



Channelrhodopsin-2 Oligomerization in Cell Membrane Revealed by Photo-Activated Localization Microscopy

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Abstract: Microbial rhodopsins are retinal membrane proteins that found a broad application in optogenetics. The oligomeric state of rhodopsins is important for their functionality and stability. Of particular interest is the oligomeric state in the cellular native membrane environment. Fluorescence microscopy provides powerful tools to determine the oligomeric state of membrane proteins directly in cells. Among these methods is quantitative photoactivated localization microscopy (qPALM) allowing the investigation of molecular organization at the level of single protein clusters. Here, we apply qPALM to investigate the oligomeric state of the first and most used optogenetic tool Channelrhodopsin-2 (ChR2) in the plasma membrane of eukaryotic cells. ChR2 appeared predominantly as a dimer in the cell membrane and did not form higher oligomers. The disulfide bonds between Cys34 and Cys36 of adjacent ChR2 monomers were not required for dimer formation and mutations disrupting these bonds resulted in only partial monomerization of ChR2. The monomeric fraction increased when the total concentration of mutant ChR2 in the membrane was low. The dissociation constant was estimated for this partially monomerized mutant ChR2 as 2.2 ± 0.9 proteins/ μm^2 . Our findings are important for understanding the mechanistic basis of ChR2 activity as well as for improving existing and developing future optogenetic tools.

Microbial rhodopsins constitute a large group of light-sensitive proteins with seven-transmembrane α -helices found in pro- and eukaryotic microorganisms. These proteins harbor a retinal cofactor which is photoisomerized upon illumination. This process is the primary event underlying the diverse functions of retinal proteins as light-driven pumps, light-gated channels and photoreceptors. Expression

of microbial rhodopsins in neurons enables to control the nerve cells' membrane potential by light with high temporal and spatial resolution.^[1] This became the basis for the emergence and development of a technique called optogenetics.^[2] It allows using light to control and study cellular activities in isolated neurons as well as complex neural systems such as the brain of living animals^[3,4].

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Promising biomedical applications of rhodopsins in optogenetics are aimed to restore vision,^[5] hearing^[6] and memory.^[7]

Microbial rhodopsins can form stable oligomers of various stoichiometry (from dimers to hexamers)^[8] that are relevant to their properties and function. The best-known example is bacteriorhodopsin (BR)—the archetypal light-driven proton pump of *Halobacterium salinarum*. BR forms trimers packed into a two-dimensional lattice in the native purple membrane.^[9] The trimeric form of BR has significantly higher proton pumping efficiency^[10] and is important for its higher thermal and photostability^[11–13] relative to the monomeric state. Other examples of oligomerization-dependent functions include Na⁺-pumping by pentameric *Krokinobacter eikastus* rhodopsin 2 (KR2),^[14] proton pumping by hexameric proteorhodopsin (PR)^[15,16] and ion channeling by trimeric ChRmine^[17] or pentameric Organic Lake Phycodnavirus rhodopsin II (OLPVR II).^[16] In most cases, the oligomeric state of microbial rhodopsins was determined by atomic force microscopy,^[18,19] circular dichroism spectroscopy,^[19–21] electron microscopy,^[22,23] crosslinking^[15,16] and X-ray crystallography.^[24–26] These methods require sophisticated sample preparation that in many cases may imprint a non-natural oligomeric state to the microbial rhodopsins under study.^[8] Hence, although numerous new microbial rhodopsins from various organisms were recently identified, the information about their oligomeric states remained sparse and largely incomplete because of a lack of approaches to determine their oligomeric state in a native cell environment.

Channelrhodopsin-2 (ChR2) is a light-gated cation channel found in the unicellular green alga *Chlamydomonas reinhardtii*.^[27] It is the first^[1] and currently the most widely used rhodopsin in optogenetics.^[2] Mass spectrometry of protein detergent solution confirmed by electron crystallography of 2D-crystals showed that ChR2 forms dimers.^[22,28,29] These findings, however, do not allow concluding unequivocally that the dimer is ChR2's functional unit, since in both sample preparations ChR2 is placed in a non-natural environment. The X-ray diffraction crystal structure of ChR2 contains two symmetrical interprotomer disulfide bridges formed by Cys34 and Cys36.^[25] Such N-terminal extracellular cysteine residues are conserved in other structurally characterized dimeric channelrhodopsins.^[30–32] Mutations of Cys34/Cys36 do not destroy ChR2 dimers when purified protein samples were analyzed by EPR^[33,34] and no effect on ChR2-induced photocurrents was detected in ChR2-expressing oocytes.^[35] The oligomeric state of ChR2 and its mutants have not been investigated in the plasma membrane of a cell so far and therefore the nature of its functional unit remains unclear.

