The outer segment serves as a default destination for the trafficking of membrane proteins in photoreceptors

Sheila A. Baker, Mohammad Haeri, Peter Yoo, Sidney M. Gospe III, Nikolai P. Skiba, Barry E. Knox, and Vadim Y. Arshavsky

1 Albert Eye Research Institute, Duke University Medical Center, Durham, NC 27710
2 Department of Biochemistry and Molecular Biology and Ophthalmology, State University of New York, Upstate Medical University, Syracuse, NY 13210

Photoreceptors are compartmentalized neurons in which all proteins responsible for evoking visual signals are confined to the outer segment. Yet, the mechanisms responsible for establishing and maintaining photoreceptor compartmentalization are poorly understood. Here we investigated the targeting of two related membrane proteins, R9AP and syntaxin 3, one residing within and the other excluded from the outer segment. Surprisingly, we have found that only syntaxin 3 has targeting information encoded in its sequence and its removal redirects this protein to the outer segment. Furthermore, proteins residing in the endoplasmic reticulum and mitochondria were similarly redirected to the outer segment after removing their targeting signals. This reveals a pattern where membrane proteins lacking specific targeting information are delivered to the outer segment, which is likely to reflect the enormous appetite of this organelle for new material necessitated by its constant renewal. This also implies that every protein residing outside the outer segment must have a means to avoid this “default” trafficking flow.

Introduction

Normal vision is dependent on the function of rod and cone photoreceptors, which are sensory neurons proven to be among the most productive model systems for studying signal transduction and sensory physiology. The key feature that makes them so amenable is their highly compartmentalized organization. Photoreceptors consist of four major compartments: a photosensitive outer segment containing highly packed disc membranes and the proteins required to initiate a response to light, an inner segment where proteins and lipids are synthesized and energy is produced, a nuclear region, and a synaptic terminal that sends information to the second order neurons in the retina (for reviews see Papermaster, 2002; Williams, 2004).

To establish this polarization, photoreceptors need to tightly control the subcellular targeting of many newly synthesized proteins. As in other cells, protein segregation into distinct membrane compartments requires regulated sorting in the ER and Golgi followed by vesicular trafficking to final destinations. The importance of correct protein targeting in photoreceptors is highlighted by the observations that mutations in targeting signals or trafficking machinery cause degenerative diseases. However, specific targeting information has been revealed for only a handful of photoreceptor-specific proteins, with most of the emphasis on rhodopsin.

One interesting aspect of photoreceptor compartmentalization is that its electrically continuous plasma membrane is separated by a diffusional barrier into two domains with distinct protein compositions, one representing the outer segment and another the rest of the cell. Notably, the densely packed discs comprising the majority of outer segment membranes are formed inside the outer segment by evaginations of the plasma membrane, requiring all disc-specific proteins to be first targeted to the outer segment compartment of the plasma membrane (Steinberg et al., 1980). By analogy with other polarized cells (for reviews see Mostov et al., 2003; Muth and Caplan, 2003; Rodriguez-Boulan and Musch, 2005),
it is likely that membrane proteins residing in each compartment are delivered by separate trafficking pathways and encode distinct targeting sequences that identify which of these pathways will be used. Indeed, targeting signals responsible for directing membrane proteins to the outer segment have been identified. One motif is shared by rhodopsin, cone opsins, and photoreceptor-specific retinol dehydrogenase (Deretic et al., 1998; Tam et al., 2000; Luo et al., 2004) and the second is found in RDS/peripherin 2 (Tam et al., 2004). However, these targeting signals are not present in any other membrane proteins resident in the outer segment and virtually nothing is known about protein targeting to the rest of the plasma membrane.

The goal of this study was to understand the differential targeting of two proteins, which are very similar structurally yet reside in the two separate membrane compartments. One was R9AP, which resides primarily in the outer segment, both in the plasma membrane and the photoreceptor discs. R9AP’s function is to anchor the GTPase-activating complex, which sets the duration of rods’ and cones’ response to light (Hu and Wensel, 2002). The second protein was syntaxin 3, which is a plasma membrane protein excluded from the outer segment and enriched at the synaptic terminal. Its function is to serve as a target membrane component of the SNARE complex mediating vesicular fusion (for review see Hay, 2001). Our approach took advantage of the experimental model of transgenic *Xenopus laevis* (Knox et al., 1998; Amaya and Kroll, 1999; Tam et al., 2000).

We analyzed a series of each protein’s mutants and chimeric constructs expressed as transgenes in *Xenopus* rods to determine the sequence information that governs the differential targeting of each protein.

