

Geometry plays a role in GPCR transmembrane signaling

September 26 2012



In a recent *JGP* article, researchers developed a novel approach to study GPCR movement within a living cell. This image shows the grid used for modeling diffusion, with tracings from the image of a rod outer segment indicating disc perimeter (white) and incisures (orange). Credit: Najafi, M., et al. 2012. *J. Gen. Physiol.* 140:249-266.

A recent study in *The Journal of General Physiology* characterizes the movement of rhodopsin, a GPCR and member of a large family of transmembrane receptors responsible for many cellular responses and involved in many human diseases.

In living Xenopus rod photoreceptor cells, rhodopsin is found in stacks



of disc membranes that are difficult to study due to their small size. Now, Peter Calvert (SUNY Upstate Medical University) and colleagues have developed an approach with two-<u>photon microscopy</u> that, for the first time, allows direct examination within the living cell of how the geometry of the micro-compartment will affect its signaling.

The team focused on the effect of micro-compartments formed by "incisures", radial membrane boundaries that segment the discs into smaller regions. These micro-compartments, defined by the rim and incisures, have a <u>surface area</u> similar to that of a human or mouse rod photoreceptor disc, so that the total number of components in the GPCR signaling cascade are conserved, and the downstream effects of a captured photon of light are similar. Calvert and colleagues used a novel high-resolution approach to monitor fluorescently labeled rhodopsin as it diffused in the disc membrane.

The team developed a model of <u>molecular diffusion</u> to interpret the results that explicitly took the measured micro-compartments into account, and found that the boundary geometry, not heterogeneity in diffusion or a "bound fraction," was sufficient to explain differences in the extent of diffusion or rhodopsin in different regions of the disc.

More information: Najafi, M., et al. 2012. J. Gen. Physiol. 140:249. Adler, E.M. 2012. *J. Gen. Physio.* 140:341.

Provided by Rockefeller University

Citation: Geometry plays a role in GPCR transmembrane signaling (2012, September 26) retrieved 3 May 2024 from https://medicalxpress.com/news/2012-09-geometry-role-gpcr-transmembrane.html



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