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**Identification of additional outer segment targeting signals
in zebrafish rod opsin**

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29 **Summary statement**

30 We identified a complex set of molecular features that function together as address labels to
31 target opsin to the outer segment.

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57 **ABSTRACT**

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59 **In vertebrate photoreceptors, opsins are highly concentrated in a morphologically distinct**
60 **ciliary compartment known as the outer segment (OS). Opsin is synthesized in the cell**
61 **body and transported to the OS at a remarkable rate of 100-1000 molecules per second.**
62 **Opsin transport defects contribute to photoreceptor loss and blindness in human**
63 **ciliopathies. Previous studies revealed that the opsin C-terminal tail, of 44 amino acids,**
64 **is sufficient to mediate OS targeting in Xenopus photoreceptors. Here we show that**
65 **although the Xenopus C-terminus retains this function in zebrafish, the homologous**
66 **zebrafish sequence is not sufficient to target opsin to the OS. This functional difference**
67 **is largely caused by a change of a single amino acid present in Xenopus, but not in other**
68 **vertebrates examined. Furthermore, we find that sequences in the 3rd intracellular**
69 **cytoplasmic loop (IC3) and adjacent regions of transmembrane helices 6 and 7 are also**
70 **necessary for opsin transport in zebrafish. Combined with the cytoplasmic tail, these**
71 **sequences are sufficient to target opsin to the ciliary compartment.**

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81 Vertebrate photoreceptors detect light via a specialized cilium that contains hundreds of parallel
82 membrane folds harboring the photosensitive pigment, opsin, and other components of the
83 phototransduction cascade. It is estimated that 10^8 - 10^9 opsin molecules are tightly packed in
84 the ciliary membranes of a single photoreceptor cell (Pugh and Lamb, 2000). Opsin is the
85 main protein component of photoreceptor cilia and in addition to detecting light its presence is
86 also essential for cilia morphogenesis (Lem et al., 1999, Pugh and Lamb, 2000).
87 Photosensitive membranes of photoreceptors are continuously turned over: they are shed from
88 the distal end of the cilium while new membrane folds are added to the cilium base (Young,
89 1967, Young and Droz, 1968). In the mouse retina, this process replaces approximately 10%
90 of the photoreceptor ciliary membranes every day (LaVail, 1973). Consequently, 10% of all
91 protein content of the cilium, including opsins and other phototransduction cascade components,
92 are transported into the cilium every day.

93

94 The complex architecture of photosensitive membranes results in an unusually large volume of
95 photoreceptor cilia (reviewed in ref. Kennedy and Malicki, 2009). The bulk of the cilium,
96 referred to as the outer segment (OS), connects to the cell body via a narrow stalk, termed the
97 connecting cilium, characterized by the typical architecture and dimensions of the ciliary
98 transition zone (Tokuyasu and Yamada, 1959, Nilsson, 1964, Besharse et al., 1985, Röhlich,
99 1975). All proteins destined for the OS, including opsins, are transported through the
100 connecting cilium (reviewed in Kennedy and Malicki, 2009, Malicki and Besharse, 2012).
101 Given the unusually large size of the OS and its high protein content, transport into the
102 photoreceptor cilium requires a particularly efficient mechanism (Malicki and Besharse, 2012).

103

104 Many attempts have been made to gain insight into the mechanism of [rod opsin transport](#). A
105 key finding in this area is that the cytoplasmic tail of 44 amino acids of [rod](#) opsin (hereafter
106 CT44) is sufficient to mediate transport into the *Xenopus* OS (Tam et al., 2000). Human
107 mutations in the homologous sequence lead to rapid vision loss (Sung et al., 1991, Bessant et
108 al., 1999, Berson et al., 2002, reviewed in Athanasiou et al., 2018, Sung et al., 1994), which

109 further speaks to its importance. To direct transport, **rod** opsin C-terminal sequences must
110 interact with protein components of cytoplasmic and/or ciliary transport machinery. Although
111 some proteins that bind CT44 have been identified using biochemical approaches (Deretic et
112 al., 2005, Mazelova et al., 2009, Wang et al., 2012, Tai et al., 1999, Chuang et al., 2007, Keady
113 et al., 2011), the understanding of this process is far from complete. How such a functionally
114 diverse group of proteins contributes to opsin transport requires further investigation. It is
115 also noteworthy that based on previous analyses, opsin appeared to differ from other ciliary
116 GPCRs, which contain a cilia targeting sequence (CTS) in the 3rd intracellular loop and do not
117 display obvious homologies to opsin C-terminal targeting motifs (Berbari et al., 2008).

118
119 Here we further characterize **rod** opsin OS targeting. Surprisingly, we find that while *Xenopus*
120 CT44 is sufficient to target an exogenous protein, such as GFP, to the ciliary compartment of
121 zebrafish photoreceptors, the homologous zebrafish sequence is unable to do so. A single
122 amino acid in the opsin helix 8 is predominantly responsible for this difference. This suggests
123 that additional sequences are necessary to target **rod opsins** to the OS. Indeed, the analysis of
124 Opsin/Sstr5 hybrid constructs reveals that a targeting sequence exists in the IC3 of rod opsin.
125 Deletion of this motif causes opsin transport deficiency. **Rod** opsin ciliary targeting also
126 requires parts of two transmembrane helices that flank the 3rd cytoplasmic loop. These
127 studies have uncovered novel motifs necessary to target **rod** opsin to the ciliary compartment
128 of the photoreceptor in zebrafish and may allow for the identification of machinery involved in
129 its transport in the future.

130

131 **Results**

132

133 **Construction of transgenic tools to monitor opsin transport.**

134

135 To monitor the dynamics of opsin transport in vivo, we generated stable zebrafish transgenic
136 lines that express an EGFP-opsin fusion specifically in rod photoreceptors from an inducible

137 promoter. This system provides both spatial and temporal control of opsin expression. It
138 consists of two transgenic lines: (1) A line that specifically expresses Cre recombinase in
139 photoreceptors from the rhodopsin promoter, and (2) a temperature-inducible line that
140 conditionally expresses EGFP-S-opsinCT44 (EGFP fused with S-peptide and Xenopus CT44)
141 from a heat-shock promoter (Fig. 1A). A lox-mCherry-STOP-lox cassette is inserted
142 upstream of EGFP-S-opsinCT44 in this line. When these two lines are crossed, Cre induces
143 recombination specifically in rod photoreceptors, placing EGFP-S-opsinCT44 under the control
144 of heat-shock promoter (Fig. 1A). When heat-shock is applied, embryos express EGFP-S-
145 opsinCT44 specifically in rods and mCherry in all other cells (Fig. 1E). The EGFP-S-
146 opsinCT44 fusion protein is efficiently transported to photoreceptor outer segments in these
147 animals (Fig. 1E, I). In a control line, which contains 44 random amino acids in place of the
148 44 amino acids from rhodopsin C-terminus, the EGFP signal is no longer enriched in the OS
149 and instead is present throughout the entire photoreceptor cytoplasm (Fig. 1F).

150

151 **Zebrafish and Xenopus CT44 differ in their ability to opsin transport.**

152

153 In Xenopus, CT44 is sufficient to mediate opsin OS targeting (Tam et al., 2000). To evaluate
154 the function of the zebrafish opsin C-terminal region in OS-directed transport, we fused the 44
155 C-terminal amino acids of zebrafish rod opsin to EGFP and expressed it from a heat-shock
156 promoter using stable transgenic lines, as previously described for Xenopus CT44 (Fig. 1).
157 We compared EGFP transport efficiency in these lines to that in animals containing Xenopus
158 CT44 transgene by assessing EGFP intensity in the cell body. We measured EGFP
159 fluorescence in the cell body rather than the OS, because the signal in the OS becomes saturated
160 quickly, decreasing sensitivity of our measurements. Surprisingly, we found that in contrast
161 to Xenopus C-terminus (XCT44), the equivalent region of zebrafish opsin (ZCT44) is very
162 inefficient at targeting opsin to the zebrafish photoreceptor outer segment (Fig. 1G-I). Similar
163 results were also obtained using a transient expression assay, in which we also used a heat-
164 shock promoter to generate a pulse of GFP-CT44 expression and monitor the rate at which

165 opsin-GFP is cleared from the photoreceptor cell body (Zhao and Malicki, 2011) (Fig. 2B-D).
166 These results indicate that, unexpectedly, opsin targeting mechanisms differ significantly
167 among vertebrate species.

