Supporting Information

Monitoring the interaction of a single G-protein key binding site with rhodopsin disk membranes upon light-activation

Tai-Yang Kim, Hiroshi Uji-i, Martina Moeller, Benoit Muls, Johan Hofkens, and Ulrike Alexiev

Experimental procedures and data analysis

Sample preparation

A variant of a high-affinity analogue (VLEDLKSCGLF (1)) from the C-terminal segment of the transducin α-subunit, VLEDLKVGLF (pG T α), was synthesized at the Institute of Biochemistry, Universitaetsmedizin-Charite Berlin, and fluorescently labeled with Atto-647N-maleimide (Atto-Tec, Germany), yielding pG T α-F. Bovine retinas were purchased from J. A. Lawson Corp (Lincoln, NE). Rod outer segment disk membranes were prepared from bovine retinas and labeled with Atto-647N-maleimide at position C316 of rhodopsin essentially as described (2). Functional binding of pG T α-F was tested by the formation of additional Meta-II (Extra-Meta-II) in a concentration dependent manner at pH 8 and 3°C (Fig. S1a), as the pH and temperature dependent equilibrium between Meta-II and its precursor state is shifted towards Meta-II under these conditions (3). Flash spectroscopy was performed as described (2). The time constant for pG T α-F dissociation from light-activated disk membranes is 1-5 s as determined by double-time-resolved fluorescence spectroscopy (4).

For microscopy experiments layers of disk membranes were deposited at the surface of a glass cover slip in a buffer filled sample compartment. Individual Atto-647N labeled disk membranes are visible in the image presented in Figure S1c. The inset shows a fully covered glass surface, where individual membranes at the top layer are visible as brighter areas. For experiments with pG T α-F unlabeled
membranes were used. The Atto647N absorption band centered at 651±1 nm (Fig. S1b, arrow 2) is well 
separated from the absorption band of rhodopsin with \( \lambda_{\text{max}} = 498 \pm 1 \) nm (Fig. S1b, arrow 1). Thus, 
fluorescence excitation of pG\(_T\alpha\)-F at 633 nm (see below) should not result in accidental rhodopsin 
activation. The emission of pG\(_T\alpha\)-F with \( \lambda_{\text{max}} = 666 \pm 1 \) nm is marked by arrow 3 (Fig. S1b).

**Fluorescence microscopy**

Wide-field fluorescence microscopy was performed using an inverted epi-fluorescence microscope (IX-
71, Olympus) equipped with a cooled electron multiplying CCD camera (C9100-13, Hamamatsu). The 
fluorescently labeled rhodopsin disk membranes and pG\(_T\alpha\)-F were excited at 632.8 nm (He-Ne laser, 
Model 1145p, JDS Uniphase Co.). Light-activation of rhodopsin was carried out with a 2 ms light-pulse of 
532 nm from a solid-state laser (CDPS532M-50, JDS Uniphase Co.) equipped with a mechanical shutter 
(Uniblitz-LS6, Vincent Associates). Both laser lines were circularly polarized using a 1/4 waveplate and 
directed onto the sample through the same dichroic mirror (Z633RDC, Chroma Technology, Inc.). The 
wide-field illumination was carried out in the TIRF (total internal reflection fluorescence) mode by 
focusing the expanded laser beams onto the back-focal plane of the objective (60x TIRF objective (PLAN 
APO, TIRF/NA1.45)). Only those pG\(_T\alpha\)-F molecules which interact with the membrane were visible. The 
emission of the fluorophore was collected with the same objective and imaged through the dichroic mirror 
and a long path filter (HQ655LP). The collected fluorescence image was further magnified 3.3x using a 
projection lens, resulting in a maximum view of 41 x 41 \( \mu \text{m}^2 \) (80.8 x 80.8 nm\(^2 \)/ pixel). Movies were 
recorded with an integration time of 31 ms per frame (32 Hz). The photobleaching time constant of 
pG\(_T\alpha\)-F was ~10 s under the experimental conditions.