Surprisingly, we have found that only one of these proteins, syntaxin 3, has targeting information encoded in its sequence. In the case of R9AP, all that is needed to get primarily to the outer segment is its transmembrane domain, which is not a sequence-specific signal. Furthermore, removing the information defining the cellular localization from syntaxin 3 redirected its localization primarily to the outer segment. This was also true for two other proteins we tested, normally residing in mitochondria or the ER; depriving them of specific targeting information redirected them to the outer segment as well. These findings reveal a pattern in which proteins lacking specific targeting sequences are delivered predominantly to the outer segment. This “default” targeting is so efficient that even key signaling proteins, such as R9AP, may reach the photoreceptor outer segment, which is traditionally viewed as a privileged cellular destination, in the absence of a targeting signal.

**Results**

**Differential subcellular localization of R9AP and syntaxin 3 in frog photoreceptors**

R9AP and syntaxin 3 are tail-anchored transmembrane proteins sharing a great degree of structural similarity (Keresztes et al., 2003; Martemyanov et al., 2003; Sharma et al., 2006). Each contains an N-terminal trihelical bundle, called the Habc domain in syntaxins, followed by the SNARE homology domain and the C-terminal transmembrane domain (Fig. 1 A). Their subcellular localization patterns are distinctly different in mammalian rods (Morgans et al., 1996; Hu and Wensel, 2002; Keresztes et al., 2003; Martemyanov et al., 2003; Sherry et al., 2006; Chuang et al., 2007) and we confirmed this in the frog retina.
(Fig. 1, B and C). R9AP was found primarily in rod and cone outer segments with a small fraction in synaptic terminals, as in other species (Fig. 1 B). Syntaxin 3 is present in both photoreceptors and the inner retina (Fig. 1 C). Within photoreceptors, it was found throughout the entire membrane of the inner segment with enrichment in synaptic terminals, but was notably absent in the outer segments. A recent study reported that in mouse photoreceptors syntaxin 3 labeling could be detected within the base of the outer segment (Chuang et al., 2007). We examined this region in the frog photoreceptor, which is best labeled by the antibody in a broken rod preparation containing the outer segment and a fragment of the inner segment (Kaplan et al., 1987), and observed robust syntaxin 3 immunostaining in the remaining portion of the inner segment and in the inner segment calycal processes surrounding the base of the outer segment, but no staining beyond this junction (Fig. 1 D).

Therefore, R9AP and syntaxin 3 could be used as markers for distinct membrane domains in photoreceptors. Our strategy in identifying the sequence information required for each of them to reach their corresponding subcellular destinations was to first perform a deletion mutagenesis on each protein individually and then to analyze the localization of chimeric constructs containing various combinations of R9AP and syntaxin 3 domains. But before presenting the analysis of the targeting information coded in their sequences, we describe how we identified the *Xenopus* homologues of these proteins to use in this study. The National Center for Biotechnology Information database contains a single *Xenopus* homologue for syntaxin 3 (Gene ID 444751) and this sequence was used in our experiments. However, the identification of the photoreceptor-specific R9AP required additional analysis.

**Identification of the photoreceptor-specific R9AP isoform in *Xenopus***

Unlike mammals, frogs have two genes homologous to R9AP (Fig. 2 A). We termed the first gene xR9AP1 (Gene ID 494990) because it is more similar to the mammalian R9AP gene (62% identical to mouse R9AP) and the second gene xR9AP2 (Gene ID 444631; 47% identical to mouse R9AP). To determine which one is expressed in rods, we first used RT-PCR on DNase-treated RNA isolated from *Xenopus* retina and found that both R9AP transcripts are present (Fig. 2 B). We next raised antibodies against each R9AP protein and found that the anti-xR9AP2 antibody recognized a robust doublet at the predicted molecular mass of 27 kD in the membrane fraction of *Xenopus* retina homogenate (Fig. 2 C), whereas the anti-xR9AP1 antibody showed no signal in the predicted molecular mass range (not depicted). Finally, we coimmunoprecipitated R9AP from frog retinas using antibodies against its photoreceptor-specific partner, RGS9-1, and confirmed the identity of the coprecipitating protein as xR9AP2 but not xR9AP1 by both Western blotting (Fig. 2 D) and mass spectrometry sequencing (not depicted). Although we cannot entirely exclude the possibility that low undetectable amounts of xR9AP1 are present in frog photoreceptors, we conclude that xR9AP2 is the major R9AP isoform in frog photoreceptors and we used it for the remainder of the study. For simplicity, we refer to it as R9AP from here on.