168

169 The 44 C-terminal amino acids of opsin sequence contain an α -helical region, referred to as the
170 helix 8, flanked on the C-terminal side by a pair of cysteines (Fig. 2A) (Palczewski et al., 2000).
171 Several studies suggested that helix 8 mediates opsin dimerization, which, in turn, may play a
172 role in opsin transport (Knepp et al., 2012, Zhang et al., 2016). We therefore hypothesized
173 that helix 8 is responsible for differences in opsin transport efficiency between zebrafish
174 (ZCT44) and Xenopus (XCT44) C-terminal regions. To test this, we replaced helix 8 in the
175 zebrafish C-terminal cytoplasmic tail with the Xenopus sequence (Fig. 2E) and vice versa, we
176 substituted helix 8 of the Xenopus C-terminal tail with the zebrafish sequence (Fig. 2F). The
177 Xenopus helix 8 very significantly improved transport efficiency of ZCT44 (Fig. 2E, I, $p <$
178 0.001). Consistent with that, replacing Xenopus CT44 helix 8 with the zebrafish sequence
179 strongly decreased targeting efficiency (Fig. 2F, I, $p < 0.001$). Xenopus sequences outside
180 helix 8 affect opsin targeting to a lesser degree: replacing the C-terminal sequence of zebrafish
181 ZCT44 with the Xenopus sequence does not result in a significant difference (compare D to F,
182 and I). Replacing the same fragment of Xenopus CT44 with the zebrafish sequence does,
183 however, somewhat decrease targeting efficiency (Fig. 2E, I, $p < 0.05$). These results
184 demonstrate that helix 8 is of paramount importance for the OS targeting of opsin.

185

186 Rod opsin helix 8 is very well conserved across vertebrate phyla (Fig. S1). Zebrafish and
187 Xenopus helix 8 sequences differ only at two positions: 315 and 317 (Fig. 2A). To narrow
188 down which amino acid contributes to opsin targeting efficiency the most, we substituted single
189 amino acids in zebrafish helix 8 with Xenopus ones. We found that while H315N substitution
190 alters transport relatively weakly ($p < 0.05$) (Fig. 2G, I), the substitution of methionine 317 with
191 leucine (M317L) significantly improves transport efficiency (Fig. 2H, I, $p < 0.01$). Sequence
192 comparison of rod opsins from over 20 vertebrate species reveals that position 317 is almost

193 invariably occupied by methionine (Fig S1). The sole exception among these species is
194 *Xenopus*. To test whether the exceptional targeting ability of XCT44 is due to the presence
195 of the leucine, we introduced the sequence of human rod opsin C-terminus (HCT38) to the
196 transient assay. In line with the zebrafish CT44, the targeting deficiency of human CT38,
197 compared to *Xenopus* CT44, was significantly improved by its M317L substitution (Fig. S2B-
198 D, G). We were curious how Leu317 would behave in non-opsin GPCRs. To address this
199 question, we replaced the entire helix 8 in XCT44 with the C-terminal sequence of porcine α
200 adrenergic receptor (Standfuss et al., 2011), which is a GPCR that shares structural similarities
201 with rod opsin. This chimera, which is supposed to localize to the *Xenopus* outer segment
202 (Tam et al., 2000), was found mainly in the cell body and the synapse of zebrafish
203 photoreceptors. Strikingly, the substitution of phenylalanine with leucine also enhanced the
204 OS targeting to a significant level (Fig. S2E-G). Given the importance of this amino acid in
205 opsin targeting (Fig. 2I), our findings show that *Xenopus* CT44 transport into the outer segment
206 is mediated by different molecular interactions, compared to that of zebrafish CT44, and highly
207 likely compared to most other vertebrate rod opsins. Importantly, these findings reveal that
208 in addition to the opsin C-terminus, other opsin sequences are necessary to target opsin to the
209 outer segment.

210

211 **Additional opsin ciliary targeting motifs**

212

213 To search for additional OS-targeting signals in the opsin sequence, we constructed a series of
214 hybrid GPCRs combining sequences from zebrafish rod opsin and another GPCR, Sstr5. We
215 chose Sstr5 because unlike its close relative, SSTR3, it is not targeted to cilia (Berbari et al.,
216 2008). All constructs were transiently expressed from a heat-shock promoter as above.
217 Compared to opsin, Sstr5 is transported to the photoreceptor ciliary outer segment very
218 inefficiently (Fig. 3A, B; compare constructs no. 1 and 2). Substitution of Sstr5 C-terminal
219 cytoplasmic tail with zebrafish CT44 did not result in a significant improvement of transport
220 efficiency (construct no. 3, Fig. 3A, B). The deficiency was preserved when we replaced the

221 Sstr5 sequence with a non-OS-targeted photopigment, melanopsin (Opn4a), sequence (Matos-
222 Cruz et al., 2011, Davies et al., 2011) (Fig. S3A, B). Analysis of 4 additional hybrid constructs
223 revealed that transport efficiency correlates with the presence of IC3, [which is the equivalent](#)
224 [region where several other GPCRs' ciliary targeting motifs are also found](#) (Berbari et al., 2008,
225 [Loktev and Jackson, 2013, Nagata et al., 2013](#)) (Fig. 3A, B; constructs 4 - 7). Consistently,
226 replacement of this loop with the Sstr5 sequence resulted in a dramatic loss of transport
227 efficiency (Fig. 3A, B; compare no. 2 to 8). These studies show that IC3 is required for opsin
228 targeting to the OS.

229

230 The assay that we use to monitor opsin transport efficiency into the ciliary compartment
231 measures the rate of GFP signal loss in the photoreceptor cell body. In addition to OS-directed
232 transport, loss of GFP signal could be caused, however, by degradation of opsin/Sstr5-GFP
233 hybrid polypeptides in the cell body. To test this, we expressed four hybrid constructs: two
234 that are efficiently transported to the ciliary outer segment (no. 4 and no. 6) and two that are
235 not (no. 5 and no. 8) in photoreceptors of *oval* (*ovl*) mutants. We showed previously that
236 *ovl^{tz288b}* strain harbors a mutation in the *ift88* gene that causes complete loss of the photoreceptor
237 outer segment (Tsujikawa and Malicki, 2004). In the absence of the outer segment,
238 opsin/Sstr5-GFP hybrids remain trapped in the cell body (Fig. 3C). We compared
239 cytoplasmic expression levels of opsin/Sstr5-GFP hybrid constructs at 5 and 12h post heat
240 shock. We did not see statistically significant loss of GFP signal for any of the fusion
241 constructs tested (Fig. 3D). Some signal loss between 5 and 12h, albeit not statistically
242 significant, was seen for no. 8. For hybrid no. 6, a small but statistically significant increase
243 of signal was observed. These findings indicate that degradation is not a factor that
244 significantly contributes to the rate of signal loss from the photoreceptor cell body in our assay.
245 Moreover, we did not observe a relationship between the overall expression level and the rate
246 of GFP signal loss from the cell body: although fusion no. 8 was expressed at a higher level
247 compared to other constructs, no. 5 was not and yet it was inefficiently transported to the outer
248 segment (Fig. 3D). Finally, we did not see a statistically significant [photoreceptor](#) cell loss

249 for any construct between 5 and 12 hours post heat-shock (Fig. S3E). Cell loss is thus unlikely
250 to account for differences in **cell body** GFP signal level. Taken together, these findings reveal
251 that opsin transport into the outer segment requires a targeting sequence in the IC3 of opsin.

