Subcellular localization of the voltage-gated K+ channel EGL-36, a member of the KV3 subfamily, in the ciliated sensory neurons in C. elegans

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Abstract
Delineated as the first cellular organelle in 1675 by Antonie van Leeuwenhoek, cilia did not receive much attention until the 2000s, when it became apparent that cilia played a key role in the development of embryos, a variety of signaling pathways. Therefore, collective efforts by many scientists have led to the identification of many novel ciliopathy and cilia genes, while we are still far from disclosing the complete components of cilia. Here we used the ciliated sensory neurons in C. elegans as a model system that revealed the voltage-gated K+ channel EGL-36 (a member of the Shaw subfamily) as a new component associated with cilia. The confocal microscopy examination of fluorescence tagged EGL-36 together with ciliary (IFT-140) or transition zone (MKS-6) markers reveal that EGL-36 is only expressed in subsets of the ciliated sensory neurons, where it partially overlaps with the basal body signals and predominantly localizes to the periciliary membrane compartment. This expression pattern along with studies of egl-36 gain-of-function variants indicates that egl-36 is not essential for ciliogenesis in C. elegans. Our data identify the voltage-gated K+ channel EGL-36 as a new cilia-associated protein, and future studies should reveal the functional significance of EGL-36 in cilia biogenesis.
**Figure 1. EGL-36 is localized to the region proximal to the ciliary base in *C. elegans***: 

**A)** EGL-36::GFP specifically localizes at the cell soma and a compartment in the distal dendrite, just proximal to the ciliary base, in PHB not in PHA sensory neurons (left and middle columns). Shown are colocalization of EGL-36::GFP and IFT-140::mCherry, a ciliary marker, (a single copy). Scale bar: 5 μm.  

**B and C)** Co-expressions of EGL-36::GFP and MKS-6::mCherry, a transition zone marker (a single copy), is displayed in head (C, right column) and tail sensory neurons (B middle columns) in *C. elegans*. EGL-36::GFP is visible in PHB, not in PHA, and does not enter into cilia. 

**D)** An illustration showing the localization of EGL-36 in relation to cilia in the tail sensory neurons (PHA/PHB). IFT-140 and MKS-6 stain the entire ciliary axoneme and the transition zone, respectively. PCMC and TZ denote the periciliary membrane compartment and transition zone, respectively. A, anterior; P, posterior; D, dorsal; V, ventral. (L) Ventral view. 

**E)** Fluorescence images display lipophilic fluorescent dye uptake for wild type, *egl-36 (n728)*, and *egl-36 (n728n398)*. No obvious Dye uptake defects for these mutants. 

**F)** Localizations of OSM-6::GFP (IFT52) in PHA/PHB cilia were shown in wild type and *egl-36 (n728)* (no obvious phenotype). 

**Description**

*C. elegans* EGL-36 displays extensive similarity to human KCNC proteins, the conserved potassium channel subfamily KV3 (the Shaw subfamily). This potassium channel subfamily contains four genes KCNC1, KCNC2, KCNC3, and KCNC4 encoding KV3.1 (Shaker), KV3.2 (Shab), KV3.3 (Shaw), and KV3.4 (Shal), respectively. Previous studies have
demonstrated that C. elegans EGL-36 is expressed in the egg-laying muscles, ventral cord neurons, and the subset of ciliated sensory neurons including ADEL, ADER, PHB through its subcellular localization in the ciliated sensory neurons remains unknown (Elkes et al. 1997; Johnstone et al. 1997). To observe the subcellular localization of EGL-36 in the ciliated sensory neurons in C. elegans, we tagged green fluorescent protein (GFP) at the C-termini of EGL-36 with a cilia-specific promoter. We observed overexpressed EGL-36::GFP in the subsets of sensory neurons in the head and tail including PHB sensory neurons in the tail but not in the PHA sensory neuron, which is in line with previous findings (Elkes et al. 1997; Johnstone et al. 1997) (Figure 1A). Confocal laser scanning microscopy shows that EGL-36::GFP is predominantly located at the base of the cilium. The EGL-36::GFP signal can be found in the neuronal cell body, with weak dendrite staining, where they may function in the dendritic excitability (Figure 1A).

Cilia have several sub-compartments including the basal body, the transition zone, the middle segment, and the ciliary tip. We further examined the exact localization of GFP-tagged EGL-36 relative to these compartments. To this end, we used the transition zone marker MKS-6/CC2D2A and the cilia marker IFT-140 (intraflagellar transport 140), and co-expressed EGL-36::GFP with either of these markers (Qin et al., 2001; Williams et al., 2011; Prevo et al. 2015 Mijalkovic et al. 2018). Our microscopy analysis revealed that some of EGL-36::GFP signal was, however, found to overlay with the proximal zone of IFT signal (marked with IFT-140::mCherry), while the majority of EGL-36::GFP signal was predominantly localized in a compartment, proximal to the ciliary base (Figure 1A). This strong EGL-36 staining likely corresponds to the periciliary membrane compartment (PCMC) (Kaplan et al. 2012). There is no overlay of EGL-36::GFP signal with MKS-6::mCherry (Figure 1B and C).