Fluorescence microscopy provides a variety of methods to analyze membrane protein oligomerization in an intact cell environment. These methods include Förster resonance electron transfer (FRET),^[36] number & brightness analysis (N&B),^[37,38] spatial intensity distribution analysis (SpIDA),^[39,40] fluorescence correlation spectroscopy (FCS),^[41,42] stepwise fluorescence photobleaching,^[43,44] single-molecule tracking,^[45,46] point accumulation in nanoscale

topography (PAINT),^[47,48] and photoactivated localization microscopy (PALM).^[49,50]

Typical cell surface density of ChR2 used in optogenetics can be estimated from photocurrent noise in patch clamp experiment as ≈ 200 proteins/ μm^2 ^[51] that also corresponds to those obtained by direct counting on the freeze-fracture electron microscopy images.^[52] This high density of protein does not allow to spatially resolve individual ChR2 oligomers via diffraction-limited fluorescence microscopy. Single-molecule localization microscopy (SMLM)^[53] overcomes this resolution barrier, and in addition to spatially resolving single protein clusters in intact cells, provides information on molecule numbers and protein oligomerization within single protein clusters.^[54] Here, we use PALM (i.e. SMLM in combination with photoswitchable fluorescent proteins), where sparsity of active emitters is controlled via photo-conversion and photobleaching of fluorescent proteins genetically fused to the protein of interest (Figure 1a). We use a kinetics-based analysis, quantitative PALM (qPALM), which relates the number of single-molecule emission events to molecule numbers^[55] (Figure 1b,c,d) and informs on protein oligomerization.^[56] In qPALM, a target protein is genetically fused with a photoconvertible fluorescent protein, which emits a limited number of fluorescence bursts after it is photoconverted and before it is irreversibly photobleached in the course of the experiment. Protein clusters containing several fluorescent proteins emit, on average, more bursts than clusters containing a monomer. The accurate analysis takes into account two challenges when using fluorescent proteins: first, some fluorescent proteins are not detected due to e.g. incomplete maturation, which results in undercounting; second, some fluorescent proteins produce more than one fluorescence burst, due to reversible transition into long-lived non-fluorescent states, which results in overcounting.^[57] Quantitative PALM analysis inherently corrects for both incomplete detection and multiple emission events.^[55] In a best-practice procedure, the experimental parameters describing the detection efficiency and the degree of multi-signal detection are determined from experiments with reference proteins with known oligomeric states.^[50] Importantly, qPALM enables the determination of the fractions of various oligomers in a mixture via fitting the histogram for the numbers of fluorescence bursts in protein clusters by a weighted sum of reference distributions.^[50,55,58]

In this work, we determined the oligomeric state of ChR2 in the intact plasma membrane of fixed human embryonic kidney (HEK293) cells using qPALM. We compare ChR2 with well-characterized monomeric and dimeric membrane protein reference samples and show that wild-type ChR2 (ChR2_{WT}) predominantly forms dimers in the plasma membrane. Furthermore, we analyze the role of two N-terminal cysteines Cys34 and Cys36 that stabilize the ChR2 dimer via disulfide bonds. We show that mutation of these cysteines partially decreases the fraction of ChR2 dimers while simultaneously increasing the fraction of ChR2 monomers. The extent of this effect depends on the ChR2 density in the plasma membrane that allows us to determine equilibrium dissociation constant of dimerization.