252

253 To define sequences sufficient to mediate outer segment transport, we inserted opsin IC3 and
254 CT44 into Sstr5 (construct no. 9) (Fig. 4). This hybrid GPCR fails to localize to the outer
255 segment, indicating that these two fragments of opsin sequence are not sufficient to target opsin
256 to the OS. Similarly, a hybrid, that in addition to IC3 and CT44, contains helix 6 from opsin
257 (Fig. 4A, C; no. 11) is not efficiently targeted to the OS and substitution of opsin helix 6 in
258 hybrid no. 6 with the Sstr5 sequence (Fig. 4A; hybrid no. 13) also impairs targeting to the ciliary
259 compartment (Fig. 4C). These results suggest that rod opsin transport mechanism involves
260 both helices 6 and 7. Coherently, based on the opsin crystal structure (Park et al., 2008,
261 Palczewski et al., 2000), these helices are close to each other and may interact to maintain opsin
262 conformation and the relative positions of IC3 and CT44. To narrow down which parts of
263 helix 6 and 7 are required for the ciliary targeting of opsin, we constructed a hybrid containing
264 the N-terminal half of helix 6 and the C-terminal half of helix 7 from rod opsin in addition to
265 opsin IC3 and CT44 (Fig. 4A, no. 14). We found that this configuration of opsin sequences
266 is sufficient to mediate outer segment targeting with the same efficiency as the full-length
267 sequence (Fig. 4C).

268

269 Studies of SSTR3 identified a conserved motif in IC3 as necessary for ciliary targeting (Berbari
270 et al., 2008). To test whether it is also required for opsin targeting, we deleted this motif
271 (AAAQQQ) from the full length rod opsin (construct no. 12, Fig. 4B). In contrast to SSTR3,
272 this deletion does not affect OS targeting in photoreceptors (Fig. 4D). Thus, we were
273 interested in which particular motif within IC3 is essential for its role in regard to OS targeting,
274 as the common ciliary targeting motif appeared to be redundant for transport. To address this
275 question, two additional deletion constructs (Fig. 4B and D; no. 16 and no. 17) were tested and
276 the OS targeting defect was only observed in the absence of RAERE sequence, which is

277 adjacent to helix 6, but the ESETTQ motif had no effect on opsin transport. It is possible, that
278 the defect in OS targeting observed in the RAERE mutant could simply be explained by a defect
279 in protein folding and transport. To address this, we expressed these opsin variants in PAC2
280 cells and visualized their localization. All of the above opsin variants are able to escape from
281 the endoplasmic reticulum (ER) and reach the plasma membrane as the full-length opsin (Fig.
282 S4), suggesting that their mislocalization was not due to protein misfolding and ER retention.
283 Taken together, these results define CT44, IC3 and parts of helixes 6 and 7 as both necessary
284 and sufficient to target opsin to the ciliary outer segment in vertebrate photoreceptors.

285

286 **Discussion**

287

288 Cilia-targeting sequences have been identified for several proteins, including Polycystins, CNG
289 channel, INPP5E and several GPCRs, including opsin (reviewed in Malicki and Avidor-Reiss,
290 2014). However, there does not appear to be a common ciliary targeting motif. GPCRs are
291 to be thought to be transported to cilia by multiple mechanisms: while the C-terminal
292 cytoplasmic tail appeared sufficient for opsin targeting to the OS, entirely different targeting
293 motifs in the third cytoplasmic loop were shown to target other GPCRs to the ciliary
294 compartment (Loktev and Jackson, 2013, Mukhopadhyay et al., 2013, Berbari et al., 2008).
295 Targeting motifs found in other GPCRs appeared irrelevant to opsin because the opsin
296 cytoplasmic tail seemed sufficient to target *Xenopus* opsin to the ciliary compartment (Tam et
297 al., 2000). Conversely, opsin C-terminal targeting motifs, such as VxPx, are poorly conserved
298 in SSTRs, if at all, supporting the idea that opsins utilize different targeting mechanisms,
299 compared to other GPCRs. In addition, the VxPx motif was shown to have negative effects
300 on primary cilia targeting (Geneva et al., 2017), suggesting distinct transport mechanisms act
301 in different cell types. Our data establish, however, that in our model system the third
302 intracellular loop of opsin is essential for the targeting towards photoreceptor ciliary
303 compartments, implying that at least some aspects of opsin transport are closely related to those
304 of other GPCRs.

305

306 The (AX[S/A]XQ) motif, previously identified in other GPCRs and conserved in the opsin
307 sequence, is not, however, necessary for opsin transport in our assay. This is not entirely
308 unexpected as ciliary targeting motifs in the IC3 of GPCRs vary substantially: the [R/K][I/L]W
309 motif of NPY2R and [I/V]KARK of GPR161 are not related to the ciliary targeting sequence
310 of SSTR3 and do not display clear similarity to each other apart from the presence of basic and
311 hydrophobic residues (Mukhopadhyay et al., 2013, Loktev and Jackson, 2013). In addition,
312 in agreement with our analysis, mutating the SSTR3-related motif does not affect the ciliary
313 trafficking of GPR83 (Loktev and Jackson, 2013). Rod opsin is thus likely to have yet another
314 set of targeting motifs in IC3. Indeed, our data suggests that the RAERE motif in the IC3 is
315 necessary for opsin OS targeting. Work of several groups' has shed the light on proteins that
316 facilitate opsin transport, including Arf4 (Deretic et al., 2005, Mazelova et al., 2009), ASAP1,
317 Rab8 and Rab11 (Wang et al., 2012) and IFTs (Keady et al., 2011, Crouse et al., 2014). It has
318 been shown that Arf4 binds to the VxPx motif and ASAP1 binds to the FR motif at the opsin C
319 termini (Fig. 5E). However, some of these components appear to be dispensable in vivo for
320 opsin transport (Pearing et al., 2017, Ying et al., 2016). Our data suggests the IC3 potentially
321 could be a new binding site for trafficking machinery. [The IC3 was suggested as a binding
322 site for TULP3 and RABL2, which have been shown to act as adaptors \(Barbeito et al., 2020,
323 Badgandi et al., 2017\) for transport of several GPCRs to the primary cilium.](#) In the future, [the
324 role of IC3](#) as an interacting site [for opsin transport](#) will need to be further addressed by
325 biochemical approaches.

326

327 Our studies show that in addition to targeting motifs in the third intracellular loop, parts of
328 helices 6 and 7 on the intracellular side of the membrane also contribute to opsin transport
329 (summarized in Fig. 5). In crystal structures (Palczewski et al., 2000, Park et al., 2008), these
330 helices localize close to each other (Fig. 5B-D) and their interactions may be necessary to
331 position IC3 and/or helix 8 in a way that facilitates interactions with components of the ciliary
332 transport machinery (Fig. 5E). Interestingly, both helix 6 and helix 7 contain residues

333 (Met253, Met257, Ile305 and others Fig. 5D), that are well conserved among ciliary but not
334 rhabdomeric opsins (Arendt et al., 2004), supporting the conclusion that these helices are
335 involved in OS targeting. Additionally, we show that segments of both helices are required
336 to facilitate efficient OS targeting, consistent with the hypothesis that their interaction with each
337 other is required.

338

339 A surprising conclusion of our analysis is that *Xenopus* C-terminal sequence functions as an
340 OS targeting signal in zebrafish photoreceptors while the homologous zebrafish sequence does
341 not. The alignment of opsins from vertebrates (Fig. S1) shows that Met317 is conserved in
342 all species examined, except *Xenopus*, where it is substituted by leucine. This suggests that
343 *Xenopus* opsin transport may be different from other vertebrates. Interestingly, we also
344 observed that the human sequence and its leucine variant behave in the same way as the
345 zebrafish constructs (Fig. S2B-D). At present it is unclear why the leucine substitution could
346 make such dramatic changes for OS targeting.

