Fluorescence Polarization and Fluctuation Analysis Monitors Subunit Proximity, Stoichiometry, and Protein Complex Hydrodynamics

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Abstract

 Förster resonance energy transfer (FRET) microscopy is frequently used to study protein interactions and conformational changes in living cells. The utility of FRET is limited by false positive and negative signals. To overcome these limitations we have developed Fluorescence Polarization and Fluctuation Analysis (FPFA), a hybrid single-molecule based method combining time-resolved fluorescence anisotropy (homo-FRET) and fluorescence correlation spectroscopy. Using FPFA, homo-FRET (a 1–10 nm proximity gauge), brightness (a measure of the number of fluorescent subunits in a complex), and correlation time (an attribute sensitive to the mass and shape of a protein complex) can be simultaneously measured. These measurements together rigorously constrain the interpretation of FRET signals. Venus based control-constructs were used to validate FPFA. The utility of FPFA was demonstrated by measuring in living cells the number of subunits in the α-isofrom of Venus-tagged calcium-calmodulin dependent protein kinase-II (CaMKIIα) holoenzyme. Brightness analysis revealed that the holoenzyme has, on average, 11.9 ± 1.2 subunits, but values ranged from 10–14 in individual cells. Homo-FRET analysis simultaneously detected that catalytic domains were arranged as dimers in the dodecameric holoenzyme, and this paired organization was confirmed by quantitative hetero-FRET analysis. In freshly prepared cell homogenates FPFA detected only 10.2 ± 1.3 subunits in the holoenzyme with values ranging from 9–12. Despite the reduction in subunit number, catalytic domains were still arranged as pairs in homogenates. Thus, FPFA suggests that while the absolute number of subunits in an auto-inhibited holoenzyme might vary from cell to cell, the organization of catalytic domains into pairs is preserved.

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Introduction

In cells, proteins rarely function individually; typically they interact with other proteins to mediate cellular processes. Determining the structure and dynamics of protein complexes in cells requires noninvasive high-resolution spatial and temporal methods, such as Förster resonance energy transfer (FRET) microscopy [1,2,3,4,5,6]. Because the efficiency of FRET is inversely proportional to the 6th power of the distance separating fluorescent donors and acceptors [2,3,7], FRET has been used to study protein interactions in cells. Energy transfer between proteins tagged with either the same fluorophore (homo-FRET [8,9,10,11,12]) or spectrally distinct fluorophores (hetero-FRET [2,3]) can be monitored using time correlated single photon counting (TCSPC [13]) to measure dynamic changes in fluorescence polarization or intensity respectively. However, the absence of FRET does not necessarily mean that proteins are not in a complex because the distance separating fluorophores in a complex may be larger than 10 nm (or their dipole orientation may be unfavorable). Similarly, a positive FRET signal may arise from non-specific FRET caused by protein over-expression rather than complex formation. These limitations are illustrated in figure 1A, in which six possible arrangements of a fluorescent protein-tagged subunit are depicted. In this diagram homo-FRET is expected for protein complexes if their attached fluorophores (yellow cylinders) are in close proximity. Thus, homo-FRET would not be observed for examples 1, 3, and 5 (even though example 3 is a dimer and example 5 is a hexamer), but FRET should be observed in examples 2 and 4. Furthermore, FRET measurements alone cannot differentiate between example 2 (a dimer) and example 4 (a hexamer). Homo-FRET should also be observed for example 6, monomers in close proximity. One potential way to differentiate all six subunit arrangements depicted in figure 1A is to measure homo-FRET and Fluorescence Correlation Spectroscopy (FCS) simultaneously.

Unlike FRET, FCS monitors the motion of fluorophore-tagged proteins, not their proximity [14,15,16,17]. In a FCS experiment temporal fluctuations of fluorescence intensity emanating from a small volume (typically less than 1 fl) are measured as individual protein complexes traverse the observation volume [16,17]. Auto- and cross-correlation analysis [16,17,18] of these fluctuations can then reveal the average number of fluorescent molecules in the observation volume, <N>. Molecular brightness, the average number of photon counts per second per fluctuating molecule (cpsm), is obtained by taking the ratio <k>/<N>, where <k> is
Figure 1. Fluorescence Polarization and Fluctuation Analysis: A method for studying the structure of protein complexes. (A) Six possible structures for a hypothetical protein (Blue hexagon) tagged with Venus (yellow cylinder). Example 1 depicts monomers. Examples 2 and 3 depict dimers. Attached Venus molecules are in close apposition for example 2 but not for example 3. Examples 4 and 5 depict hexamers. Pairs of attached Venus molecules are in close apposition for example 4 but not for example 5. Example 6 depicts monomers that are confined to a sub-compartment (dashed line) where they are expressed at a high concentration. (B) Schematic for microscope to measure FPFA. (C) Micro-time data measured for a homogenate prepared from cells expressing Venus monomers are used to calculate fluorescence lifetime histograms for photons detected by either the parallel or perpendicular hybrid-detectors as a function of time after the laser excitation pulse (left panel). These two decay curves are then used to calculate the time-resolved anisotropy (right panel top), or fluorescence lifetime (right panel bottom) of Venus monomers. Red dashed lines show fitting to a single exponential model. (D) Macro-time data measured for a homogenate prepared from cells expressing Venus monomers are used to calculate auto- and cross-correlation functions for photons detected in the parallel and perpendicular hybrid-detectors. A single diffusible-component 3-D Gaussian model was used to fit the cross-correlation (red dashed line), and to estimate the correlation time and the average number of molecules in the observation volume. The excitation power used was 10.2 mW.

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the average fluorescent count rate [17]. With control experiments to measure the brightness of free fluorophore, the molecular brightness, can be used to calculate the Normalized Brightness - the average number of fluorophores in a protein complex. If each subunit is tagged with only a single fluorophore, the normalized brightness will reflect the number of subunits in a complex. This is called Brightness Analysis [19]. In addition, FCS can also determine the average amount of time a protein complex remains in the observation volume [17]; a value related to the lateral diffusion coefficient of the complex, itself a function of viscosity, mass and hydrodynamic volume (and hence the conformation of the complex). Another technical advantage of FCS analysis is that single molecule fluctuations in the fluorescent intensity can only be observed at very low fluorophore concentrations (typically nanomolar) [17], thus, many of the problems associated with over expression of exogenous proteins cannot exist in successful FCS experiments. By considering both the presence (or absence) of homo-FRET, as well as the normalized brightness, all 6 examples depicted in figure 1A could be differentiated. Furthermore, often these interpretations could be corroborated using the lateral diffusion time, an additional parameter measured.