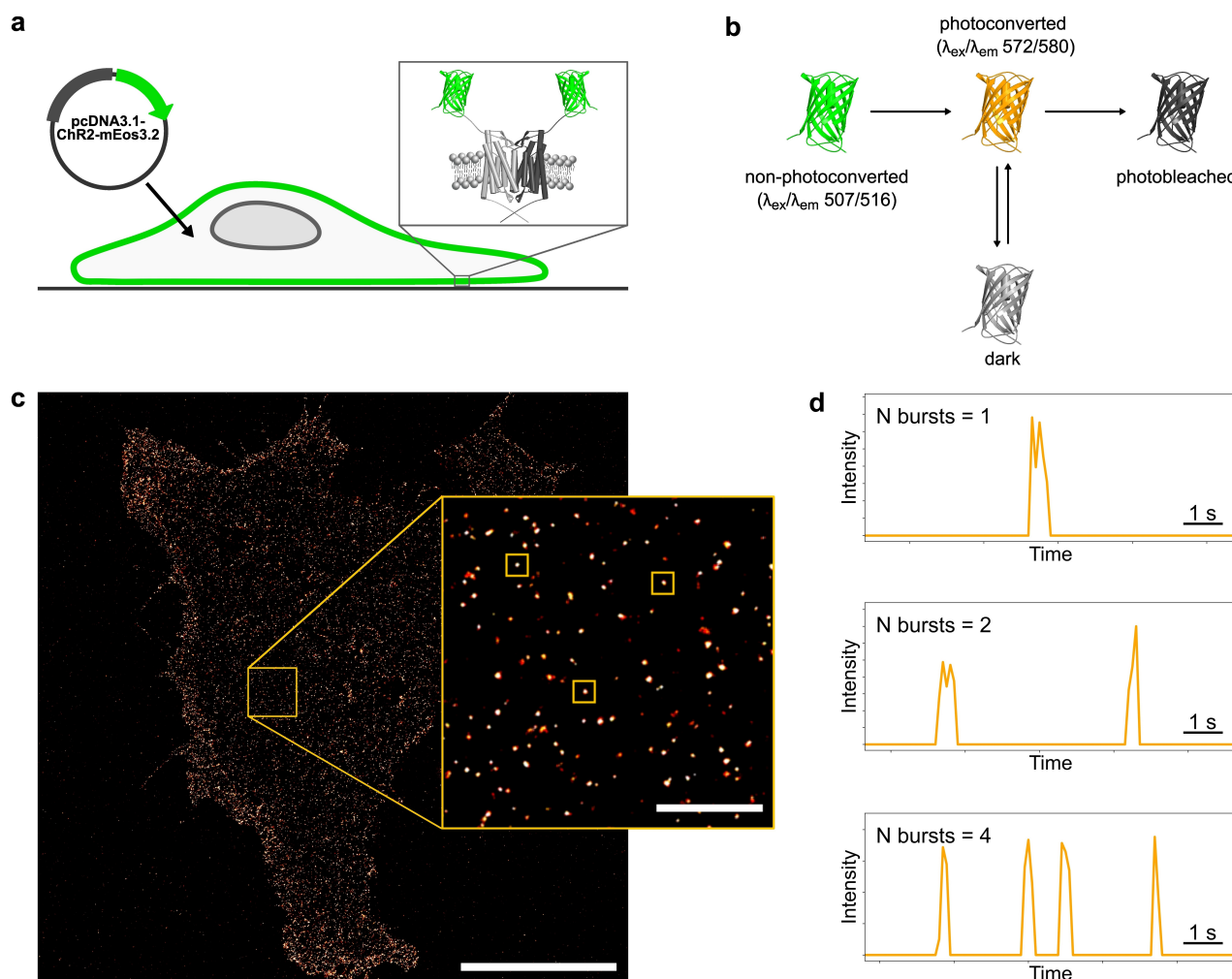


Figure 1. Illustration of general principles underlying quantitative PALM experiment. (a) HEK293 cells were transiently transfected with a pcDNA3.1 plasmid coding ChR2 fused to mEos3.2. The fluorescence from proteins in the basal plasma membrane was analyzed via quantitative PALM. (b) A basic 4-state photophysical model^[59] for the photoconvertible fluorescent protein mEos3.2. In a PALM experiment mEos3.2 first is irreversibly photoconverted by violet light (405 nm) from a shorter wavelength (non-photoconverted form; $\lambda_{\text{ex}}/\lambda_{\text{em}}$ (nm) 507/516) to a longer wavelength emission state (photoconverted form; $\lambda_{\text{ex}}/\lambda_{\text{em}}$ (nm) 572/580). Photoconverted mEos3.2 can photoblink, i.e. reversibly switch between a non-emissive dark state and the emissive photoconverted state until irreversible photobleaching. (c) An example of a super-resolved cell membrane image obtained from a PALM experiment. PALM experiments are performed on cells expressing membrane proteins fused with mEos3.2 (ChR2_{WT}-mEos3.2 in this particular case). The fluorescence of a single photoconverted mEos3.2 is detected in TIRF mode allowing high-precision visualization of the proteins localized in the basal plasma membrane; scale bar 10 μm . Zoomed inset: individual protein clusters can be selected and the number of fluorescence bursts in individual clusters can be determined; scale bar 1 μm . (d) Examples of fluorescence time courses measured in individual clusters are provided to reveal the number of bursts. All bursts emitted by selected clusters during the entire data acquisition are contained within the shown time intervals. Before the first burst is emitted, all mEos3.2 molecules in the cluster remain in non-photoconverted state, and after the last burst, all mEos3.2 molecules are photobleached.