347

348 One possibility is that the *Xenopus* C-terminal tail has unusually strong propensity to bind
349 components of opsin transport machinery, whereas additional sequences are required for
350 zebrafish and other opsins to achieve sufficient interactions. This would explain why
351 *Xenopus* CT44 is efficiently targeted to the outer segment in zebrafish, but the homologous
352 sequence from zebrafish does not behave in this way. *Xenopus* opsin transport may need to
353 be significantly more efficient because the *Xenopus* rod OS is ~30 fold larger than the mouse
354 rod OS. In addition, the replenishment rate of the OS is higher in *Xenopus* (~8 fold higher
355 than in the mouse) compared to other species (Pearing et al., 2013).

356

357 An additional explanation for the ability of the *Xenopus* opsin C terminus to more efficiently
358 target to the OS may be linked to the likelihood of opsin dimer formation. Several lines of
359 evidence show that helix 8 together with helix 1 form one of the possible interfaces that mediate
360 interactions between opsin molecules. First, an opsin homodimer crystal structure (PDB ID

361 3CAP) shows that helices 8 and 1 are positioned at the dimerization interface (Park et al., 2008).
362 Second, chemical cross-linking in bovine rod outer segment membranes identified a Cys316-
363 Cys316 cross link between helices 8 of two monomers (Knepp et al., 2012), again suggesting
364 that helix 8 mediates dimerization. Notably, Cys316 is adjacent to Leu317 that we identified
365 as essential for efficient XCT44 targeting to the OS. Finally, delivery of helix 8 peptide to
366 mouse photoreceptors mislocalizes opsin, most likely due to interference with dimerization
367 (Zhang et al., 2016). The alignment of opsins from vertebrates (Fig. S1) shows that Met317
368 is conserved in all species examined, except *Xenopus*, where it is substituted by leucine. This
369 reveals that *Xenopus* opsin transport involves molecular features not shared with other
370 vertebrates. In the light of evidence that helix 8 mediates opsin dimerization, an intriguing
371 possibility is that the *Xenopus* C-terminal tail has unusually strong propensity to dimerize and
372 consequently is co-transported, or “hitch-hikes,” with the full-length wild-type protein. This
373 would explain why *Xenopus* CT44 is efficiently targeted to the outer segment in zebrafish, but
374 the homologous sequence from zebrafish does not behave in this way. A caveat is that
375 multiple intradimeric contacts, including helix 1 and 2, 1 and 5, 1 and 8, 4 and 5 etc., have been
376 identified by studies using electron microscopy, atomic force microscopy as well as
377 biochemical tools (Knepp et al., 2012, Jastrzebska et al., 2015, Liang et al., 2003, Ploier et al.,
378 2016, Salom et al., 2006). Thus, whether opsin transport is in the form of dimer/oligomer,
379 and whether helix 8 participates in, or is sufficient for, dimerization, are questions that remain
380 to be addressed.

381

382 It is also possible that zebrafish opsin transport is different to that of higher vertebrates. For
383 example, the zebrafish C-terminal sequence does not function well on its own or fused to the
384 transmembrane domain of other GPCRs. However, the mouse rod opsin C-terminal sequence
385 is able to direct part of peripherin, which lacks the C-terminal targeting signal, to the OS
386 (Salinas et al., 2017, Tam et al., 2004, Salinas et al., 2013). At present, no study has directly
387 tested how an isolated C-terminus behaves in mouse photoreceptors. However, it is worth
388 noting that the RAERE motif is conserved in higher vertebrates as KAEKE and it is commonly

389 shared within ciliary opsins but not within rhabdomeric opsins (Arendt et al., 2004). In the
390 future, it will be important to directly test whether these residues are critical for opsin transport
391 in higher vertebrates and if they play a role in binding transport machinery.

392

393 **Materials and Methods**

394

395 **Zebrafish strains and maintenance**

396 Zebrafish were maintained in the University of Sheffield aquarium facility in accordance with
397 UK Home Office requirements. Embryos of AB, LWT (London Wild-type) or Nacre lines
398 were used for transient expression assays and transgenesis. The *ov1^{tz288b}* strain was described
399 by us previously (TsujiKawa and Malicki, 2004).

400

401 **DNA constructs**

402 Two transgenes were used to obtain spatial and temporal control of expression in photoreceptor
403 cells: the Cre expression transgene, and a conditional EGFP-CT44 transgene. To obtain rod
404 specific Cre expression, we cloned the 1.3 kb BglII-BamHI fragment from the XopsEGFP-N1
405 plasmid (Fadool, 2003), containing the *Xenopus* rod opsin promoter, into the p5E-MCS vector.
406 The resulting entry clone: p5E-XOPS-1324, pME-Cre, and p3E-PolyA, were assembled into a
407 pDestTol2CG2 destination vector, using Gateway reaction as described (Kwan et al., 2007).
408 To construct conditional EGFP-S-CT44 transgenes, we synthesized a DNA fragment encoding
409 a C-terminal fragment of EGFP fused to a 2xTEV-S-tag-CT44-polyA sequence (Integrated
410 DNA Technologies, IDT) and combined it by overlap extension PCR with a lox2272-EGFP
411 sequence. The PCR amplification product was inserted into the Gateway entry vector p3E-
412 MCS (Invitrogen, Inc.). The resulting entry clone was combined with p5E-hsp70 (Kwan et
413 al., 2007), and pME-lox2272-mCherry(stop) in the pDestTol2pA2 destination vector, using the
414 Gateway reaction. Three versions of the conditional EGFP-CT44 transgene were made,
415 containing zebrafish CT44 (CT44), *Xenopus* CT44 (XCT44), or a random 44 amino acid
416 control sequence (CTC). The transgenes are termed as follows:

417 Tg(XOPS-1324:Cre),
418 Tg(hsp70: lox2272-mCherry-lox2272-EGFP-2xTEV-S-tag-XCT44),
419 Tg(hsp70: lox2272-mCherry-lox2272-EGFP-2xTEV-S-tag-ZCT44),
420 Tg(hsp70:lox2272-mCherry-lox2272-EGFP-2xTEV-S-tag-CTC)

421

422 To test zebrafish and Xenopus CT44 cilia targeting sequences in transient expression assays,
423 we first inserted EGFP between HindIII and EcoRI sites in the pME-MCS Gateway vector from
424 the zebrafish Tol2 transgenesis kit (Kwan et al., 2007). The zebrafish CT44 sequence was
425 then inserted downstream of EGFP between EcoRI and BamHI sites. The heat-shock
426 promoter and the polyA site were added using p5E-hsp70 and p3E-polyA entry clones from the
427 same kit. We used a Gateway reaction to assemble the entry clones into the pDestTol2-CG2
428 destination vector, which includes *cmlc2:eGFP* as a transgenesis marker. Zebrafish/Xenopus
429 hybrid CT44 sequences were assembled by PCR using forward primers encoding Xenopus or
430 Zebrafish Helix 8 or their mutant variants (Fig. 2). Zebrafish or Xenopus CT44 sequence was
431 used as a template. Human constructs, AAR and their variants were made following the above
432 strategy. Human and AAR sequences were obtained through DNA synthesis.