**Figure 2. Identification of the photoreceptor-specific R9AP isoform in *Xenopus***

(A) Phylogeny tree of R9AP protein sequences built using the Neighbor-Joining method. Accession nos. are as follows: bovine, AAM67417; chicken, NP_989836; chimpanzee, XP_512566; dog, XP_855518; human, NP_997274; macaque, XP_001087093; mouse, NP_665839; pufferfish, CAF92756; rat, XP_001079782; zebrafish, XP_687644. *Xenopus tropicalis* R9AP1 is deduced from genome v4.1, scaffold 94 (920566/921276), and *X. tropicalis* R9AP2 is deduced from genome v4.1, scaffold 761 (402303-408685). (B) RT-PCR amplification of xR9AP1 and xR9AP2 transcripts from Xenopus retina; omission of reverse transcription (RT) in parallel samples served as a negative control for DNA contamination. (C) Western blot of Xenopus retina extract probed with the antibody specific to xR9AP2. Lane T, total extract; lane S, soluble protein fraction; lane M, membrane fraction. Note that the xR9AP2 band is a doublet, which reflects the previously documented phosphorylation of R9AP in photoreceptors (Martemyanov et al., 2003). (D) xR9AP2 coimmunoprecipitates with RGS9-1 from *Xenopus* retina. Beads coated with sheep IgG were used for the negative control.

The transmembrane domain of R9AP is sufficient for outer segment targeting

We first identified the regions of R9AP required for its targeting to rod outer segments. A series of R9AP fragments were fused to GFP and expressed under the rod-specific rhodopsin promoter (Knox et al., 1998); a summary of all transgenically expressed proteins used in this study is shown in Table I. To establish the baseline, we studied the distribution of full-length R9AP, which was found to be similar to that of the endogenous protein (Fig. 3 A). The majority of GFP signal originated from the outer segment, with some also detected in the synaptic terminal and plasma membrane of the rod. Note that the GFP signal in the outer segments often appears as distinct bands of various thicknesses and that the transgene expression is mosaic from cell to cell. This phenomenon has
Table I. Summary of all transgenic constructs presented in this study

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<th>Construct</th>
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ER, endoplasmic reticulum; gg, geranylgeranylated; IS, inner segment; ISPM, inner segment plasma membrane; MT, mitochondria; OS, outer segment.
tested two Myc-tagged R9AP deletion mutants and found no difference between the targeting of these molecules and the GFP-tagged variants (Fig. S1, B and C). Finally, removal of the transmembrane domain from GFP-R9AP resulted in a soluble protein localized primarily to the inner segment (Fig. 3 E), including the nucleoplasmic space, in a pattern indistinguishable from that of GFP alone (see Fig. 5 A; Knox et al., 1998; Peet et al., 2004). Overall, these data indicate that the transmembrane domain of R9AP is both necessary and sufficient to achieve outer segment localization.

The subcellular localization of syntaxin 3 requires the SNARE domain

We next conducted a similar deletion mutagenesis with syntaxin 3 constructs. First, we looked at the full-length GFP-tagged Xenopus syntaxin 3 (Fig. 4 A) and found its subcellular localization pattern to be the same as that of the endogenous protein (Fig. 1, C and D). Indeed, this resulted in far fewer aggresomes with the majority of Myc-R9AP detected in the outer segment and a low level of signal found in the synaptic terminal as is the case with endogenous R9AP. Note that the overexpression of R9AP or any of the other proteins we expressed in transgenic rods in the course of this study did not result in any signs of retinal cell death.

The data shown in Fig. 3 (B–D) demonstrate that the pattern of R9AP distribution remains essentially unchanged in a series of deletion mutants lacking either some or all of its cytoplasmic domains. As in the case of full-length R9AP, we also tested two Myc-tagged R9AP deletion mutants and found no difference between the targeting of these molecules and the GFP-tagged variants (Fig. S1, B and C). Finally, removal of the transmembrane domain from GFP-R9AP resulted in a soluble protein localized primarily to the inner segment (Fig. 3 E), including the nucleoplasmic space, in a pattern indistinguishable from that of GFP alone (see Fig. 5 A; Knox et al., 1998; Peet et al., 2004). Overall, these data indicate that the transmembrane domain of R9AP is both necessary and sufficient to achieve outer segment localization.

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was found in internal membranes of the inner segment (Fig. 4 B). However, the deletion of the SNARE domain redirected a large portion of the expressed protein to the outer segment (Fig. 4 C), with an overall pattern mirroring that of R9AP. The deletion of the entire cytoplasmic domain of syntaxin 3, leaving just the transmembrane domain, also resulted in robust outer segment localization (Fig. 4 D). This analysis suggests that the intracellular targeting information for syntaxin 3 is coded within the SNARE domain.