Membrane proteins with known oligomeric states were used to calibrate ChR2 molecular counting. As references, we chose monomeric β_1 -adrenergic receptor ($\beta_1\text{AR}$) and constitutively dimeric CD28, which have been used as standard controls in previous studies.^[60,61] Both proteins were tagged with photoconvertible fluorescent protein mEos3.2. It is a brighter and truly monomeric version of mEos2^[62] successfully used in other qPALM-based studies.^[63] Both reference constructs $\beta_1\text{AR}$ -mEos3.2 and CD28-mEos3.2 were expressed in human embryonic kidney 293 (HEK293) cells and mostly localized in the plasma membrane (Figure 2a). The HEK293 cell line was chosen as

a common system for optogenetic functional tests of microbial rhodopsins.^[31,64] Additionally, this cell line does not express either $\beta_1\text{AR}$ or CD28 endogenously.^[65] We used PALM images to visually separate individual protein oligomers on the cell membrane. On super-resolved images obtained from PALM, individual oligomers appear as bright round-shaped clusters of less than 100 nm in size (Figure 1c). We counted the number of fluorescence bursts of mEos3.2 in individual clusters. As expected, the clusters of dimeric CD28 contain more fluorescence bursts than monomeric $\beta_1\text{AR}$: the most frequent numbers of bursts in a cluster are two and one for CD28 and $\beta_1\text{AR}$, respectively (Figure 2b,c).