433

434 For opsin/Sstr5 hybrid experiments, the opsin reference construct was prepared by inserting
435 EGFP after T336 in the ASTTASK sequence in zebrafish rod opsin (UniProtKB - P35359).
436 An additional serine was introduced in this process generating the following insertion: AST-
437 (EGFP)-STASK. This resulted in a fusion similar to that previously used to test bovine opsin
438 transport (Moritz et al., 2001). The sstr5 reference construct was generated by appending
439 EGFP to the C-terminus of the zebrafish sstr5 sequence (UniProtKB - Q6NV10), which was
440 amplified from zebrafish cDNA. The melanopsin reference construct was cloned using the
441 same procedure as for sstr5 one. The zebrafish melanopsin gene (Gene ID: 571624) was
442 amplified from a cDNA clone (Source Bioscience, IRCYp5023J172D). Opsin-EGFP, Sstr5-
443 EGFP and melanopsin-EGFP constructs were inserted into EcoRI/BamHI (opsin),
444 HindIII/BamHI (sstr5) or XhoI/NotI (melanopsin) sites of the pME-MCS vector. All

445 opsin/sstr5 and opsin/melanopsin hybrid constructs were made by overlap extension PCR and
446 inserted into either EcoRI/BamHI (No5, 8, 12, 16, 17) or HindIII/BamHI (No3, 4, 6, 7, 9, 11,
447 13, 14) or XhoI/BamHI (ML-No3) sites of the same vector. To generate the final heat shock
448 promoter expression constructs, three entry clones, p5E-hsp70, pME-opsin/sstr5 and p3E-
449 polyA were combined as above. The structure of all final constructs was confirmed by
450 sequencing. Full amino acid sequences of opsin/Sstr5 and opsin/melanopsin are listed in
451 supplementary [Table S1](#).

452

453 **Embryo Injections and Transgenesis**

454 To generate stable transgenic lines and to perform transient expression assays, constructs were
455 microinjected with Tol2 transposase as described (Kwan et al., 2007). While performing
456 transient expression assays, in each experiment all constructs analyzed were injected during a
457 single session. To obtain stable transgenic lines, XOPS-1324-Cre transgenic fish were
458 selected at 3 dpf on the basis of *cmlc:GFP* expression. To generate fish transgenic for
459 conditional EGFP-S-CT44 constructs, G0 larvae were screened 3-4 days after injection by heat
460 shocking and selecting for mCherry expression 8 hours post heat shock, using a high power
461 fluorescence dissecting microscope. The same procedure was used in the F1 generation and
462 mCherry-positive fish were raised to adulthood. Single transgene insertions that segregate in
463 a Mendelian fashion were identified in subsequent generations.

464

465 **Transient expression assays**

466 In transient expression assays, four-day wild-type larvae were heat-shocked at 39°C for 30
467 (CT44 analysis) or 10 (opsin/Sstr5 assays) or 20 min. (opsin/melanopsin). Larvae with GFP-
468 positive hearts were selected 5-6 h after heat shock and fixed 24h post heat shock. Fixation,
469 cryosectioning and imaging was performed as described (Malicki et al., 2016). Prior to
470 imaging, sections were counterstained for 10 min. in DAPI solution. For analysis in *ovl*^{-/-}
471 mutant background, opsin/sstr5 hybrid constructs were injected into embryos originating from

472 a *ovl*^{+/-} heterozygotes cross. Homozygous mutants were heat shocked at 3 dpf for 10 min. and
473 fixed 5 or 12 hours post heat shock.

474

475 **Assays using stable transgenic lines.**

476 For the analysis of opsin transport in wild-type retinæ, single or double transgenics were heat
477 shocked at 4 dpf for 30 (EGFP-ZCT44) or 5 (EGFP-XCT44) minutes and analyzed at 5 hr and
478 24 hr post heat shock. Different heat shock times were used to compensate for the difference
479 in the expression level of ZCT44 and XCT44 transgenes due to positional effects of transgene
480 insertion. mCherry-positive larvae with green hearts were selected and processed for
481 cryosectioning as above.

482

483 **Image analysis**

484 Single optical sections in the plane of retinal cryosections were acquired on an Olympus
485 FV1000 Confocal microscope and saved as .oif format files, which were then processed using
486 Fiji software. Signal intensity from the GFP channel was color-coded using a thermal lookup
487 table. To measure signal intensity in photoreceptor cell bodies, a line, approximately 2.5 μm
488 wide, was drawn from the synapse to the outer segment base for each GFP-positive
489 photoreceptor using Fiji software. Mean signal intensity in the area covered by the line was
490 quantified and the resulting images were flattened and stored as TIFF files. To avoid signal
491 overlap from neighboring cells, isolated GFP-positive cells were preferred for this analysis.

492

493 **Statistical Analysis**

494 To analyze cytoplasmic GFP intensity in stable transgenic lines (Fig. 1), we log transformed
495 signal intensity data and analyzed them using 2-way ANOVA. Similarly, all other GFP
496 intensity data were log transformed. The resulting data sets approximated normal distribution
497 and were analyzed using ANOVA of mixed effects models assuming fish as a random effect.
498 Mann-Whitney test was used for data sets containing signal intensity, which was averaged per
499 animal. * < 0.05; ** < 0.01; *** < 0.001; **** < 0.0001.

500

501 **Cell culture and immunofluorescence**

502 PAC2 cells (Martin Lowe, University of Manchester, UK) were cultured at 28 °C with 0.1%
503 CO₂ and L-15 medium containing 10% FBS and 1% Penicillin/Streptomycin. Transfections
504 were performed according to the manufacturer's protocol for FuGENE HD transfection reagent
505 (Promega E2311). Heatshock was applied 4 hours post transfection at 39 °C and cells were
506 fixed by 4% PFA/PBS 48 hours after heatshock followed by 0.1% Triton X-100/PBS
507 permeabilization. Permeabilization was not applied if followed by 4D2 staining. 1%
508 BSA/PBS was used for blocking and cells were stained by the following antibodies: anti-
509 GRP78/Bip (Eames et al., 2013) (1:200, Abcam), anti-Rhodopsin Rho-4D2 (Salinas et al., 2017)
510 (1:100, Abcam) and Alexa Fluor 568 anti mouse or rabbit conjugated secondary antibody
511 (1:500, Invitrogen). Anti-GRP78/Bip antibody was applied to label the ER. Anti-Rho-4D2
512 antibody recognizes the N terminus of rod opsin. Without permeabilization, anti-Rho-4D2
513 only detects opsin molecules that reach the plasma membrane.