We next tested whether the targeting role of the SNARE domain could be fully preserved when the adjacent transmembrane domain is removed. However, just deleting the transmembrane domain (or both SNARE and transmembrane domains) from GFP-tagged syntaxin 3 resulted in protein distributions similar to that of soluble GFP, which reflects the availability of the rod cytoplasm (Fig. 5, compare B and C to A; Knox et al., 1998; Peet et al., 2004). To make these constructs membrane associated we placed a CaaX box (CIIL) on the C termini of these proteins to enable their geranylgeranylation. This modification dramatically changed the intracellular localization of both constructs lacking the SNARE domain to the outer segment (Fig. 5, D and F), whereas the construct containing the SNARE domain was localized to the synaptic terminus and the plasma membrane of the inner segment (Fig. 5 E), just as the full-length syntaxin 3 in Fig. 4 A.

Together, the data from Figs. 4 and 5 indicate that the SNARE domain plays a decisive role in setting the subcellular localization of syntaxin 3. They further demonstrate that the removal of targeting information releases this protein primarily to the outer segment (as far as the resulting construct remains membrane associated). This suggests that the outer segment may serve as a default destination for membrane protein delivery in this cell type, a suggestion tested in the subsequent experiments.

It should be also noted that a study of syntaxin 3 in epithelial cells did not reveal that the SNARE domain was required for its normal subcellular localization to the apical rather than the basolateral plasma membrane and instead demonstrated that a four–amino acid FMDE motif within the Habc domain was required (Sharma et al., 2006). To determine if this motif plays any role in the localization of syntaxin 3 in photoreceptors, we replaced the FMDE sequence with four alanines and found that this mutation had no effect on syntaxin 3 targeting in rods (Fig. 4 E), illustrating that different cell types can use different information for targeting the same protein.

Subcellular targeting of R9AP/syntaxin 3 chimeras

An alternative approach we used to identify regions within the R9AP and syntaxin 3 molecules involved in their targeting was to analyze subcellular distributions of a series of chimeric constructs containing various combinations of domains from each of these two proteins (Fig. 6). Because of the propensity for several of the GFP-tagged chimeric molecules to form aggresomes (not depicted), we instead used the Myc-tagged versions of these constructs. Replacement of R9AP’s transmembrane domain (sufficient for its primarily outer segment localization) with the one from syntaxin 3 had no effect on the chimera’s

Figure 5. Localization of soluble and geranylgeranylated GFP-tagged syntaxin 3 mutants in transgenic Xenopus rods. (A) Soluble GFP. (B) GFP-tagged syntaxin 3 lacking both the SNARE and transmembrane domains. (C) GFP-tagged syntaxin 3 lacking the transmembrane domain. (D) Geranylgeranylated GFP. (E) Geranylgeranylated GFP-tagged syntaxin 3 lacking the SNARE homology domain and transmembrane domain. (F) Geranylgeranylated GFP-tagged syntaxin 3 lacking the transmembrane domain. OS, outer segment; IS, inner segment; N, outer nuclear layer; ST, synaptic terminal. Bars, 5 μm.
localization (Fig. 6, compare C to A), once again confirming that the transmembrane domain is not involved in syntaxin 3 targeting. However, the replacement of the SNARE homology domain of syntaxin 3 (sufficient for its exclusion from the outer segment) with the one from R9AP completely shifted the chimera’s localization to the R9AP-like pattern (Fig. 6 D). The replacement of the transmembrane domain from syntaxin 3 with that of R9AP resulted in a molecule maintaining the syntaxin 3 localization pattern (Fig. 6, compare E to B). However, replacement of the SNARE homology domain from R9AP with that of syntaxin 3 prevented the molecule from targeting to the outer segments (Fig. 6 F). A detailed description of each chimeric molecule is provided in Table S1 (available at http://www.jcb.org/cgi/content/full/jcb.200806009/DC1).

Consistent with the results of the deletion mutagenesis, these data provide complementary evidence that the SNARE homology domain in syntaxin 3 plays an active role in excluding this protein from the outer segment. To the contrary, the transmembrane domain of R9AP does not appear to be unique in directing outer segment targeting. But are the transmembrane domains from R9AP and syntaxin 3 interchangeable because they belong to structurally related proteins or would any transmembrane domain lacking specific targeting information be targeted to the outer segment?