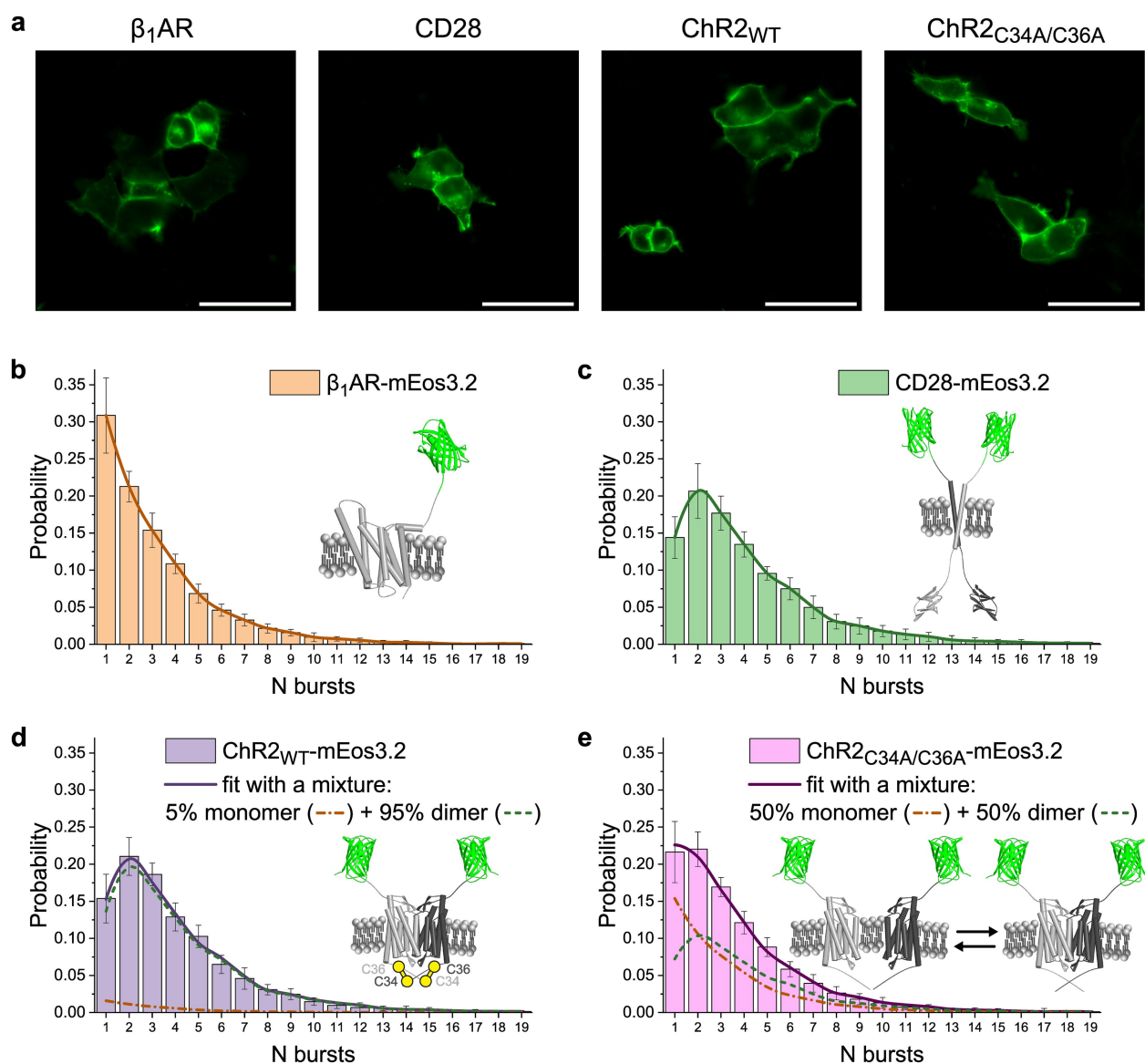


Figure 2. Confocal images of transfected cells and burst distributions for four proteins used in the study: β_1 AR, CD28, ChR2_{WT}, and ChR2_{C34A/C36A}. (a) Confocal images of HEK293 cells expressing membrane proteins fused with mEos3.2. The green form of mEos3.2 was excited with an argon ion laser at 488 nm. Scale bar 50 μ m. (b,c,d,e) Numbers of fluorescence bursts in individual clusters of membrane proteins fused with mEos3.2. Distributions were averaged over individual cells. Error bars represent the standard deviation. (b,c) Burst distributions obtained for β_1 AR-mEos3.2 (14 cells, 10,631 clusters) (b) and CD28-mEos3.2 (15 cells, 7,844 clusters) (c), which were used as monomeric and dimeric references, respectively. (d) The burst distribution obtained for ChR2_{WT}-mEos3.2 (12 cells, 7,510 clusters) was fitted to the weighted sum of reference distributions (weighted components are indicated by dashed lines). ChR2_{WT} showed predominant dimerization (95% of dimers) in the cell membrane. (e) The burst distribution obtained for ChR2_{C34A/C36A}-mEos3.2 (11 cells, 6,966 clusters) was fitted to the weighted sum of reference distributions (weighted components are indicated by dashed lines). The mutation of the two cysteines caused the partial decrease of ChR2 dimer fraction accompanied by the increase of monomer fraction (from 5% of monomers to 50% of monomers).