514

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520

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525

526 **Conflict of Interest Statement**

527 The authors declare no conflict of interest.

528

529 **Data Availability**

530 All data are made available in the paper.

531

532

533

534 **References**

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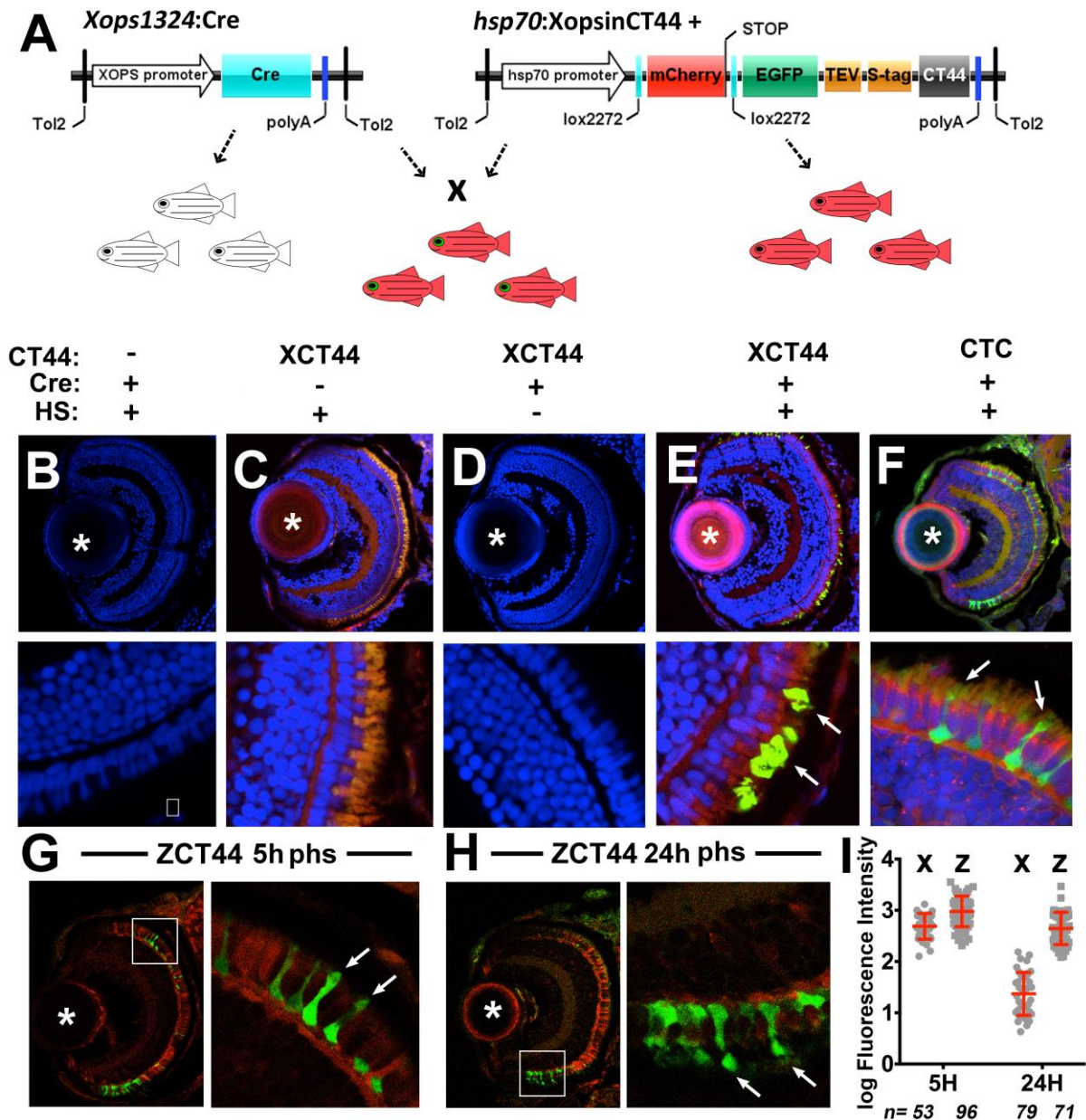
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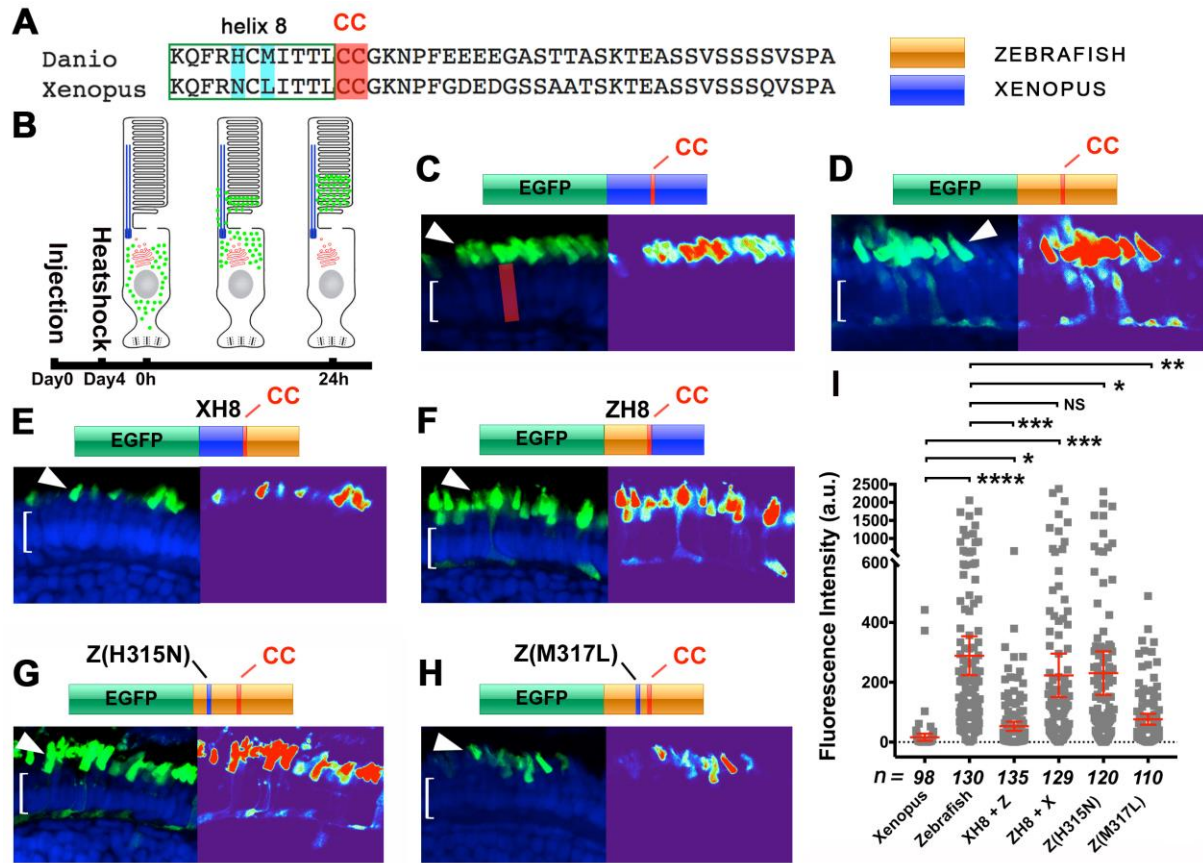
701

702 **Figures**



703
 704 **Fig. 1** (A) Schematic of transgenes used to obtain spatial and temporal control of expression
 705 for EGFP-S-opsinCT44 fusion proteins. (B-H) Confocal images of cryosections through the
 706 retinae of transgenic zebrafish larvae at 5 dpf. In (B-F), lower panels show higher
 707 magnification images. Transgene genotypes and heat shock treatment are indicated above.
 708 CTC, control random 44 amino acid peptide. (B) No fluorescent signal is observed in the Cre
 709 line even when heat shock is applied. (C) mCherry but not GFP expression is present when
 710 heat shock is applied to the *hsp70:(mCherry)EGFP-S-opsinXCT44* transgenic line. (D) No
 711 fluorescent signal is observed in the double transgenic line when heat shock is not applied. (E)