The length of a transmembrane domain is critical for outer segment targeting

To address this question, we analyzed the targeting of other GFP-tagged transmembrane domains from proteins either not expressed in photoreceptors or not present in the outer segments. For the former, we chose the first transmembrane segment of the metabotropic glutamate receptor 1 (mGluR1), which is not expressed in rods (Koulen et al., 1997; Sen and Gleason, 2006). As shown in Fig. 7, this “random” transmembrane construct accumulated in the outer segments, similarly to R9AP.

For the non-outter segment proteins, we choose microsomal cytochrome b5 and BclXL, an antiapoptotic protein that resides in the mitochondrial outer membrane. Like R9AP and syntaxin 3, these proteins contain a single pass transmembrane domain at their C termini. In the case of cytochrome b5 (Pedrazzini et al., 2000), the transmembrane domain is sufficient to ensure the localization of reporter molecules to the ER. This property is dictated by the relatively short length of the cytochrome b5 transmembrane domain, predicted to contain 14 amino acids (Table S2, available at http://www.jcb.org/cgi/content/full/jcb.200806009/DC1). Increasing this length by several hydrophobic amino acids shifts the intracellular localization from the ER to the plasma membrane in the fibroblast cell line CV-1 (Pedrazzini et al., 1996). The data from Fig. 7 A demonstrate that in Xenopus rods the construct containing a GFP-tagged cytochrome b5 transmembrane domain is targeted to internal membranes of the inner segment where it overlaps with the ER marker protein, calnexin (Fig. S2). However, increasing the length of this transmembrane domain by four amino acids redirected it almost exclusively to the outer segment (Fig. 8 B).

In the case of BclXL, its mitochondrial localization is determined by the positively charged residues in the sequence immediately flanking the transmembrane domain (Kaufmann et al., 2003). We confirmed that the corresponding GFP-tagged...
length are predisposed to traffic to the outer segment, regardless of their specific amino acid composition.

Do proteins targeted by default to photoreceptor outer segments target to cilia in other cell types?

The rod outer segment is a modified primary cilium (for review see Besharse et al., 2003). Therefore, in the last set of control experiments, we tested whether the pattern of preferential membrane protein localization to the outer segment is a property of the photoreceptor cell or ciliary structures in general. We generated transgenic tadpoles expressing either GFP-tagged R9AP or the GFP-tagged transmembrane domain from syntaxin 3 (both targeted to rod outer segments by default) in a nontissue-specific manner under the control of a ubiquitous CMV promoter. In general, these reporters displayed essentially identical localization patterns throughout the tadpole body, although the latter construct had a much lower propensity to form aggresomes. Three examples are shown in Fig. 10. In photoreceptors, the CMV promoter drove expression in cones much more abundantly than in rods, providing us with a chance to show that both types of photoreceptors handle these proteins similarly by targeting them to the outer segments (Fig. 10 A). In the cells of the nasal pit, which at this developmental stage consists of both ciliated and nonciliated olfactory neurons and support cells (Hansen et al., 1998), the reporter proteins were found throughout the cell with the notable exception that they were absent from the cilia, visualized by labeling with acetylated α-tubulin (Fig. 10 B). The epidermis of the tadpole consists of mucous-secreting cells regularly interspaced with epidermal cells containing

Collectively, our data from five different examples suggest that proteins containing transmembrane domains of sufficient length are predisposed to traffic to the outer segment, regardless of their specific amino acid composition.
multiple motile cilia (Steinman, 1968). In these cells, the reporter proteins were localized nonspecifically to the plasma membrane but were absent from the cilia (Fig. 10 C). Together, these data argue that the membrane protein trafficking pattern in rods and cones is not determined simply by the presence of a ciliary organelle.

Discussion

The compartmentalized organization of vertebrate photoreceptors is elegantly suited to serve their function as highly sensitive light detectors. Yet, little is known about the mechanisms targeting individual proteins to their respective intracellular compartments in these cells. Most recently, substantial effort has been directed toward studying proteins targeting to the outer segment, with a strong emphasis on the trafficking of rhodopsin. In this study, we took a more general look at this problem by comparing the targeting of two related proteins, syntaxin 3 and R9AP, into the two distinctly separate plasma membrane domains, the outer segment and the remainder of the cell. Our major finding is that, whereas specific targeting information was required for achieving protein localization outside the outer segment, no sequence-specific signals were required for the outer segment targeting as far as a protein contained a transmembrane domain of sufficient length and lacked overriding sequence signals targeting it elsewhere. This suggests that the outer segment protein composition could be achieved by having only one or a few proteins that actively direct vesicular trafficking to this compartment. Other membrane proteins, such as R9AP, may potentially be cotransported in the same vesicles. Here we will discuss how this idea fits the current understanding of photoreceptor outer segment trafficking.