To determine the oligomerization of ChR2_{WT} in the plasma membrane we transfected HEK293 cells with a ChR2_{WT}-mEos3.2 construct. ChR2_{WT} was predominantly localized in the plasma membrane as it was the case for the reference membrane proteins β_1 AR and CD28 (Figure 2a). We visualized individual ChR2_{WT} clusters in the plasma membrane using PALM and counted the number of mEos3.2 fluorescence bursts in individual clusters (Figure 2d). The histogram of fluorescence bursts in individual

clusters for ChR2_{WT} appears very similar to the burst distribution of the dimer reference CD28. ChR2_{WT} burst distribution was well fitted by the weighted sum of distributions pre-recorded for monomeric β_1 AR and dimeric CD28, showing no evidence of higher oligomerization (no right shift of the distribution maximum compared to the distribution of dimeric CD28). According to this approximation, ChR2_{WT} is almost fully dimerized in the cell membrane (95% of dimers and 5% of monomers). To estimate the

significance of this approximation, we evaluated cell-to-cell variations. The data from individual cells were processed in the same manner (fitted by the weighted sum of reference distributions; Figures S1–S3). Despite slight variations in the dimer fraction between individual cells, statistical analysis showed no significant difference between Chr2_{WT} and reference dimer CD28 (Figure 3a).

Our PALM measurements confirm that most of the Chr2_{WT} in cell plasma forms dimers as it was expected from previous studies with purified Chr2, where stable dimers were observed via crystallography,^[22,25,29] mass-spectrometry^[28] and gel electrophoresis.^[22,28] While these earlier results pointed towards a strong dimerization tendency, the formation of higher Chr2 oligomers in the plasma membrane could not be excluded. The absence of higher oligomers in the cell membrane was confirmed by our qPALM experiments.

In a crystal structure,^[25] the Chr2 dimer is stabilized by two covalent interprotein bonds between cysteines at positions 34 and 36. To test the effect of these bonds on the oligomeric state, we prepared a double-mutant Chr2 version ($\text{Chr2}_{\text{C34A/C36A}}$), tagged with mEos3.2. The mutations of Chr2 did not affect the predominant localization of the protein in the plasma membrane (Figure 2a). As before, we counted the number of bursts in individual $\text{Chr2}_{\text{C34A/C36A}}$ clusters visualized with PALM. The burst distribution averaged across all measured cells showed a larger fraction of 1-burst clusters for $\text{Chr2}_{\text{C34A/C36A}}$ compared to Chr2_{WT} (Figure 2d, e). Again, the distribution for the number of fluorescence bursts was fitted to the weighted sum of reference monomeric and dimeric distributions (Figure 2e).

The dimeric fraction of Chr2 decreased from 95 % in Chr2_{WT} to 50 % in $\text{Chr2}_{\text{C34A/C36A}}$, while the monomer fraction increased concomitantly from 5 % to 50 %. The significance of this change in monomer-dimer equilibrium was estimated by evaluating cell-to-cell variations in our PALM measurements. To calculate the fraction of dimeric $\text{Chr2}_{\text{C34A/C36A}}$ in single cells we fitted the individual burst distributions per cell with the weighted sum of reference distributions (Figure S4, Figure 3a). The fraction of dimers was significantly lower in $\text{Chr2}_{\text{C34A/C36A}}$ than in Chr2_{WT} ($p < 0.05$).

In contrast to the stable dimerization of Chr2_{WT} , the mutant $\text{Chr2}_{\text{C34A/C36A}}$ was, on average, a mixture of 50 % monomers and 50 % dimers in the cell membrane. This result suggests that inter-protomer disulfide bonds are important but not necessary for the dimer formation. Based on the crystallographic structure,^[25] it can be inferred that the dimer is stabilized through a combination of backbone H-bonds (Y145/Y145' of TM4/TM4', A111'/Y109 from loop-region), specific interactions between side chains (Y184/Y85' of TM5/TM2', W118/T165' of TM3/TM4') as well as several water-mediated interactions. The mutation of these amino acids may contribute to a further increase of monomeric Chr2 fraction. In the previous EPR-based studies, purified Chr2 remained dimeric even when inter-protomer disulfide bonds were removed.^[33,34,66] The higher fraction of dimers in purified Chr2 compared to our study can be explained by high density of the protein due to overexpression in a heterologous expression system, altered lipid environment after solubilization and high protein concentrations required for EPR measurements.