While FRET and FCS are usually considered alternative approaches for studying protein complexes [20], figure 1A illustrates that combining them could be advantageous. New instrumentation for TCSPC can now record both micro-time (the elapsed time between photon detection and the laser excitation pulse) and macro-time (the elapsed time between the start of an experiment and when an individual photon is detected) [13,21]. This technology enables combining FCS (based on macro-time data) and either hetero- or homo-FRET (which both use micro-time data) with high photon efficiency. While combining hetero-FRET and FCS is problematic (donor emission may bleed into the acceptor signal, and hetero-FRET itself will cause reciprocal intensity fluctuations in donor and acceptor channels) [20], combining homo-FRET and FCS is more straightforward (because only a single fluorophore is used). An additional benefit of combining homo-FRET and FCS is that spectral bandwidth for fluorophore excitation and emission is kept narrow enabling the use of other fluorophores in more complicated multiplexed experiments.

Here we describe a new method, Fluorescence Polarization and Fluctuation Analysis (FPFA), that combines Förster resonance energy transfer microscopy (FRET) [1,2,3,4,5,6] and Fluorescence Correlation Spectroscopy (FCS) [14,15,16,17] as a tool to study the structure of protein complexes inside living cells. We developed this method specifically to characterize the structure of the calcium-calmodulin dependent protein kinase-II (CaMKII) holoenzyme under physiological conditions. CaMKII is an excellent example of a protein whose function depends on protein-protein interactions. CaMKII subunits assemble to form a multimeric kinase involved in memory and synaptic modulation [22,23]. The number of subunits that comprise the holoenzyme is uncertain with estimates ranging from 3 to 14 [24,25,26,27,28,29,30]. In cells the kinase is transiently activated in response to a rise in intracellular calcium. Calcium binds to calmodulin (CaM) to form calcium/calmodulin (Ca2+/CaM) [23], and Ca2+/CaM in turn binds to the regulatory domain of CaMKII. Ca2+/CaM binding to CaMKII is thought to trigger a large conformational change that opens the substrate-binding site on the catalytic domain [30]. Unlike most other biological calcium sensors, CaMKII is sensitive to the frequency of calcium spikes [31]. This unique ability is thought to be an emergent feature of its holoenzyme organization [32], and is thought to play a role in memory storage [22]. Little is known about the structure or stoichiometry of the holoenzyme under physiological conditions, in part because few methods exist to monitor the structure of protein complexes in cells.

CaMKII monomers have three domains, the N-terminal catalytic domain, a regulatory domain, and a C-terminal association domain responsible for holoenzyme oligomerization [23,33]. In the auto-inhibited holoenzyme, catalytic domains are thought to dimerize [10,30,34], resulting in a holoenzyme structure with multiple catalytic domain pairs distributed around a central association domain core. X-ray crystallography was first to indicate that isolated CaMKIIΔc catalytic domains are arranged as dimers [34], but the validity of this conclusion is controversial, as this dimeric structure was not observed in X-ray data sets for isolated catalytic domains from several other CaMKII isoforms [30]. Nonetheless, analytic ultracentrifugation of isolated catalytic domains from multiple isoforms did detect catalytic domain dimerization [30], as did homo-FRET experiments with fluorescent protein-tagged CaMKIIΔc in hippocampal neurons [10]. Catalytic domain pairing was not observed in a recent crystal structure of a mutated CaMKIIΔb holoenzyme. In this new model for the holoenzyme structure individual catalytic domains are docked onto the central association domain core as monomers to form a compact holoenzyme structure [35]. Because Ca2+/CaM binding sites on the regulatory domain are inaccessible in this compact structure, it was proposed that a more extended auto-inhibited structure must also exist to allow for activation. Mutation in the CaMKII regulatory domain (TT305/306DD for the α isoform) are known to prevent Ca2+/CaM from binding to CaMKII [36], and based on this new model are now predicted to also disrupt the docking of catalytic domains to the association domain core [35]. Thus it is possible that catalytic domains might form pairs after they un-dock from the association domain core. It should be noted that unlike most CaMKII isoforms, the CaMKIIΔb isoform had dramatically reduced levels of catalytic activity under physiological conditions [35,37]. Thus, it is unclear if this new model for the intact CaMKIIΔb holoenzyme reflects the structure of active CaMKII isoforms in cells.

In this study we develop a new method for characterizing protein complexes in living cells, FPFA. We first validate this method with two sets of Venus [38] based control constructs, and then apply FPFA to simultaneously measure the number of subunits in the CaMKIIΔc holoenzyme in cells, as well as the organization of its catalytic domains.

Results

The experimental setup used for linear-polarized two-photon excitation of fluorophores in a small focal volume is shown in figure 1B. Both micro- and macro-times of emitted photons detected through orthogonally oriented polarization pathways (parallel and perpendicular) were recorded. Calibration experiments with fluorescein and Venus monomers were used to identify excitation powers in vitro where bleaching of Venus was not detected, and to then estimate the volume and dimensions of our observation volume under these conditions (see materials and methods). For in vivo measurements laser powers ranging from 9.0-10.2 mW were used. Time-resolved anisotropy (TRA) and lifetime were calculated from histograms of parallel and perpendicular detector micro-times (Fig 1C), while polarization corrected correlation times and brightness were calculated by fitting the cross-correlation of these detectors macro-time signals [39] (Fig 1D).