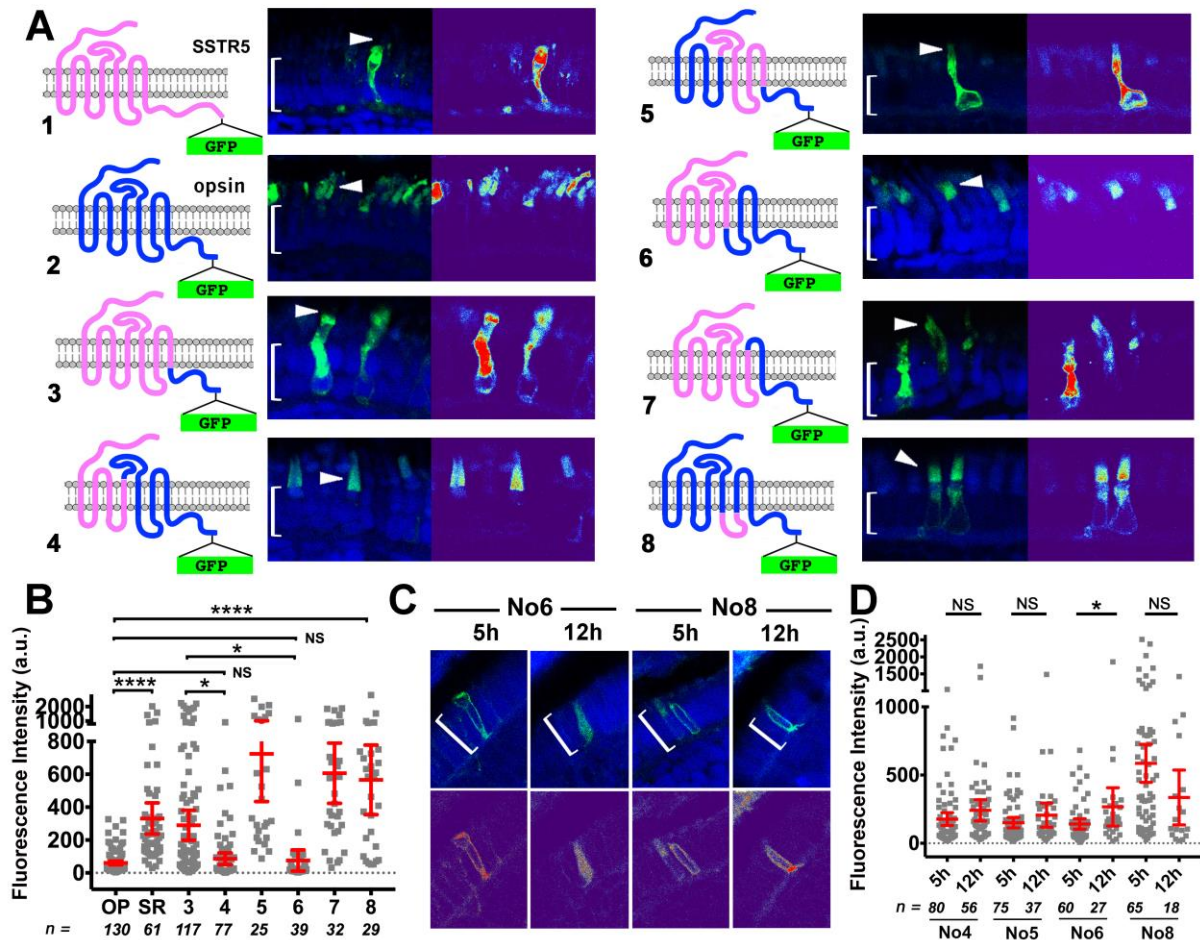
712 EGFP is expressed in photoreceptors but not in other retinal cells when heat shock is applied to
713 the double transgenic line containing Tg(XOPS-1324:Cre) and Tg(hsp70: lox2272-mCherry-
714 lox2272-EGFP-2xTEV-S-tag-XCT44) transgenes. EGFP signal localizes to outer segments
715 in this line (arrows in lower panels). (F) In contrast, EGFP signal remains in the cell body in
716 a double transgenic line that contains a control EGFP construct in which CT44 cilia targeting
717 signal was replaced with a random peptide. (G, H) High level of EGFP-S-opsinZCT44
718 persists in the photoreceptor cell body at 5 (G) and 12 (H) hours after heat shock. Panels to
719 the right show enlargements of photoreceptors from images presented to the left. (I) GFP
720 signal intensity in the cell body at 5h and 24h post heat shock for Xenopus (X) and zebrafish
721 (Z) GFP-CT44 fusions. [Log-transformed data from 2 independent experiments](#) are shown.
722 Each dot represents a measurement from a single photoreceptor cell. GFP is efficiently
723 transported from the cell body to the outer segment when using the Xenopus but not the
724 zebrafish CT44 ($p < 0.001$, ANOVA). In (B-F) sections are counterstained with DAPI (blue).
725 Asterisks indicate the lens.
726



727

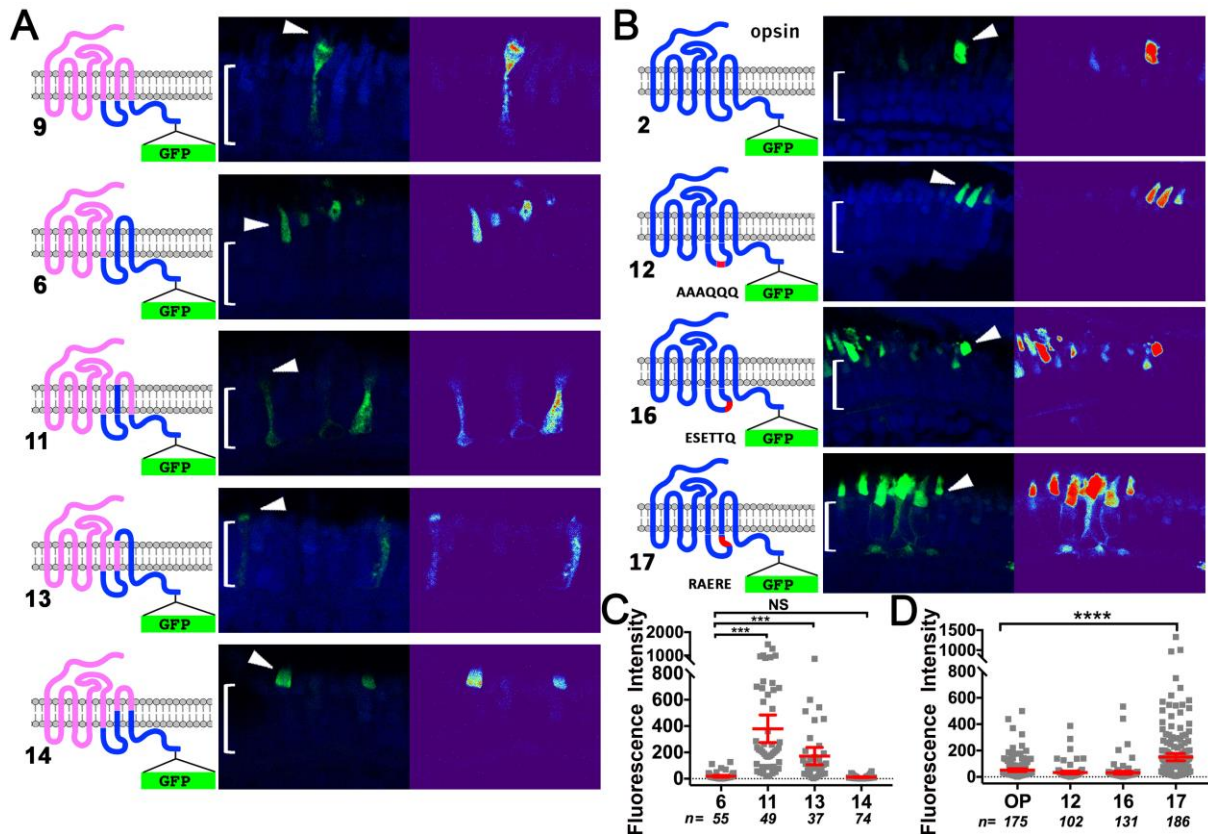
728 **Fig. 2** Comparison of OS-targeting efficiency of Xenopus and zebrafish opsin C-terminal tails
 729 (CT44). (A) Alignment of rod opsin 44 C terminal amino acids from zebrafish and Xenopus.
 730 Palmitoylated cysteines are highlighted in red. (B) A schematic demonstrates the process of
 731 the transient assay. (C-H) Confocal images of cryosections through the photoreceptor cell
 732 layer of wild-type zebrafish retinae, that transiently express the following variants of the EGFP-
 733 CT44 construct at 5 dpf: (C) XCT44; (D) ZCT44; (E) ZCT H8 + X; (F) XCT H8 + Z; (G) Z-
 734 H315N; (H) Z-M317L. Confocal image (left) and a heat map (right) of GFP signal intensity
 735 are juxtaposed. Brackets indicate the photoreceptor cell layer, arrowheads point to outer
 736 segments. (I) Quantification of fluorescence intensity in photoreceptor cell bodies. The red
 737 rectangle in (C) approximately shows the soma of a single photoreceptor cell. Each dot
 738 represents a measurement from a single photoreceptor cell. Data are from 3 independent
 739 experiments. Mean and 95% confidence interval are indicated. Sample sizes are provided
 740 in italics below the horizontal axis. In schematic drawings above panels (C-H), zebrafish and
 741 Xenopus sequences are color-coded in orange and blue, respectively. Red bars indicate