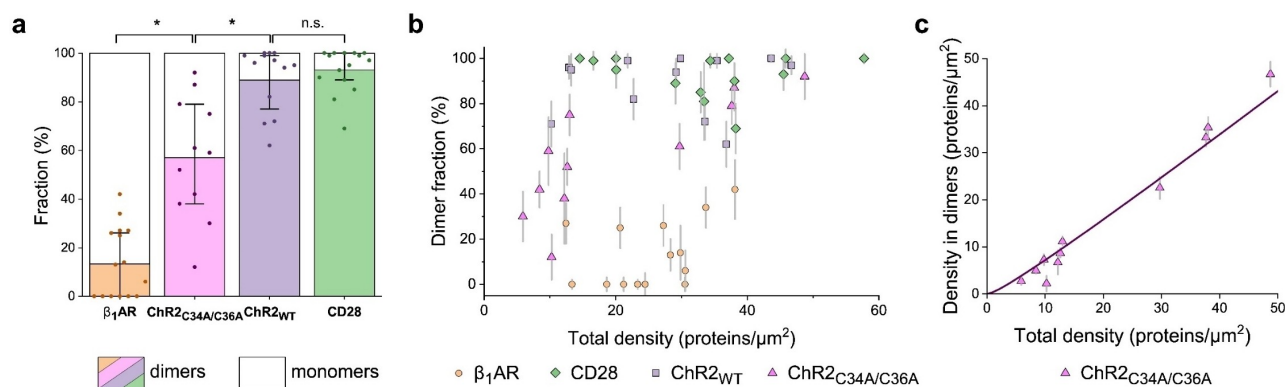


Figure 3. Analysis of cell-to-cell variations in PALM measurements. (a) Distributions of numbers of fluorescence bursts in individual clusters obtained from individual cells were fitted with the weighted sum of reference monomeric and dimeric distributions. Fractions of monomer and dimer clusters in single cells are shown as individual data points; the mean and the inter-quartile range are shown as bars with whiskers. Taking into account deviations caused by poorer statistics available from individual cells and their non-ideal homogeneity, Chr2_{WT} does not significantly differ from reference dimer CD28. At the same time, $\text{Chr2}_{\text{C34A/C36A}}$ showed the largest cell-to-cell variability and on average significantly differed both from reference monomer $\beta_1\text{AR}$ and Chr2_{WT} ; * $p < 0.05$ (Kruskal-Wallis test with post-hoc Dunn's test). (b) Dependence between the fraction of dimers and average protein density in the plasma membrane. Protein density is proportional to the number of bursts per unit area in the PALM experiment. From burst distributions for $\beta_1\text{AR}$ we estimated that one protein emits on average 3.25 bursts. Every point represents one cell expressing monomeric $\beta_1\text{AR}$ (orange circles), dimeric CD28 (green rhombuses), Chr2_{WT} (purple squares) or $\text{Chr2}_{\text{C34A/C36A}}$ (pink triangles). Errors are estimated with a bootstrapping algorithm. As well as reference proteins, Chr2_{WT} demonstrates no clear tendency to change oligomers ratio depending on protein density in the plasma membrane. The mutant $\text{Chr2}_{\text{C34A/C36A}}$ is likely to increase dimer fraction with increasing protein density. (c) For $\text{Chr2}_{\text{C34A/C36A}}$, the dependence of density of proteins residing in dimers on the total density of proteins is fitted to a binding model and the dissociation constant for $\text{Chr2}_{\text{C34A/C36A}}$ dimers ($K_D = 2.2 \pm 0.9$ proteins/ μm^2) is determined from the fit (solid line; see "Determination of dissociation constant K_D for $\text{Chr2}_{\text{C34A/C36A}}$ dimers" in Supporting Information).

In comparison to ChR2_{WT}, the mutant ChR2_{C34A/C36A} showed higher cell-to-cell variability of the dimeric fraction ranging from 12 % to 92 % in individual cells (Figure 3a). We proposed that in the absence of stabilizing disulfide bonds, dimerization of ChR2_{C34A/C36A} is transient and the fraction of dimers varies depending on the density of proteins per unit area of the plasma membrane. To test this hypothesis, we looked at the dependence of the dimer fraction on the density of proteins in the plasma membrane (Figure 3b). Since membrane proteins are stoichiometrically labeled, the density of fluorescence bursts on the membrane images is proportional to the protein density. Each protein emits, on average, 3.25 fluorescence bursts, as can be inferred from the blinking distribution of a reference monomer β_1 AR (Figure 2b). The mutant ChR2_{C34A/C36A} showed a statistically significant trend from predominant monomeric fraction at low protein densities towards predominant dimeric fraction at higher densities (Spearman's rank correlation coefficient $r = -0.80$, $p = 0.003$). On the contrary, there was no statistically significant correlation between the monomeric fraction and protein density for ChR2_{WT} or reference proteins ($p > 0.1$).