Validation of FPFA

FPFA was validated using 6 fluorescent oligomers composed of between 1 and 6 concatenated Venus molecules (V1–V6). Venus,
a yellow GFP spectral variant, was used because it matures rapidly [30], and has a Förster distance of 5.3 nm [40] supporting efficient homo-FRET. To avoid nonspecific aggregation, a monomeric variant of Venus was used in all experiments [41]. Immunoblot analysis using a GFP specific antibody confirmed that the molecular weight of these constructs increased linearly with the number of Venus molecules (Fig 2A). Micro-time data was used to calculate the TRA (Fig 2B) of homogenates prepared from cells expressing V1–V6. The V1 anisotropy decayed as a single-exponential ($\tau = 15.3 \pm 0.8$ ns). Like V1, V2–V6 also had a slow decay component ($\tau > 15$ ns), as well as additional fast decay components occurring primarily within the first 2 ns. The similarity between the V1 decay constant and the rotational time constant of purified Venus (16.4 ± 0.6 ns) [42] indicates that V1 is a monomer in solution. The additional fast depolarization observed for V2–V6, much faster than monomer rotation, is the hallmark of homo-FRET between fluorescent proteins [10,11]. While the amplitude of these fast anisotropy decay components is a function of the number of Venus molecules exchanging energy by FRET [10,11], the similarity between the V4–V6 anisotropy decay curves reveals a major limitation for using homo-FRET alone to estimate subunit stoichiometry in a complex with four or more subunits.

The same micro-time data was also used to calculate the lifetime of Venus in these control constructs (Fig 2C). The lifetime of Venus was 3.1 ± 0.0 ns, similar to values measured previously (3.03 ± 0.01 ns) [42], and its lifetime was not appreciably altered by concatenation or by homo-FRET. Because the photon count rate emitted from a molecule depends on the molecule’s lifetime, this observation also indicates that subsequent FCS based brightness analysis will not be significantly influenced by homo-FRET. Thus, the primary source of intensity fluctuations in the FPPA experiments described here should be the diffusion of Venus-tagged molecules in and out of the excitation volume.

Macro-time data was used to calculate the cross-correlation between photons detected in parallel and perpendicular channels [39]. Cross-correlations for Venus concatamers were all well fit using a single-component 3-dimensional Gaussian model [17], and these fits were used to derive the correlation time and the average number of diffusing molecules in the observation volume. The later was used to normalize the photon count rate to calculate molecular brightness as mentioned earlier. Figures 2D & E show the correlation time and molecular brightness obtained for each concatemer. A systematic increase in correlation time was observed as the number of Venus molecules per concatemer increased, but as expected, correlation time was not very sensitive to changes in mass. Molecular brightness also increased with the number of Venus molecules in a concatemer, but unlike the correlation time, molecular brightness was linearly proportional to the number of Venus molecules in a concatemer (Fig 2E). Thus, while changes in correlation time may serve as an indicator of complex formation, the ratio of the brightness of a construct and the brightness of a Venus monomer can directly reveal the number of Venus molecules in a construct. This normalized brightness analysis can potentially be used to specify the number of fluorophore-tagged subunits in a protein complex, as will be shown later when FPPA is used to investigate the holoenzyme structure of Venus-tagged CaMKII. 

Because FPPA can measure homo-FRET, it should be able to detect structural changes that FCS cannot. To test this capability, FPPA was performed on three different Venus dimers, V5V (the V2 construct), V17V and V32V, where 5, 17 and 32 amino acids linkers separate Venus molecules respectively. TRA analysis revealed that these 3 dimers all had fast decay components indicative of homo-FRET [11] (Fig 2F). The decay rate of these fast components should be proportional to the FRET transfer rate, itself a function of the distance between Venus molecules [11]. Consistent with this, the anisotropy of V5V decayed fastest while V32V decayed the slowest. In contrast, the correlation times and brightness of V5V, V17V, and V32V were comparable (Fig 2F inset).

**FPPA Studies of CaMKIIα Holoenzyme in vitro**

To measure the number of subunits in the CaMKIIα holoenzyme with FPPA, cells were transfected with DNA encoding Venus-CaMKIIα [V-CaMKIIα]. Previous studies have shown that tagging the N-terminus of CaMKII with GFP does not alter its catalytic activity, auto-phosphorylation, or the ability of the kinase to assemble [36,43]. Furthermore, we have shown that V-CaMKIIα expressed in neurons undergoes a change in anisotropy correlated with calcium influx through NMDA receptors [10]. The next day, transfected cells were harvested and homogenates prepared for FPPA. Supernatants from homogenates of cells expressing V-CaMKIIα had a normalized brightness of 10.2 ± 1.3 ($n = 5$, mean ± SD) (Fig 2A). FPPA experiments were repeated using V-CaMKIIα[TT305/306DD] a mutant that cannot bind Ca$^{2+}$/CaM to determine if the auto-inhibited holoenzyme is also a decamer. V-CaMKIIα[TT305/306DD] had a mean normalized brightness of 10.7 ± 1.3 ($n = 5$, mean ± SD). These values were not statistically different [paired Student’s t test, P = 0.2967]. Nearly an identical number of subunits was determined using electron microscopy to count catalytic-domain protrusions on wild-type holoenzyme isolated from rat forebrain [predominantly CaMKIIα] [24]. Thus, in solution there is on average 10 V-CaMKIIα subunits in the auto-inhibited holoenzyme.

The correlation time for V-CaMKIIα holoenzyme was 1.2 ± 0.1 ms (mean ± SD, $n = 5$) and for V-CaMKIIα[TT305/306DD] it was 1.3 ± 0.1 ms (mean ± SD, $n = 5$) (Fig 3B). As expected for a large holoenzyme, these samples dispersed much slower than the Venus monomer (Fig 2D). TRA analysis revealed a fast homo-FRET component for V-CaMKIIα and V-CaMKIIα[TT305/306DD] indicative of catalytic domain pairing (Fig 3). Hetero-FRET analysis was used to confirm the existence of catalytic domain pairing in the CaMKIIα holoenzyme, as well as to measure the FRET efficiency detected between fluorescent protein tagged catalytic domains. Cells were transfected with DNA constructs encoding CaMKIIα tagged on the catalytic domain with either Cerulean [44] acting as a FRET donor, or Venus [38] as a FRET acceptor. Cells transfected with Cerulean-CaMKIIα (C-CaMKIIα) & free Venus monomers were used as a negative FRET control, while cells transfected with C5V, a construct with a Cerulean donor tethered to a Venus acceptor using a 5 amino-acid linker [45,46], was used as a positive FRET control. FRET efficiency values and the fraction donor were measured the following day from live cells using E-FRET microscopy [47,48]. Cells transfected with C-CaMKIIα & V-CaMKIIα had FRET efficiencies that increased linearly as the donor fraction decreased (Fig 3D). A linear relationship between the observed FRET efficiency and the fraction donor is indicative of FRET occurring between 1 donor and 1 acceptor [49], and suggests that catalytic domains form pairs within the holoenzyme structure in living cells. The FRET efficiency (the y-intercept when the donor fraction is zero) between fluorescent protein-tagged CaMKIIα catalytic domains was 39.4 ± 0.8%, while the FRET efficiency for our negative control was 5.6 ± 0.8%. The FRET efficiency for the C5V positive control was 48.9 ± 3.8% (mean ± SD, n = 34 cells), and C5V’s fraction
donor was 0.50±0.01 as expected for a construct with one donor covalently linked to one acceptor.