Outer segment protein targeting in photoreceptors

Membrane protein trafficking to the outer segment has been studied most extensively for rhodopsin and is thought to consist of several discrete steps (for reviews see Papamaster, 2002; Williams, 2004; Deretic, 2006). After synthesis at the ER and passing through the Golgi, rhodopsin is sorted into newly budding vesicular transport carriers. This is mediated by the interaction between the VXPX targeting sequence in rhodopsin’s C terminus (Deretic et al., 1998; Tam et al., 2000) and the small GTP binding protein ARF4 (Deretic et al., 2005). Other small GTP binding proteins, rab6 and rab11, also play a role in formation of the rhodopsin carriers in the trans-Golgi (for review see Deretic, 2006). The carriers are then delivered to the base of the cilium connecting the inner and outer segments with the aid of the motor cytoplasmic dynein (Tai et al., 1999). There the carriers dock and fuse with a specialized region of the plasma membrane separated from the plasma membrane of the inner segment via a diffusional barrier. This docking is assisted by another set of small GTP binding proteins, rac1 and rab8 (Moritz et al., 2001a; Deretic et al., 2004). Rhodopsin then moves laterally through the membrane of the connecting cilium into the outer segment, possibly assisted by cytoskeletal motors (Liu et al., 1999; Marszalek et al., 2000). The V(I)XPX signal is also present in cone opsins and in photoreceptor-specific retinol dehydrogenase (Deretic et al., 1998; Tam et al., 2000; Luo et al., 2004) but not in any other proteins resident in the outer segment.

This major transport pathway is necessary for the outer segment to form and for the cell to survive. For example, the lack of outer segment formation and severe degeneration observed in rhodopsin knockout mice (Humphries et al., 1997; Lem et al., 1999) can be rescued by the expression of normal rod or cone opsin (Concepcion et al., 2002; Guang Shi, 2004), but not by rhodopsin in which the VXPX sequence has been deleted (Concepcion et al., 2002; Lee and Flannery, 2007). However, when rhodopsin lacking the VXPX sequence is expressed in rods in addition to normal rhodopsin, outer segments are formed with most mutant molecules observed inside the outer segment and a smaller fraction in other cellular compartments,
resembling the normal R9AP distribution (Sung et al., 1994; Li et al., 1996; Green et al., 2000; Moritz et al., 2001b; Concepcion et al., 2002; Kraft et al., 2005; Tam et al., 2006; Lee and Flannery, 2007). The V(1)XPX-dependent pathway may not be the only one targeting proteins to the outer segment because peripheralin 2 has an alternative targeting sequence, and it has been suggested that delivery of this protein relies on a distinct set of transport vesicles (Fariss et al., 1997; Tam et al., 2004; Lee et al., 2006). Most recently, it has been speculated that guanylate cyclase is another protein that marks independent carriers to the outer segment, yet no details of how these putative vesicles are assembled and targeted are known (Karan et al., 2007).

**Outer segment protein localization can be achieved without specific targeting signals**

Here we have shown that a key signaling protein, R9AP, can be delivered to the outer segment without containing any specific targeting signals coded in its sequence. Furthermore, every other transmembrane protein construct that we tested was delivered to the rod outer segment as far as it obeyed two simple rules. First, the transmembrane domain had to be of a sufficient length. This most likely reflects the phenomenon of hydrophobic matching whereby short transmembrane domains partition into the thin bilayer of the ER and the longer transmembrane domains preferentially partition into the thicker bilayers of the Golgi and post-Golgi components of the secretory pathway, which contain more sterols and complex sphingolipids (for review see Killian, 1998; Andersen and Koepe, 2007; van Meer et al., 2008). Second, the protein had to be stripped of any targeting information directing it to other subcellular compartments.

The first rule was established by showing that increasing the length of cytochrome b5’s transmembrane domain redirected it from the ER to the outer segment and, conversely, the reduction in the length of R9AP’s transmembrane domain had the opposite effect of shifting its localization from primarily outer segments to ER. The second rule was demonstrated with syntaxin 3, which was redirected from the plasma membrane of the inner segment and synaptic terminal to the outer segment upon the loss of its SNARE homology domain. BclXl bridged both rules: its redirection from mitochondria to outer segments required both an increase in the length of the transmembrane domain and a reduction in the positive charge in the flanking region. Finally, a randomly selected transmembrane segment from metabotropic glutamate receptor 1, which obeyed both rules, localized predominantly to the outer segment.