We next went on characterizing potential ciliary functions of EGL-36 in C. elegans, we therefore obtained two mutants for egl-36 including egl-36(n728) and egl-36(n728n398). egl-36(n728) mutant contains a missense variant, conversion of glutamic acid at the 142nd position of EGL-36 to lysine (p.E142K). EGL-36(p.E142K) represents a gain of function variant of egl-36, which displays defects in egg-laying in C. elegans. On the other hand, proline to serine conversion (p.P439S) in egl-36 suppresses the egg-laying defect of p.E142K variant (Elkes et al., 1997; Johnstone et al., 1997). egl-36(n728n398) mutant contains both p.E142K and p.P439S variants. First, we performed the DiI-filling assay, which has been used to test ciliary structural defects in C. elegans. The lipophilic fluorescent dye is not taken up by C. elegans with abnormal cilia structures, while wild types always stain their ciliated sensory neurons with this fluorescent dye (Herman and Hedgecock 1990). The dye analysis coupled with microscopy imaging revealed that similar to wild type, both egl-36 mutants fill up their ciliated sensory neurons with the lipophilic fluorescent dye, suggesting that egl-36 mutants have no gross cilia abnormalities in C. elegans (Figure 1E). We next examined the cilia morphologies of phasmid (tail) sensory neurons in egl-36(n728) mutant. We generated transgenic strains expressing OSM-6/IFT52::GFP in egl-36(n728) mutants, and our microscopy analysis revealed that PHB (tail) displayed no abnormality in cilia morphology in egl-36(n728) mutants (Figure 1F).

**Methods**

Request a detailed protocol

**C. elegans** strains and maintenance

All strains used were grown in the nematode growth medium (NGM) at 20°C as previously described (Brenner, 1974).

**Generation of transgenic strains with microinjection**

The transgenic strain OIK904, N2;turEx12[Parl-13::EGL-36::GFP::unc-54 3’UTR+pRF4] was generated with microinjection of the Parl-13::EGL-36 (cDNA)::GFP::unc-54 3’ UTR construct (25 ng/μl) along with the rol-6 co-injection marker (50 ng/μl) by our laboratory.

**Dye-filling assays**

Dye filling assay previously described was used (Cevik et al., 2010; Herman and Hedgecock 1990).

**Fluorescent and Confocal microscopy**

Fluorescent images for the dye assay were obtained by a fully automated upright microscope system (Leica DM6 B) with a Plan ApoChromat 100x/1.40 NA and an electron-multiplying charge-coupled device camera (Andor iXon Ultra 897 EMCCD camera and iQ3.6.2 Andor software) that is attached to the microscope while confocal images were acquired with a laser-scanning confocal inverted microscope (Zeiss LSM 900 with Airyscan 2 and ZEN 3 Blue edition software) with a Plan ApoChromat 63x/1.40 NA objective.

**Reagents**

Strains and plasmid are available upon request to sebiha.cevik(at)agu.edu.tr

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N2  Caenorhabditis elegans  CGC
MT1540  egl-36(n728) X.  CGC
KP100  egl-36(n728n398) X.  CGC
SP2101  osm-6(p811);mnIs17(osm-6::gfp; unc-36(+).  CGC
OIK912  egl-36 (n728)x.; mnIs17(osm-6::gfp; unc-36(+).  This study
OIK904  turEx12[Parl-13::EGL-36::GFP::unc-54 3’UTR+pRF4]  This study
EJP81  vuaSi24 [pBP43; Pche-11::che-11::mCherry; cb-unc-119(+)]II; unc-119(ed3) III; che-11(tm3433)V. (This is referred IFT-140 throughout to the text)  Peterman Lab
DAM954  vuaSi24[pBP39; Pmks-6::mks-6::mCherry; cb-unc-119(+)II];mks-6(gk674) I (This is referred MKS-6 throughout to the text)  Dammermann Lab
OIK952  N2;turEx12[Parl-13::EGL-36::GFP::unc-54 3’UTR+pRF4];vuaSi21[pBP39; Pmks-6::mks-6::mCherry; cb-unc-119(+)]II. (This is referred IF1-40 throughout to the text)  This study
OIK953  N2;turEx12[Parl-13::EGL-36::GFP::unc-54 3’UTR+pRF4];vuaSi24 [pBP43; Pche-11::che-11::mCherry; cb-unc-119(+)]II; unc-119(ed3) III; che-11(tm3433)V. (This is referred IFT-140 throughout to the text)  This study

Plasmids  Genotype  Description
OK133  Parl-13::EGL-36 (cDNA) ::GFP::unc-54 3’UTR  Inserted Parl-13 (300 bp)- egl-36 (cDNA) with Sphl and Age I restriction enzymes by Sunybiotech
pRF4  rol-6(sa1006)  (Mello C et al. 1991)

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References


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