FPFA Measurements in Cells

To measure the number of subunits that comprise the CaMKIIα holoenzyme in living cells using FPFA required expressing V-CaMKIIα at low enough concentrations to detect fluctuation, while also allowing sufficient time for protein expression, Venus maturation, and holoenzyme assembly. Because the amount of protein expression at 24 hours could be controlled by titrating the amount of RNA used in a transfection [50], RNA encoding V-CaMKIIα was used for live cell FPFA. Additionally, the laser power for live cell FPFA was reduced to 6 mW to prevent bleaching. The TRA of V-CaMKIIα in cells (Fig 4A) was similar to the signals observed in solution being more depolarized than the V1 monomer, but less depolarized than the V3 Venus trimer (Fig 3C), and together with hetero-FRET experiments in cells (Fig 3D) indicates that catalytic domain pairing occurs in cells as well as in solution. The brightness of cells transfected with RNA encoding the V1–V6 controls are shown in figure 4B. In cells, the brightness of these concatemers was still linear with the number of Venus molecules with the exception that V1 showed a slightly elevated value. This can be attributed to endogenous autofluorescence. The V2–V6 brightness values were well fit to a linear model so a linear interpolation was used to calculate normalized brightness of Venus-CaMKIIα holoenzyme in cells. The normalized brightness for V-CaMKIIα in cells was 11.9±1.2 (mean±SD, n = 11 cells) (Fig 4B). Surprisingly, the holoenzyme in cells had approximately 2 more subunits than holoenzyme in solution. The variance of this measurement was twice as large as the 95% prediction bands of the linear fit of our controls, suggesting that in cells V-CaMKIIα has on average 12 subunits, but ranges from 8–14 subunits.
Discussion

FPFA was validated using two sets of controls. The first set consisted of 6 Venus concatemers. FPFA readily differentiated these molecules by their brightness, and to a limited extent by their anisotropy and correlation time (Fig 2). Importantly, brightness analysis unambiguously determined the number of Venus molecules in each construct (Fig 2E). The second set of controls consisted of 3 Venus dimers separated by different length amino-acid linkers. Here, based on different fluorophore separation distances, anisotropy could differentiate between these molecules (Fig 2F). Thus, FPFA can detect subtle structural differences based on homo-FRET, changes in mass and hydrodynamics using correlation times, while simultaneously monitoring the number of fluorophores in a complex using brightness analysis.

Figure 3. FPFA of CaMKIIα holoenzyme. (A) The normalized brightness for V-CaMKIIXα holoenzyme, and mutant that cannot bind Ca\(^{2+}\)/CaM. Bars represent the means with n = 5. The excitation power used was 10.2 mW. (B) The correlation times for samples in panel A. (C) Average TRA for samples in panel A. TRA traces for Venus monomers (V1) and Venus trimers (V3) from figure 2B are overlaid as a reference, and to illustrate that the Venus-tagged catalytic domains in V-CaMKIIαs produce an anisotropy signal most consistent with Venus-dimers. (D) Hetero-FRET analysis of CaMKIIα catalytic domain pairing in living cells. Cells were transfected with DNA constructs encoding CaMKIIα tagged on the catalytic domain with either Cerulean or Venus. Cells transfected with Cerulean-CaMKIIα & free Venus monomers were used as a negative FRET control. Cells transfected with CSV was used as a positive FRET control. Each point is the average FRET efficiency and fraction donor value measured for an individual cell. Dashed lines are linear fits.

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Figure 4. FPFA of V-CaMKII\(\alpha\) expressed in HEK cells. (A) Average TRA for V-CaMKII\(\alpha\) (Black) holoenzyme in cells. TRA traces for Venus monomers (V1, Yellow) and Venus trimers (V3, Green) expressed in HEK cells are also plotted as a reference, and to illustrate that the Venus-tagged catalytic domains in V-CaMKII\(\alpha\) produce an anisotropy signal most consistent with Venus-dimers. The excitation power used was 6 mW. (B) The brightness values for V1–V6 expressed and measured in HEK cells are plotted as a function of the number of Venus molecules in each construct. Each
Several potential sources of error in FPFA measurements warrant consideration. First, polarization artifacts can be avoided by measuring the cross-correlation of parallel and perpendicular polarized detectors [39]. Because the molecular brightness of V5V, V17V and V32V yielded comparable values (although their time-resolved anisotropies were visibly different) this conclusion is supported (Fig 2F). A second potential source of error is the impact of flickering on brightness measurements [19]. Because fluorescent protein flickering has not been observed with two-photon excitation, flickering should not impact FPFA measurements as implemented here. This conclusion is supported by the observed linear relationship between molecular brightness and the number of Venus molecules in a concatemer (Fig 2E). Finally, for live cell FPFA measurements, auto-fluorescence can corrupt brightness measurements [16,17]. The apparent brightness of a sample with more than one fluorophore species is thought to be a nonlinear function weighted by the abundance of each fluorescent species multiplied by the square of its brightness [18,51] (see discussion in Materials and Methods). In typical live cell experiments the fluorophore tag and the wavelength used to excite it are both selected to maximize the fluorescent signal of the fluorophore, and to minimize cellular auto-fluorescence. Under these conditions, the brightness of the sample will typically be much brighter than the brightness of auto-fluorescent species. Even if the concentration of these dim auto-fluorescent species are significantly higher than the concentration of the fluorophore-tagged species, if a sample has or can be tagged with multiple copies of the fluorophore, eventually the contribution of auto-fluorescence to the apparent brightness of the sample will become negligible. This was observed in figure 4B for Venus concatemers.