742 conserved cysteine residues in opsin C terminus. Data are log transformed for statistical
 743 analysis as described in methods.
 744

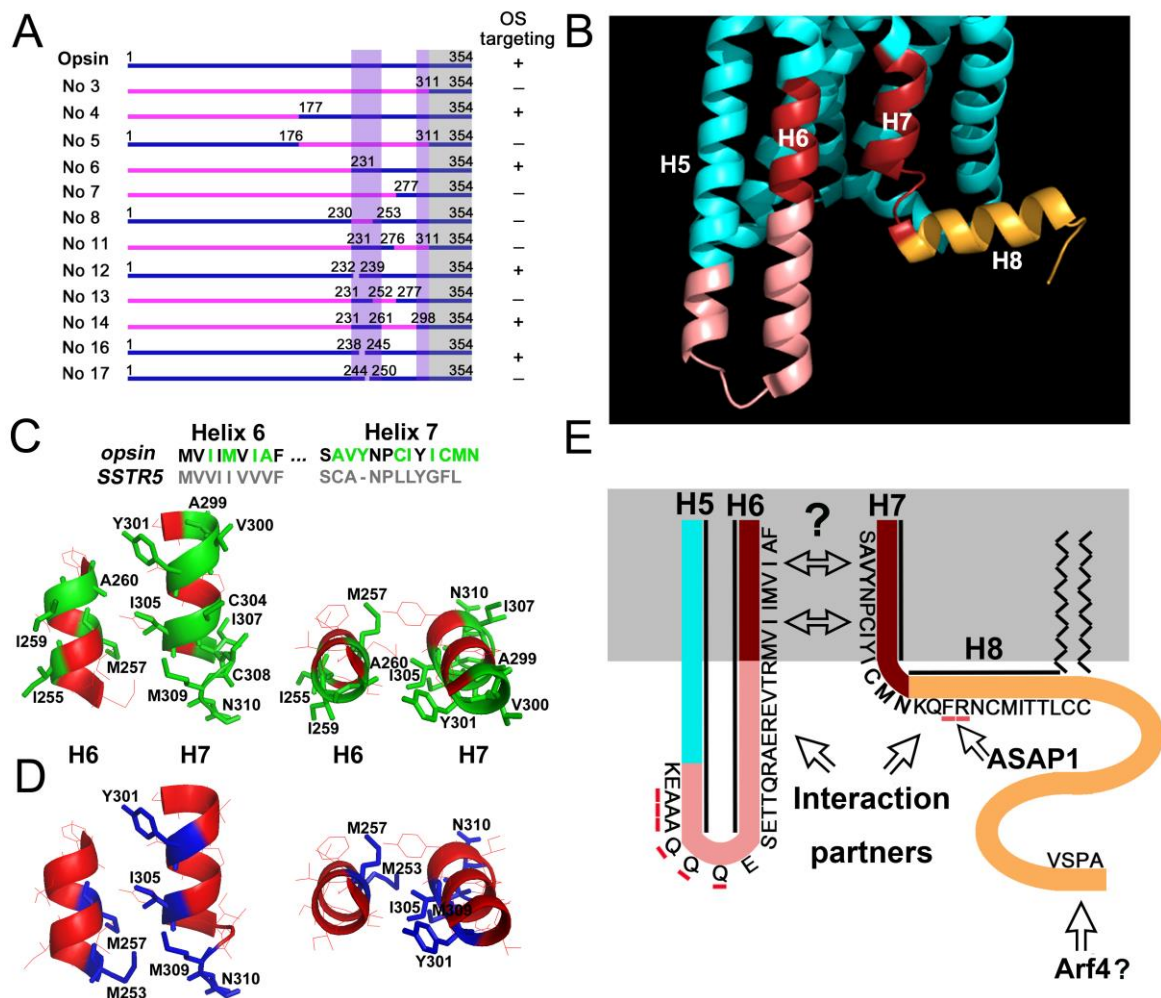


745
 746 **Fig.3** Transport efficiency of opsin/Sstr5 hybrid GPCRs into photoreceptor ciliary
 747 compartment. (A) Images of cryosections through zebrafish retinas expressing wild-type or
 748 hybrid GPCRs schematically shown to the left of each image. Brackets indicate the
 749 photoreceptor cell layer, arrowheads point to outer segments. In each panel, a confocal image
 750 (left) and a heat map (right) of GFP signal intensity are shown. (B) Quantification of the GFP
 751 signal intensity in photoreceptor cell bodies for wild-type and hybrid GPCRs shown in (A).
 752 Data from 4 independent experiments are provided. (C) Images of cryosections showing the
 753 expression of two hybrid GPCRs in *oval*^{-/-} mutant photoreceptors at 5 and 12 h post heat shock
 754 as indicated in above panels. A confocal image (top) and a heat map (bottom) of GFP signal
 755 intensity are shown. (D) Quantification of GFP signal intensity in the cell bodies of *oval*^{-/-}

756 mutant photoreceptors expressing 4 hybrid GPCRs at 5 and 12 h post heat shock as indicated.
 757 Data are from 4 independent experiments. In (B, D) each dot represents a single photoreceptor,
 758 mean and 95% confidence interval are provided, sample sizes are in italics below the horizontal
 759 axis.
 760



761
 762 **Fig. 4** Transport efficiency of opsin/Sstr5 hybrid GPCRs (continued from Fig. 3).
 763 (A, B) Images of cryosections through zebrafish retinæ expressing wild-type or hybrid GPCRs
 764 schematically shown to the left of each image. Brackets indicate the photoreceptor cell layer,
 765 arrowheads point to outer segments. In each panel, a confocal image (left) and a heat map
 766 (right) of GFP signal intensity are shown. (C) Quantification of the GFP signal intensity in
 767 photoreceptor cell bodies for hybrid GPCRs shown in (A). (D) Quantification of the GFP
 768 signal intensity in photoreceptor cell bodies for hybrid GPCRs shown in (B). In (C, D), data
 769 originate from 3 independent experiments, each dot represents a single photoreceptor, mean
 770 and 95% confidence interval are shown, sample sizes are in italics below the horizontal
 771



772

773 **Fig. 5** Summary of opsin OS targeting motifs identified in this and previous studies. (A)

774 Summary of results from Figs. 3 and 4. Rod opsin sequences included in each construct are

775 represented as blue horizontal bars, Sstr5 sequences are depicted in pink. The first and the

776 last amino acid of each opsin fragment are numbered. Sequences necessary for the ciliary

777 targeting of opsin are highlighted by vertical bars. The CT44 is indicated by a grey vertical

778 bar. (B) The structure of opsin monomer based on X-ray crystallography (PDB: 3CAP).

779 Fragments of transmembrane helices near the cytoplasmic face of the membrane and

780 cytoplasmic loops are shown. Helix 8 is highlighted in orange, IC3 is in pink, and the portions

781 of helix 6 and 7 that we found necessary for opsin transport are red. The remaining sequences

782 are in cyan. (C, D) Regions of helix 6 and 7 necessary for opsin transport in side view and

783 viewed along helix axis. (C) Amino acid side chains that differ between zebrafish rod opsin

784 and zebrafish Sstr5 are numbered and depicted in green. (D) Amino acids that are conserved

785 in ciliary but not in rhabdomeric opsins are numbered and depicted in blue. (E) Schematic
786 representation of opsin features that are necessary and sufficient for the ciliary targeting of
787 opsin. Sequences are color coded as in (B). Helixes are indicated with solid black lines.
788 The SSTR3 ciliary targeting motif and the FR motif are underlined in red. Arrows indicate
789 potential interaction sites with proteins that facilitate opsin transport based on work of several
790 groups. In (C, D), two amino acid substitutions were made: V_{Bos304} to C_{Danio} and V_{Bos308} to
791 C_{Danio} to match the zebrafish sequence. In (B-D), protein structure is visualized using PyMOL
792 software.