Without stabilizing inter-subunit disulfide bonds, the ratio between monomers and dimers in ChR2_{C34A/C36A} shifts towards pure monomers at low total protein densities in the membrane. This correlation between dimeric fraction and protein density suggests that a dynamic balance exists between monomers and transient dimers in ChR2_{C34A/C36A}. We determined the dissociation constant for ChR2_{C34A/C36A} dimers via fitting the dependence between density of proteins residing in dimers and total density of proteins with a binding model ($K_D = 2.2 \pm 0.9$ proteins/ μm^2 , Figure 3c, see "Determination of dissociation constant K_D for ChR2_{C34A/C36A} dimers" in Supporting Information). This finding implies that mutant ChR2_{C34A/C36A} must be predominantly dimeric at the expression levels common in optogenetic applications (~ 200 proteins/ μm^2 ,^[51]).

It is particularly intriguing whether the monomeric form of the ChR2_{C34A/C36A} is functional. Structural^[25] and mutagenesis^[35] studies of ChR2 showed that the monomer alone contains all necessary structural elements (gates and cavities) for ion channeling. It is therefore generally accepted that a monomer rather than a dimer is a functional unit of ChR2. However, even for the functional monomer, oligomerization may be a necessary requirement (similar to ChRmine^[17]) or at least significantly enhance its efficiency (similar to BR^[10]). Oligomerization might well be important for operative orientation of monomers in the lipid bilayer and stabilize their structural elements in functional states as it was shown for KR2.^[67] Although previous studies showed that ChR2 without inter-subunit disulfide bonds retains photocurrent activity in oocytes,^[35,66] the accurate correlative analysis of the ChR2 activity, density and oligomerization will be required to determine whether ChR2 monomers are functional.

The estimated dissociation constant for ChR2_{C34A/C36A} dimers is similar to those previously reported for other transiently dimerizing seven transmembrane helix proteins, class A G-protein coupled receptors (GPCRs).^[68–73] In

GPCRs, dimerization is tightly coupled to the functioning of receptors. Dimerization impacts the trafficking of receptors to the plasma membrane, regulates receptor internalization,^[74] affects ligand binding and downstream signaling.^[75,76] On the other side, the equilibrium between monomers and dimers depends on the activation state of the proteins.^[61,71] Given the structural similarity between retinal proteins and class A GPCRs and similarity between their dimer dissociation constants, it is reasonable to expect similar effects of dimerization for retinal proteins. Indeed, oligomerization was proposed to hinder the development of fused tandem rhodopsins due to undesired clustering of proteins.^[77] Multimerization properties were also suspected to completely deteriorate channelrhodopsin-derived tandem rhodopsins function in lysosomes.^[78] These findings highlight the importance of understanding the effects of dimerization and oligomerization when developing optogenetic tools.

Our study demonstrates the successful application of qPALM to determine the oligomeric state of rhodopsins in intact cells. This proof-of-concept study paves the way for further qPALM-based investigations of rhodopsins' oligomerization. In combination with functional tests, qPALM is potentially useful for mechanism-of-action studies that can reveal whether minimal functional units of specific rhodopsins are monomers or oligomers. Future qPALM studies can facilitate engineering of optogenetic tools whose functionality is affected by rhodopsin oligomerization.

Supporting Information

The authors give additional information on experimental procedures, DNA sequences and single cell burst distributions and have cited additional references within the Supporting Information.^[79–84]

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Conflict of interest

The authors declare no conflict of interest.

Data Availability Statement

The data that support the findings of this study are openly available in the public repository Zenodo at <https://doi.org/10.5281/zenodo.8349310>, <https://doi.org/10.5281/zenodo.8349425>, <https://doi.org/10.5281/zenodo.8349192>, <https://doi.org/10.5281/zenodo.8349352>.

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