Interestingly, the ability of the SNARE domain of syntaxin 3 to mediate exclusion from the outer segment was retained when the transmembrane domain was replaced by geranylgeranylation. Intracellular localization of a protein containing the SNARE domain of syntaxin 3 was indistinguishable from normal syntaxin 3, whereas protein constructs lacking this targeting domain were localized predominantly to the outer segment, similar to transmembrane domain containing proteins carried by the default pathway. This is consistent with previous results that GFP-tagged constructs that were myristoylated or palmitoylated predominantly localized to the outer segment as well (Tam et al., 2000). It is intriguing to speculate that peripheral membrane proteins may use the same default targeting pathway as the proteins we analyzed in this study. However, in the absence of a systematic analysis this idea should be treated cautiously particularly because all cells, including photoreceptors, express an array of proteins specifically binding to lipidated proteins and regulating their intracellular localization. Examples of such proteins actively studied in photoreceptors include the prenyl binding protein PrBP/δ, which regulates the outer segment localization of rhodopsin kinase and the cyclic guanosine monophosphate phosphodiesterase (Norton et al., 2005; Zhang et al., 2007), and phosphducin, which regulates the inner–outer segment translocation of the transducin βγ subunit (Sokolov et al., 2004).

Notably, we have also found that the targeting information of syntaxin 3 in photoreceptors is coded differently than in epithelial cells, where its normal localization in the apical plasma membrane is determined by the FMDE motif (Sharma et al., 2006). Substitutions of these amino acids in epithelial cells caused a randomized syntaxin 3 distribution throughout both apical and basolateral membranes. However, the substitution of the entire motif with alanines in transgenic frog rods did not affect its normal inner segment plasma membrane distribution. This example illustrates that different cell types may use different mechanisms for achieving specific intracellular targeting.
The actual mechanism by which the SNARE domain directs syntaxin 3 targeting in rods remains beyond the scope of this study. Because the SNARE domain is the region involved in almost all known protein–protein interactions, there are several possibilities including that this domain encodes a specific signal (interacting with a component of vesicular trafficking such as a Rab GTPase or coat protein), which could be conformation-dependent (Mancias and Goldberg, 2007), or that syntaxin 3 must assemble into a SNARE protein complex, which again could either reveal a conformation-dependent targeting signal or provide an interaction partner that binds directly to a component of the vesicular trafficking machinery (Mossessova et al., 2003).

In summary, our findings indicate that the default route of membrane protein trafficking in photoreceptors is directed to the outer segment. This mode of delivery is so reliable that it is used for trafficking R9AP, a protein whose expression level is tightly regulated in photoreceptors and sets the dominant rate constant at which the cell recovers from the photoresponse (Krispel et al., 2006). This raises important questions concerning the interplay between targeted and default protein flow in these cells.

The relation between default and targeted protein trafficking in photoreceptors

Essential to understanding default trafficking patterns mechanistically is the recognition that intracellular vesicular transport always requires at least one cargo protein to contain specific signals that regulate their delivery to individual subcellular locations (for reviews see Muth and Caplan, 2003; Rodriguez-Boulan and Musch, 2005). This suggests that proteins lacking such signals and carried by default pathways have to be copackaged into the same vesicles as proteins that do carry specific targeting information. Therefore a straightforward hypothesis to explain the outer segment delivery of R9AP is that it is transported in the same post-Golgi vesicles as rhodopsin, the most abundant transport vesicles in this cell. In this case, even a random incorporation of proteins lacking a targeting signal into all types of transport carrier vesicles would result in the majority reaching the outer segment and only a small fraction delivered elsewhere. Consistently, a small fraction of R9AP is always found outside the outer segment (Hu and Wensel, 2002; Martemyanov et al., 2003; this study). Similarly, when rhodopsin mutants lacking the VXPX sequence are expressed in rods containing normal rhodopsin, they display a localization pattern very similar to that of R9AP, with the majority found in the outer segment and a minor fraction in the inner segment membranes (compare Fig. 6 from Moritz et al. [2001b] with Fig. 2). This was interpreted as the mutant rhodopsin cotransporting with the wild-type rhodopsin (Concepcion et al., 2002). The new development in this study is that such cotransport may serve as the normal route of outer segment delivery for a functional protein, R9AP.