Using FPFA we show that in cells V-CaMKII\(\alpha\) holoenzyme has on average 12 subunits (Fig 4B), with its catalytic-domains arranged as pairs (Fig 4A). The pairing of catalytic domains in the holoenzyme was previously observed in hippocampal neurons using anisotropy measurements [10], and this structural feature of the auto-inhibited holoenzyme was confirmed using hetero-FRET measurements in cells (Fig 3D). It is unlikely that attached fluorescent proteins caused this dimerization because 1) a monomeric form of Cerulean and Venus were used in all experiments [41], 2) homo-FRET was not observed between Venus monomers (V1, Figs 2B, & 4A), and 3) isolated non-tagged catalytic domains formed dimers in solution [30].

It is likely that these dimers exist specifically in the postulated auto-inhibited extended form of the CaMKII\(\alpha\) holoenzyme [35] because homo-FRET between Venus-tagged catalytic domains was observed in the V-CaMKII\(\alpha\)TT305/306DD construct (Fig 3C). As mentioned previously, the TT\(\text{305/306}\) to DD\(\text{305/306}\) mutations are expected to disrupt the compact auto-inhibited holoenzyme structure [35] but also block the activation of the enzyme by preventing Ca\(^{2+}\)/CaM binding [36]. Thus, our FPFA data supports the postulated Ca\(^{2+}\)/CaM accessible auto-inhibited holoenzyme structure, but with catalytic domains organized as pairs. Presumably, when catalytic domains un-lock from the association domain core they reorganize to form dimers.

Like in living cells, catalytic domain pairing was also observed in freshly prepared homogenates (Fig 3C). In contrast, the number of subunits in the holoenzymes was reduced from 11.9 ± 1.2 in cells to 10.2 ± 1.3 in homogenates. Thus, catalytic domain pairing as observed by homo-FRET occurs in cells as well as in vitro and persists despite a reduction in holoenzyme subunits stoichiometry. This would be expected if the reduction in subunits represent a loss of a pair of subunits from the holoenzyme with their paired catalytic domains, rather than a loss of two independent subunits. This type of paired subunit organization has been previously proposed based on subunit stoichiometry distributions from electron microscopy imaging of native holoenzyme [24].

An implicit assumption of using brightness analysis to measure the number of subunits in a CaMKII holoenzyme is that only subunits tagged with a fluorophore are incorporated into a complex. This assumption may not always be warranted, particularly for cells that express a high level of non-tagged endogenous subunits. Under these conditions it is prudent to interpret normalized brightness as a lower limit for the number of subunits in a complex. In these studies for Venus-CaMKII\(\alpha\) expression in HEK cells this is unlikely to be a problem because the CaMKII subunit stoichiometry predicted by FPFA analysis was in good agreement with estimates based on non-tagged native holoenzyme [23,24]. Nonetheless, a conservative interpretation of our data would conclude that in cells CaMKII\(\alpha\) holoenzyme typically has ≥12 subunits.

The impact of fluorophore-tagging must be considered when interpreting structural findings based on fluorescence. Here FPFA detected 2 distinct V-CaMKII\(\alpha\) protein interactions which were previously observed using untagged-CaMKII. First, in cells V-CaMKII\(\alpha\) subunits assembled to form a dodecamer (Fig 4B), a value similar to in vitro estimates using native holoenzyme [23,24], and in vivo measurements based on GFP-tagged CaMKII\(\alpha\) translocation [25]. Second, catalytic domains were arranged as pairs in the holoenzyme (Figs 3C & 4A), consistent with homo-FRET [10], hetero-FRET (Fig 3D), and ultra-centrifugation studies using untagged catalytic domains [30]. Thus, it is unlikely that Venus-tagging caused or significantly perturbed these interactions.

It is worth comparing the relative merits of CaMKII\(\alpha\) holoenzyme structural studies inside living cells based on considering only hetero-FRET data (Fig 3D), only homo-FRET data ([10], and Fig 4A), as compared to considering both homo-FRET and brightness analysis from FPFA (Fig 4A & B). Hetero-FRET analysis (Fig 3D) readily demonstrated that fluorescent protein-tagged CaMKII\(\alpha\) catalytic domains were in close proximity. The ensemble FRET efficiency for the 171 cells transfected with C-CaMKII\(\alpha\) and V-CaMKII\(\alpha\) was 25.2 ± 7.9%, significantly higher than our negative control. Nonetheless, FRET efficiencies ranging from 0 to 40% were observed depending on the relative expression of Cerulean- and Venus-tagged CaMKII\(\alpha\) subunits. Thus, if the fraction donor were not simultaneously measured, as is often the case for many FRET approaches, interpretation of this hetero-FRET data would be problematic. For example, our interpretation of CaMKII\(\alpha\) hetero-FRET in cells, as energy transfer between paired donors and acceptors with a FRET efficiency of 39.4 ± 0.8% was based on modeling how the FRET efficiency changed from cell to cell as a function of the fraction donor. While we feel this interpretation is compelling, we also note that these experiments cannot differentiate between holoenzyme structures having a single set of paired subunits, and various other possible configurations comprised of multiple sets of paired
subunits, and even our negative control showed a small amount of FRET, perhaps as a result of over expression.

Homo-FRET analysis (Fig 4A) could also readily detect energy transfer between Venus-tagged catalytic domains, again indicating that in cells Venus-tagged catalytic domains are in close proximity to other Venus-tagged catalytic domains. Based on the amplitude of V-CaMKIIz’s fast anisotropy decay component, compared to the anisotropy decays of Venus monomer and trimer controls, it is most likely that V-CaMKIIz’s catalytic domains are organized as pairs [10]. Like the previous hetero-FRET analysis, homo-FRET analysis alone cannot differentiate between a holoenzyme structure having a single set of paired subunits, and configurations comprised of multiple sets of paired subunits. One advantage of homo-FRET over hetero-FRET in that only a single fluorescent protein tagged CaMKII subunit, V-CaMKIIz, was needed for these measurements. Thus, possible errors related to the relative expression of donor and acceptor-tagged subunits are eliminated.

