (46). They reported a different dissociation pattern when urea was used to facilitate the dissociation compared with pressure-induced dissociation in the absence of urea. The authors concluded that urea loosens the interaction between the monomers, but at the same time it fundamentally changed the interfacial binding sites. Their work, and the work presented here, suggests that care must be taken using these extrapolation methods in determining free energies for even relatively well-defined two-state processes.

Changes in the tryptophan emission when PLA\(_2\) interacts with organized lipid structures, such as micelles, have been reported. The question is whether these changes are related or not with the dissociation of the enzyme. C. atrox PLA\(_2\) has six tryptophans (45), three tryptophan residues per subunit, two of which are located near the surface of the molecule. The third one is located near the calcium binding site, protected from the solvent by the companion PLA\(_2\) subunit. It is well-known that tryptophan emission can be affected by the polarity of the surroundings. Spectral blue shift is characteristic when the tryptophan environment becomes less polar (47). Given the number of the tryptophans present, the spectral shift and intensity changes cannot be unambiguously assigned to specific residues in the protein.

Unlike the tryptophan emission, the fluctuation amplitude measured by FCS derives from fluorescein molecules covalently bound to the protein. The fluctuations, as outlined in the Materials and Methods section, can give us information about the dissociation state of the molecule. In this way, the combination of tryptophan signal and the measurement of the number of molecules in the excitation volume gives us a handle to separate the effects of protein dissociation from protein/lipid interactions.

On the basis of our combined FCS and tryptophan fluorescence results, we propose the following model based on four states for the interactions of C. atrox PLA\(_2\) dimer with lipid micelles in the absence and presence of Ca\(^{2+}\) (Figure 5).

State a: In the absence of lipids, the enzyme is a dimer and remains in this state until at least 10\(^{-11}\) M according to our dilution studies. The addition of calcium at this stage produces an increase in intensity and a red shift (Figure 4) of the tryptophan emission spectra. Even if the changes in tryptophan intensity represent the average behavior of all the six tryptophans in the dimer protein, we can assume that the changes produced after the addition of Ca\(^{2+}\) are due to Ca\(^{2+}\) binding to the enzyme’s active site.

State b: C12-PN below its CMC of 1.1 mM (42). In this state, the C12-PN does not significantly interact with the protein as inferred by the lack of spectroscopic changes.

State c: C12-PN above its CMC. The lipid micelles are the primary units interacting with the protein. We propose an intermediate state where the enzyme is fully associated with the C12-PN, but it remains in the micelle still as a dimer, with weaker interaction between its monomers due to the presence of the lipids (Figure 5c). The micelle here is a protein–lipid comicelle with a much reduced number of lipid monomers, with respect to pure C12-PN micelles. In this case, we observe changes in the tryptophan emission mainly due to the exposure of the tryptophan to a more hydrophobic media (blue shift and increase in intensity), but we do not see changes in the number of particles in the excitation volume from FCS measurements. Our data indicate that the
changes in tryptophan emission when PLA$_{2S}$ interact with lipids are due to the interaction with the lipids and not to the dissociation of the dimer. A similar state has also been described for a monomeric PLA$_2$ porcine pancreas (42, 48, 49).

State d: Excess C12-PN well above its CMC. As more C12-PN is added, changes in the tryptophan spectra (spectral shift and intensity) plateau. At the same time, the migration of PLA$_2$ monomers to different micelles occurs, as reported by the FCS results (particle number). Each monomer is surrounded by lipids, and no more changes could be observed in tryptophan emission. At these higher C12-PN concentrations the complex diffusion constant, 60 $\mu$m$^2$/s (55 $\mu$m$^2$/s in the presence of Ca$^{2+}$), suggests the association of PLA$_2$ with a large C12-PN micelle. This result indicates that we are observing the association of the PLA$_2$ monomers with the micelle interface and that the PLA$_2$ is not (no longer) part of a micelle.

FCS directly measures the dissociation of the dimer (state d), and tryptophan emission changes are the manifestation of two processes, the association with lipids (state c) and the dissociation of the enzyme (state d). There are two important differences revealed by the tryptophan fluorescence in the presence of calcium. First, with calcium the changes in tryptophan emission start to occur before 1 mM lipid concentration, indicating that the interaction between the lipids and the protein starts earlier in the presence of Ca$^{2+}$. Second, the number of fluorescent particles in the excitation volume starts to increase when the changes in tryptophan emission have almost finished, indicating that calcium stabilizes the intermediate state (state c in our model). Lipids are competing for the monomer interface. It would be reasonable to assume that a tighter dimer would require a higher lipid concentration for dimer dissociation. With this in mind, our results (Figure 4) would indicate that Ca$^{2+}$ increases the affinity of the dimer subunits and stabilizes the intermediate state (state c).

In summary, we have used tryptophan fluorescence and FCS to study the behavior of $C$. atrox venom PLA$_2$. By combining both techniques, we gain complementary information. FCS directly monitors the dissociation process of the PLA$_2$ dimer while changes in tryptophan emission reflect binding of lipid to PLA$_2$. In solution the $C$. atrox venom PLA$_2$ is a tight dimer with a dissociation constant at or below 0.01 nM, either in the presence or in the absence of Ca$^{2+}$ cofactor (Table 1). In the presence of the lipid analogue C12-PN two distinct states can be described: one in which a protein and lipid micelle form and a second in which the interaction appears to be more in the nature of interfacial binding of the enzyme to an intact C12-PN micelle. The relevant point in terms of PLA$_2$ activity is the clear indication that the PLA$_2$ is monomeric at high lipid concentrations. We observed a condition at C12-PN concentrations of 20 mM in which single PLA$_2$ subunits are bound to individual micelles of C12-PN. It is well documented that PLA$_2$ is highly active against mixed micellar substrates, such as the Triton X-100 and egg phosphatidylcholine assay system used here. It seems likely that substrate mixed micelles and C12-PN micelles would act toward PLA$_2$ in a similar fashion. Thus, our observation that the nonhydrolyzable substrate analogue C12-PN dissociates a dimeric PLA$_2$ supports the single enzyme hypothesis for PLA$_2$ action on organized lipid substrate.

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The authors dedicate this work to the memory of Professor Gregorio Weber (1916–1997).

REFERENCES

Aggregation States of C. atrox PLA₂