The next interesting consideration is whether the incorporation of proteins lacking a targeting signal into all types of transport carrier vesicles is random or if it is possible that vesicles carrying proteins to destinations other than the outer segment are less permissive for random protein incorporation than those responsible for the default trafficking flow. Additionally, the degree of this permissiveness may be affected by the specific properties of individual proteins. Indeed, for the various protein constructs using the default pathway, we observed a significant variation in the amounts present outside the outer segment (including an apparent lack for mutated BclXI or cytochrome b5). Although some of this variation may relate to the differences in the constructs’ expression levels, it could also reflect the relative permissiveness of different types of transport vesicles to incorporate individual proteins. Addressing these questions requires identification of all the major classes of transport carriers in photoreceptors, which is an exciting direction of future studies.

Importantly, the default mode of protein delivery can be used only on the condition that minor mislocalization is not harmful to the cell. Accordingly, the overexpression of the proteins using the default trafficking flow that we examined here did not cause any overt signs of photoreceptor stress or lead to death in the time frame of this study. Furthermore, the normal “leak” of R9AP appears nonconsequential. This is in contrast to rhodopsin, where even minor mislocalization leads to photoreceptor degeneration, although not nearly as severe as in the absence of normal rhodopsin (Lee and Flamery, 2007).

Another logical question is whether other integral or even peripheral membrane proteins use the default trafficking pathway in photoreceptors. This is an exciting topic to explore, particularly given that nearly nothing is known about the delivery of many proteins responsible for signaling and maintenance in the outer segment (such as guanylate cyclase, ABCR transporter, or cyclic guanosine monophosphate–gated channel, just to mention a few). But perhaps an even more significant consequence of our finding that the default protein flow in photoreceptors is directed to outer segments is the implication that every membrane protein residing in other cellular compartments has to use specific mechanisms to avoid this destination. Disruption of the targeting information in each non-out segment protein did not simply yield a randomized distribution among all cellular membranes, but resulted in substantial redirection to the outer segment. This calls for expanding the questions in addressing protein targeting in photoreceptors from what allows a restricted subset of cellular proteins to reach the outer segment to what allows the greater number of membrane proteins residing in other cellular compartments to avoid the outer segment.

Lastly, it is interesting to consider the possibility that other differentiated cells may use an interplay between specific targeting mechanisms and default trafficking patterns to meet their compartmentalization needs. Traditionally, a constitutive (or default) trafficking pathway was considered to account for the bulk of protein trafficking to the plasma membrane, at least in nonpolarized cells (for review see Ponnambalam and Baldwin, 2003). Polarized cells are more complex and many proteins that were initially thought to be delivered by default have now been shown to be delivered in a signal-dependent manner (for review see Rodriguez-Boulan and Musch, 2005). Our findings in photoreceptors indicate that default trafficking may be reliable enough for the delivery of a key signaling protein, which may call for reconsidering the relative contributions of default and targeted trafficking in various cells. However, the example of photoreceptors also illustrates that this should be explored in the context of each specific cell type.
Materials and methods

Generation of transgenic tadpoles

Transgenic Xenopus tadpoles were produced using restriction enzyme–mediated integration development previously (Kroll and Amaya, 1996; Amaya and Kroll, 1999) with modifications (described in Batni et al., 2000; Whitaker and Knox, 2004). A minimum of four individual transgenic animals were evaluated for every DNA construct.

The constructs were generated using standard PCR-based subcloning methods. Fusion constructs and large deletions were generated using splicing by overhang extension PCR primers; point mutations and small deletions or insertions were generated using the QuikChange II XL kit (Stratagene). PCR products were subcloned into the XOP5.5 vector (Knox et al., 1998) and confirmed by direct sequencing. DNA templates were obtained from a wild-type tadpole by gentle agitation of the retina in frog Ringer’s solution [115 mM NaCl, 2 mM KCl, 1 mM CaCl2, 1 mM MgCl2, and 5 mM Hepes, pH 7.4]. The fragments were allowed to settle onto a concavalin A [Sigma-Aldrich]-coated glass slide and immunostained as described in the Histology section.

Online supplemental material

Fig. S1 shows the localization of Myc-tagged R9AP and two Myc-tagged R9AP deletion constructs to the outer segments of transgenic tadpole photoreceptors. Fig. S2 shows colocalization of the expressed GFP-tagged cytochrome b5 transmembrane domain with the ER marker calnexin. Fig. S3 shows the localization of a Myc-tagged R9AP variant with a shortened transmembrane domain in transgenic tadpole photoreceptors. Table S1 describes the exact breakpoints of the R9AP/syntaxin 3 chimeric molecules presented in Fig. 6. Table S2 describes the sequence and length of the GFP-tagged transmembrane domains used in this study. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.200806009/DC1.

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