Both homo- and hetero-FRET analyses are susceptible to false positive FRET signals due to over expression of fluorophore-tagged subunits in cells. One advantage of homo-FRET analysis of data collected by FPFA is that the possibility of false positives is almost completely eliminated. For soluble fluorophores, non-specific FRET typically requires concentrations greater than 1 mM [4,52]. With the FPFA instrumental design reported here this would correspond to having in excess of 210,000 Venus molecules in the 0.35 fl observation volume of our microscope. At these high concentrations fluorescence intensity fluctuations (due to individual fluorophores moving in and out of the observation volume) are too small relative to the average fluorescence intensity to be measured [17]. For this reason, FCS and FPFA measurements, by necessity, are limited to samples with concentrations much smaller. For example, in figure 1D the number of Venus monomer and trimer controls, it is much less than 1 mM (~210 molecules). Furthermore, to avoid TCSPC pile-up errors [13] in time-resolved anisotropy measurements, samples for FPFA was further restricted to those with count-rate less than 100,000 cps. Within these limits, the maximum number of Venus molecules in the observation volume that we could measure is ~120 (~570 nM), but was typically much smaller. For example, in figure 1D the number of Venus molecules in the observation volume was ~65. Thus, observing non-specific FRET in a FPFA experiment is highly unlikely. Another major advantage of FPFA analysis over both hetero- and homo-FRET analysis is that FPFA simultaneously measures normalized brightness. Thus we can deduce that the dimeric homo-FRET signal we observed in cells for V-CaMKIIz (Fig 4A) was coming from protein complexes having on average at least 12 subunits (Fig 4B). This observation eliminates the possibility that tagging CaMKIIz on its N-terminus (catalytic domain) prevented holoenzyme oligomerization via its association domain. We conclude that the advantages of being able to simultaneously measuring homo-FRET, brightness, and correlation time to study protein interactions in living cells justifies the use of FPFA over other FRET approaches, and in addition to being a useful tool for understanding the structure of the CaMKII holoenzyme; FPFA can also be used to study protein interactions in other complexes as well.

Materials and Methods

Cell Culture, Transfection and Homogenate Preparation

HEK 293 cells (ATCC) were cultured as a monolayer in a T-75 Flask (Corning) in a humidified incubator containing 5% CO2 in air at 37°C in High Glucose DMEM media containing L-Glutamine, sodium pyruvate and 10% fetal bovine serum (all from Gibco). A day prior to FPFA measurement, cells were resuspended using TrypLE Express (Invitrogen) and washed with DPBS (with calcium and magnesium; Mediatech). For in vitro measurements, plasmid DNA encoding Venus-tagged constructs (typically 1 μg/250,000 cells) were transfected into the cells using electroporation (Digital Bio/BTX MicroPorator). Transfected cells were plated on 60 mm culture dishes (Falcon) and incubated overnight. On the following day, cells were harvested and lysed using passive lysis buffer (Promega). Homogenates were centrifuged at 100,000 x g for 1 hour to remove membranes and particulate matter. Supernatants were diluted for FPFA to yield a photon count rate between ~20,000 cps (>25 x the dark count rate) and <100,000 cps (to avoid TCSPC pile-up artifacts [13]). The clarified homogenates were then further loaded into 35 nm glass bottom dishes (MatTek) and micro- and macro-times were measured by FPFA the same day. For live-cell measurements, cells were transfected with RNA encoding Venus-tagged constructs since the amount of protein expression after 24 hours could be controlled by titrating the amount of RNA used during transfection, typically around 500 μg of RNA/250,000 cells was used. Transfected cells were plated on 35 mm glass bottom dishes and incubated overnight in phenol-red free DMEM media and FPFA was measured the next day.

Molecular Biology and Immunoblot Analysis

DNA clones encoding V1 (Addgene ID 277794), V2/V3/V (Addgene ID 29423), V3 (Addgene ID 27814), V4 (Addgene ID 29425), V5 (Addgene ID 29426), V6 (Addgene ID 27813), V17V (Addgene ID 29424), V32V (Addgene ID 29561), and V-CaMKIIz (Addgene ID 29428) are available from Addgene (http://www.addgene.org/Steven_Vogel/). DNA clones encoding V-CaMKIIz [TT305/306DD], and C-CaMKIIz were generated as follows: Sense (5\'-GGAGGCATCTCCGAGCA-CATGCTGGCCACCGAG-3\') and antisense (5\'-CCTTGGTGCCAGCATGTCGTGAGATGGCTCC-3\') primers were used to perform site directed mutagenesis PCR (as previously described [10]) to generate V-CaMKIIz[TT305/306DD]. Note that the mutated bases are underlined in the primer sequences. C-CaMKIIz was generated by excising the open reading frame of mouse CaMKIIz from V-CaMKIIz (Addgene ID 29428) and inserting it into a Salmon/HindI digested Cereulan C1 (Addgene ID 27796). The resulting C-CaMKIIz construct was confirmed by sequencing. For RNA transfection, DNA encoding full length V-CaMKIIz was excised and inserted into the P22 vector [50]. This clone was digested using Not-I and the linearized plasmid was used to generate cRNA using an mMachine mMessage RNA kit (Ambion).

Immuno blot analysis. HEK 293 cells were transfected with plasmid DNA encoding the V1–V6 constructs. Cells (80–100% confluent) expressing Venus concatamers were scrapped off the dish, washed twice with PBS, and pelleted. The pellet was then lysed in denaturing Laemmli buffer (BioRad) with 1% SDS and heated to 90°C for 10 mins), and 20 μl of the solution was loaded into a precast gradient SDS PAGE gel (4–20%; BioRad). Immunoblotting with a GFP specific antibody (NeuroMab Anti-GFP N86/8) was performed as previously described [42].

Fluorescence Polarization and Fluctuation Analysis

Experimental Setup

An 80-MHz, 200-fs mode locked Ti:sapphire laser (Coherent Chameleon Ultra-2) was tuned to 950 nm to provide pulsed two-photon excitation. The power of the laser beam was adjusted using a variable attenuator consisted of a half-wave plate and a Glan-
laser polarizer (Thorlabs). The excitation beam was filtered and expanded using a spatial filter system (KT310, Thorlabs), and then passed through a near-IR linear polarizer (100,000:1 extinction ratio, Thorlabs) to enter the back port of a Zeiss Axio Observer microscope. A multiphoton short-pass dichroic beamsplitter (FF670-SD01-25×36, Semrock) was used to reflect the excitation beam to a Zeiss 63×1.2 NA water objective (back aperture slightly overfilled) that focused the beam to a diffraction-limited spot (~0.4 μm in diameter). Fluorescence from the observation volume in the sample was filtered through a BG39 filter (to block residual near-IR photons), a high throughput band-pass filter (FF01-540/50–25, Semrock), and then guided to a polarizing beam splitter (Thorlabs) that was augmented with two orthogonally oriented linear polarizers (Thorlabs) to increase the extinction ratio. At the beam splitter, parallel and perpendicular emitted photons were separated and focused onto two HPM-100-40 hybrid detectors (Becker & Hickl). The dark count rate for these detectors was typically 400–750 cps at room temperature. Photons detected by each detector were processed by a SPC-132 TCSPC card (Becker & Hickl). The SPC-132 recorded both micro- and macro-time for each parallel and perpendicular detected photon. For synchronization between excitation pulses and detected photons, a small fraction of the excitation beam was extracted and focused onto a fast photodiode that was powered by battery to avoid crosstalk. Note that all optics used in the excitation pathway was selected to minimize group delay dispersion.