**Data analysis**

Fluorescence images were analyzed using self-written routines in MATLAB (Mathworks). After filtering, 
background subtraction, and applying the triangle algorithm for thresholding, the signals originating from 
individual molecules in the original image were fitted to a two-dimensional Gaussian profile (5). The two-
dimensional trajectories of molecules interacting with the membrane were constructed from the correlated images of identical molecules in subsequent observations. Tests with simulated data showed that the positional accuracy was ~18 nm (analyzed according to reference 6,7). Additional simulations were performed to discriminate between immobilization and confined diffusion for the slow component (subpopulation 1). The upper bound of positional accuracy for low fluorescent molecules was ~21 nm, the lower bound for bright fluorescent molecules was ~15 nm. Bright and low fluorescent molecules were found within both subpopulations.

Individual trajectories were analyzed by determining the mean square displacement $<r^2>$ as a function of elapsed time, where the lag time is $t_{lag} = n\Delta t$ with n being the frame index and $\Delta t$ denotes the frame interval. For the experiments presented in this study $\Delta t = 31$ ms. The mean square displacement of a set of coordinates $\{x_i, y_i\}$ was calculated as the average over all overlapping pairs

$$\left<r^2(t_{lag})\right> = \frac{1}{N-n} \sum_{i=1}^{N-n} [(x_{i+n} - x_i)^2 + (y_{i+n} - y_i)^2]$$

(1)

with $N$ being the total number of frames in the trajectory. For a two-dimensional random diffusion the diffusion coefficient $D$ is related to the mean square displacement $<r^2>$ by

$$\left<r^2(t_{lag})\right> = 4Dt_{lag}.$$

(2)

However, for non-homogeneous populations multiple diffusion coefficients can be introduced. This is better described by probability distribution functions (7-11). A characteristic radial probability distribution of displacements $r$, $p(r,t_{lag})$ for multiple subpopulations (8,9) is given by:

$$p(r,t_{lag})dr = \sum_{i=1}^{N_i} A_i \frac{2}{4D_i t_{lag}} e^{-\frac{r^2}{4D_i t_{lag}}} r dr$$

(3)
with $N_D$ being the number of subpopulations, and $A_i$ the fractional amplitudes with the corresponding diffusion constants $D_i$ and $4D_i t_{\text{lag}} = r_i^2(t_{\text{lag}})$.

Alternatively, a cumulative radial distribution function $P(r, t_{\text{lag}})$ can be obtained by integration of $p(r, t_{\text{lag}})$ over all $r$. For the case of $P(r^2, t_{\text{lag}})$ and $N_D=2$ the probability distribution is

$$P(r^2, t_{\text{lag}}) = 1 - \left( A \cdot e^{-r^2/r_1^2} + (1 - A) \cdot e^{-r^2/r_2^2} \right)$$  \hspace{1cm} (4)

with $A$ being the fractional amplitude (7).

For restricted diffusion, the confinement length $L$, i.e. diffusion within a square of side length $L$, and the initial diffusion constant $D_0$ can be extracted from the data using (12):

$$r^2(t_{\text{lag}}) = \frac{L^2}{3} \left( 1 - \exp \left( \frac{12D_0 t_{\text{lag}}}{L^2} \right) \right).$$  \hspace{1cm} (5)

**Structure presentation**

The structure shown in Figure 1C (main text) was prepared with VMD (13).

**References**


**Figures**

**Figure S1**

a) Extra-Meta-II formation at different concentrations of pGT\(\alpha\)-F. Inset: Half maximum concentration of pGT\(\alpha\)-F for the formation of Extra-Meta-II. The binding affinity of pGT\(\alpha\)-F is in the same range as the parent peptide (HAA) (3).

b) Absorption spectra of rhodopsin (arrow 1) and pGT\(\alpha\)-F (arrow 2) and emission spectrum of pGT\(\alpha\)-F (arrow 3).

c) Fluorescence image to visualize individual rhodopsin membranes. Scale bar: 2 \(\mu\)m. Inset: Glass surface fully covered with layers of rhodopsin membranes.
**Figure S2**
Control of instrument drift. Drift analysis on experimental data shows no indication for a systematic drift. Lines indicate particle movement (100x amplified) integrated over 310ms (10 frames). 5 consecutive groups are shown in different colors. A given color shows displacements occurring simultaneously.

![Image of particle movement](image1.png)

**Figure S3**
Residence times.

a) Residence time histogram of 2517 pG\(_{\alpha}\)-F molecules interacting with dark rhodopsin membranes.

b) Residence time histogram of 614 pG\(_{\alpha}\)-F molecules interacting with light-activated rhodopsin membranes.

![Histograms of residence times](image2.png)