SPCM software (Becker & Hickl) running in FIFO mode was used for data acquisition and to calculate time-resolved fluorescence and auto-/cross-correlation functions from measured micro- and macro-time data, respectively. Excitation power was kept low (typically 10.2 mW in vitro and 6 mW in living cells) to avoid bleaching during acquisition (~120 s in vitro and 20 s in living cells). For each homogenate, three to five replicate measurements were performed and these were averaged for each point. For live-cell measurements ten to fifteen replicate measurements were averaged for each cell. All measurements were performed at room temperature.

Time Resolved Anisotropy and Lifetime Analysis

Time-resolved anisotropy was calculated based on fluorescence decay of parallel and perpendicular channels using the following equation [10,11]:

\[
r(t) = \frac{I_\parallel(t) - g \cdot I_\perp(t)}{I_\parallel(t) + 2g \cdot I_\perp(t)}
\]

where \(I_\parallel(t)\) and \(I_\perp(t)\) are fluorescence intensity of parallel and perpendicular channels (dark noise subtracted) respectively, and \(g\) is the instrument correction factor which for our microscope had a value of 1 as determined by calibration using fluorescein tail fitting.

The total time-dependent fluorescence intensity decay or lifetime was calculated using the following relationship [13]:

\[
I(t) = I_\parallel(t) + 2g \cdot I_\perp(t)
\]

Polarized Fluorescence Fluctuation Analysis

For FPFA, a cross-correlation curve is fitted to a single component three-dimensional Gaussian function [17] \(G(t)\) to estimate the values \(\langle N \rangle\), the average number of fluorescent molecules in the excitation volume, and \(\tau_D\), the correlation time, the average time that a molecule spends in the detection volume:

\[
G(t) = \frac{1}{\langle N \rangle} \cdot \frac{1}{1 + (\frac{t}{\tau_D})^2} \left(\frac{1}{\sqrt{1 + (\omega/z)^2 (t/\tau_D)}}\right)
\]

Where \(\omega\) and \(z\) are the radial and axial beam waists respectively, and the constant \(\gamma\) has a value of 0.35 for a two-photon three-dimensional Gaussian PSF [17].

The molecular brightness \(\eta\) is the average number of photon emitted per second per molecule (cpsm):

\[
\eta = \frac{\langle k \rangle}{\langle N \rangle}
\]

where \(\langle k \rangle\) is the average photon count rate recorded during data acquisition.

The normalized brightness, \(\rho\), of a Venus-tagged protein complex was determined by dividing the molecular brightness \(\eta_{\text{complex}}\) of a complex composed of Venus-tagged subunits, by the molecular brightness of a Venus monomer \(\eta_{\text{Venus}}\):

\[
\rho = \frac{\eta_{\text{complex}}}{\eta_{\text{Venus}}}
\]

Note that \(\eta_{\text{complex}}\) and \(\eta_{\text{Venus}}\) should be measured using similar conditions (primarily using the same laser excitation power, filters, and optics), and that \(\eta_{\text{Venus}}\) can be measured by running a Venus monomer control. In cells, where auto-fluorescence was an issue, the normalized brightness was calculated using the measured slope \(m\) and y-intercept \(b\) of the linear portion of the \(V1–V6\) brightness curve:

\[
\rho = \frac{\eta_{\text{complex}}}{m} - \frac{b}{m}
\]

With two-photon excitation the relationship between correlation time \(\tau_D\) and the diffusion coefficient \(D\) is given by [53]:

\[
\tau_D = \frac{\omega^2}{8D}
\]

Calibration

The instrument correction factor \(g\) for calculating time-resolved anisotropy (equation 1) was measured using tail-fitting [11] of fluorescein samples and found to be 1. At high pH, fluorescein has a constant quantum yield and its diffusion coefficient \(D\) is 300 μm²/s at room temperature [17]. Thus, equation 7 can be used to estimate the value of \(\omega\) (at a specific excitation power) by measuring fluorescein’s correlation time (at the same power). For example, with \(D = 300 \text{ μm}^2/\text{s}\), and a measured correlation times of \(74.8 ± 9.3 \mu\text{s}\) \((n = 5)\), the value of \(\omega\) was 424 ± 26 nm with 10.2 mW excitation power (at 950 nm). The ratio \(\omega/z\) (equations 3 and 4) was measured by global fitting (to equation 3) of cross-correlation curves obtained from known dilutions of fluorescein. In this calibration it is assumed that with constant excitation power for all fluorescein dilutions, only the value \(\langle N \rangle\) will change with dilution. At 10.2 mW excitation power the \(\omega/z\) ratio was \(0.13 ± 0.00\), and taken together with our estimate for \(\omega\) predicts a \(z\) value of 2.8 ± 0.2 μm. The validity of this calibration procedure was confirmed by measuring the diffusion coefficient of Venus monomers under identical conditions. Using \(\omega = 424\) nm, and \(\omega/z = 0.15\) the measured correlation time for Venus monomers with
10.2 mW excitation power was 345±16 μs (n = 3). This, corresponds to a diffusion coefficient for Venus monomers in solution of 65±3 μm²/s (n = 3) in good agreement with the diffusion coefficient measured for GFP [51]. Because fluorescein was poorly excited with 6 mW excitation power at 950 nm, calibration for live cell FPFA used a slightly different procedure. Global fitting of cross-correlation curves obtained from dilutions of Venus monomers were used to measure the ratio ω/z (0.10±0.00), as well as the correlation time of Venus monomers (299±65 μs, n = 3). These values indicated that with 6 mW excitation power ω = 367±63 nm and z = 3.7±0.9 nm. Note that these values had large errors because of the low excitation power. We used ω and z values measured at 6 and 10.2 mW excitation power to determine if the two-photon observation volume changed appreciably with increased excitation power. The two-photon observation volume (V) at any specific excitation power can be calculated using the following equation [17]:

\[ V = \pi \sigma^2 z \frac{m}{8} \]  

(8)

Accordingly, the observation volume with 10.2 mW excitation power was 0.35 μl, only slightly larger than the volume measured at lower power (6 mW, 0.34 μl). Note that these volumes, and the value of <N> from a FPFA measurement can be used to calculate the concentration of Venus or of Venus-tagged protein complexes, a key factor for determining if non-specific FRET (due to molecular crowding) can occur.

**Hetero-FRET Measurements**

DNA encoding CaMKIIα tagged on the catalytic domain with Cerulean [44] acting as a FRET donor, and Venus [38] as a FRET acceptor were transfected into HeLa cells (ATCC). Cells transfected with Cerulean-CaMKIIα & free Venus monomers were used as a negative FRET control, and cells transfected with C5V [45] was used as a positive control. Images were acquired on an automated wide-field microscope and FRET analysis was performed using the E-FRET method [48] calibrated with Cerulean and Venus FRET standards [45]. Each point is the average FRET efficiency and fraction donor value measured for an individual cell. Dashed lines are linear fits. A linear relationship between the observed FRET efficiency (Eobs) when plotted as a function of the fraction donor (p) indicates that FRET is only occurring between 1 donor and 1 acceptor [49]. This linear relationship can be understood as follows: If CaMKII subunits form random pairs based on their abundance, then if p is the fraction of Cerulean-tagged CaMKII subunits in the population, then (1-p) will be the fraction of Venus-tagged CaMKII subunits in the population. The random pairing of Cerulean- and Venus-tagged catalytic domains can be modeled using a binomial distribution. In a population the fraction of Cerulean-tagged CaMKII subunits paired with other Cerulean-tagged CaMKII subunits will be p², and each of the 2 Cerulean molecules in these pairings will have a FRET efficiency of zero because they are both not paired with a Venus acceptor. Similarly, the fraction of Venus-tagged CaMKII subunits paired with another Venus-tagged CaMKII subunits will be (1-p)², but because these pairs lack donors, they have no direct impact on the FRET efficiency of the population. Finally, the fraction of Cerulean-tagged CaMKII subunits paired with Venus-tagged CaMKII subunits will be 2p(1-p), and because each has a single donor (and acceptor), they will each have a FRET efficiency of E. Thus the observed ensemble FRET efficiency for the population Eobs, as a function of the fraction donor (p) will be:

\[ E_{\text{obs}}(p) = \frac{2p(1-p)E}{2p^2 + 2p(1-p)} = (1-p)E \]  

(9)

Note, that Eobs is a linear function of p, and that the slope and y-intercept (when p = 0) are both equal to the FRET efficiency between a single donor and acceptor.

**The Impact of Autofluorescence on Brightness Measurements for Live Cell FPFA**

For live cell FCS measurements, auto-fluorescence and other background light sources can impact on brightness measurements [16,17,18,51]. In addition to the fluorophore of interest, endogenous fluorophores can also emit photons resulting in an altered apparent brightness. This problem can also occur for FPFA based brightness measurements. Corrections for two different types of background signals have been described, these are: 1. Background signals that only alter the photon-count rate, and 2. Background fluorescence from endogenous fluorophores that fluctuate. For background signals that only alter the photon-count rate the apparent molecular brightness (ηapp) of the sample is [16,18]:

\[ \eta_{\text{app}} = \eta \left( \frac{\langle k \rangle - \langle k_B \rangle}{\langle k \rangle} \right)^2 \]  

(10)

Where η is the brightness of the exogenous fluorophore, <k> is the average measured count rate (sample plus background), and <kB> is the average measured count rate of the background. The apparent brightness of a sample with fluctuating endogenous fluorophores is [18,51]:

\[ \eta_{\text{app}} = \frac{\eta^2 \langle N \rangle + \sum_k \eta_k^2 \langle N_k \rangle}{\eta \langle N \rangle + \sum_k \eta_k \langle N_k \rangle} \]  

(11)

Where η and <N> are the molecular brightness and the average number of exogenous fluorophore molecules in the observation volume, and ηB and <NB> are the molecular brightness and the average number of molecules in the observation volume of the bth fluorescence background species. In living cells it is possible that both of these types of background signals can adversely contribute to the apparent brightness. Furthermore, the relative impact of these types of background signals on the apparent brightness is rarely known. Moreover, many of the factors in equation 11 are either difficult to measure or are themselves unknown, and it is worth noting that the validity of these corrections has not been rigorously tested. An alternative empirical strategy for interpreting the apparent brightness of a complex composed of FP-tagged subunits, in terms of subunit stoichiometry, is to compare an unknown samples brightness to the measured brightness of a series of FP-concatemers expressed in the same cell type, such as the use of the V1–V6 series used in this study. The underlying assumption for this calibration strategy is that the brightness, abundance and fluctuation behavior of the sources of cellular background signals are not altered by the expression of different FP-tagged constructs. Under ideal conditions, experimental samples will have brightness values falling within the range of a control set (1–6 subunits in this instance). Under these conditions control brightness values can be used directly as a standard curve to interpret the brightness of an unknown. For samples with an apparent brightness greater than the largest concatemer in a control set (in this example those with
>6 subunits) the subunit stoichiometry of the sample can be estimated by extrapolating brightness values based on the standard brightness curve. While extrapolations beyond control values is not ideal, by using proper error propagation [54] errors in interpretation can be minimized.

Curve Fitting and Statistics

IGOR Pro software (Vs 6.22) was used for standard and global fitting of time-resolved anisotropy, cross-correlation curves, and linear fits for brightness controls. GraphPad Prism 5 was used to calculate means and standard deviations (SD). Values are presented throughout the text as mean±SD, deviations of ±0.00 indicate a value of less than 0.005, while deviations of ±0.05 indicate a value of less than 0.005. GraphPad Prism was also used to calculate Student’s t-test, which was paired and two-tailed.

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Author Contributions

Conceived and designed the experiments: SSV TAN. Performed the experiments: TAN PS JVV SVK. Analyzed the data: SSV TAN. Contributed reagents/materials/analysis tools: JVV SVK. Wrote the paper: SSV TAN.

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