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## **Data Sheet**

### **TMEM16B (ANO2) - HEK293 Recombinant Cell line**

**Cat #: 90332**

#### **Product Description**

Recombinant HEK293 cell line expressing human TMEM16B (transmembrane protein 16B, also called as anoctamin 2, calcium-activated chloride channel (ANO2), accession number NM\_001278596).

#### **Format**

Each vial contains  $1.5 \times 10^6$  cells in 1 ml of 10% DMSO.

#### **Storage**

Immediately upon receipt, store vials in liquid nitrogen.

#### **Mycoplasma testing**

The cell line has been screened using the PCR-based Venor™GeM Mycoplasma Detection kit (Sigma-Aldrich) to confirm the absence of Mycoplasma species.

#### **Introduction**

Calcium-activated chloride channels (CaCCs) are involved in a variety of physiological functions including smooth muscle contraction and olfaction. TMEM16B (ANO2) has been identified as a CaCC that is activated by intracellular  $Ca^{2+}$  and  $Ca^{2+}$ -mobilizing stimuli. It has eight putative transmembrane segments and is permeable to monovalent anions.

#### **Functional validation**

Human TMEM16B channel has been stably expressed in HEK293 cell line and its expression was confirmed by Western blotting.

The CaCC activity of TMEM16B was characterized by an assay based on a halide-sensitive yellow fluorescent protein (YFP) mutant whose fluorescence is quenched by increasing halide concentration. When TMEM16B-expressed HEK293 cells were stimulated with ionomycin to raise the intracellular level of  $Ca^{2+}$ , TMEM16B produced  $I^-$  influx in HEK293, triggering a rapid decrease of fluorescence by the transfected YFP mutant. The ionomycin-induced  $I^-$  influx through TMEM16B was blocked by niflumic acid, a CaCC channel blocker.

This data shows the stable expression of TMEM16B channel in HEK293 cells.

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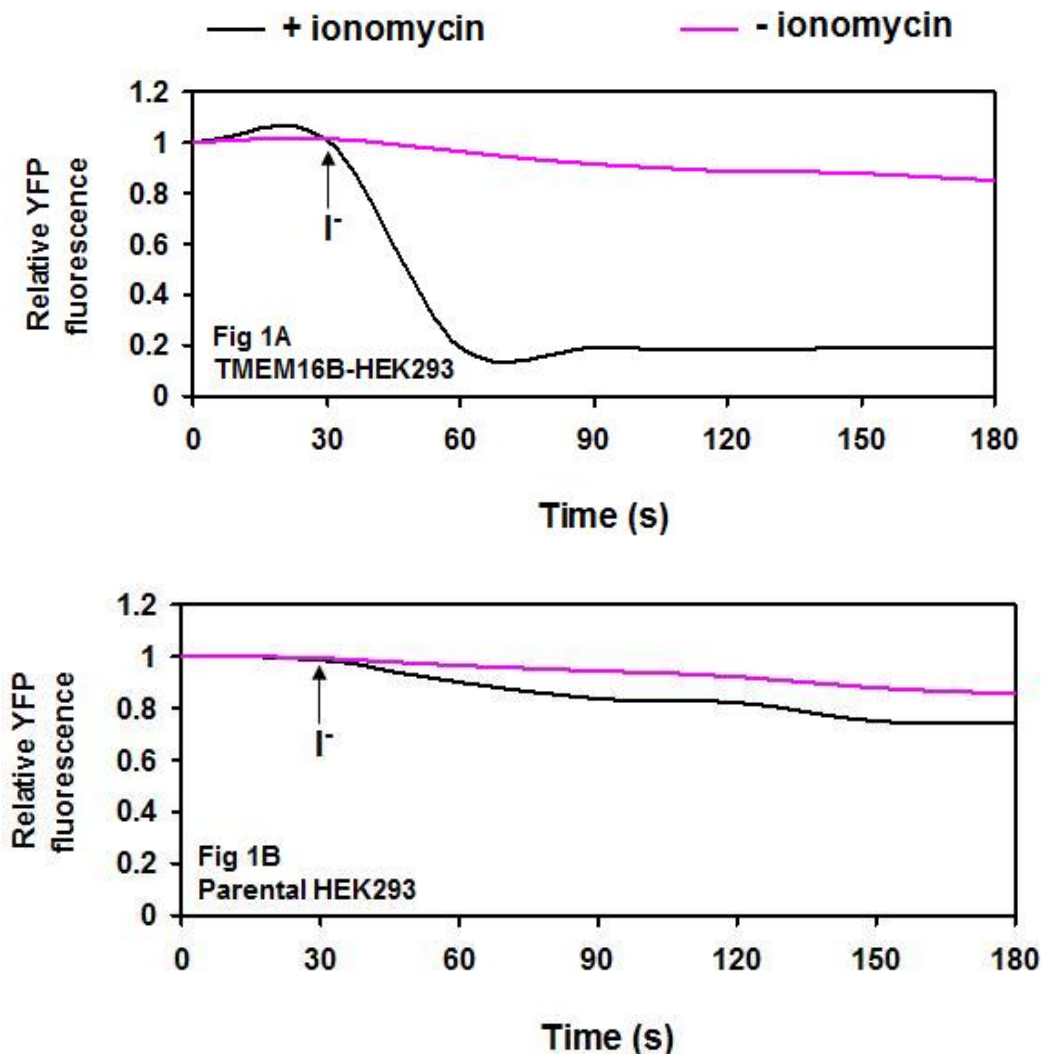
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**Figure 1. TMEM16B expressed in HEK293 produced I<sup>-</sup> influx after extracellular addition of I<sup>-</sup> with ionomycin. A) TMEM16B-HEK293; B) parental HEK293 cells.**

TMEM16B-HEK293 or parental HEK293 cells were transiently transfected with halide-sensitive YFP-H148Q/I152L mutant, then treated with I<sup>-</sup> (100 mM) saline solution (arrow) with (black) or without (pink) ionomycin (1 μM). I<sup>-</sup> influx was measured by YFP fluorescence (excited at 485±10 nm and emission at 528±10 nm). Results showed that following iodide addition, YFP fluorescence declined rapidly with ionomycin treatment in TMEM16B-HEK293 cells (but not parental HEK293 cells) due to I<sup>-</sup> influx through the TMEM16B channel.



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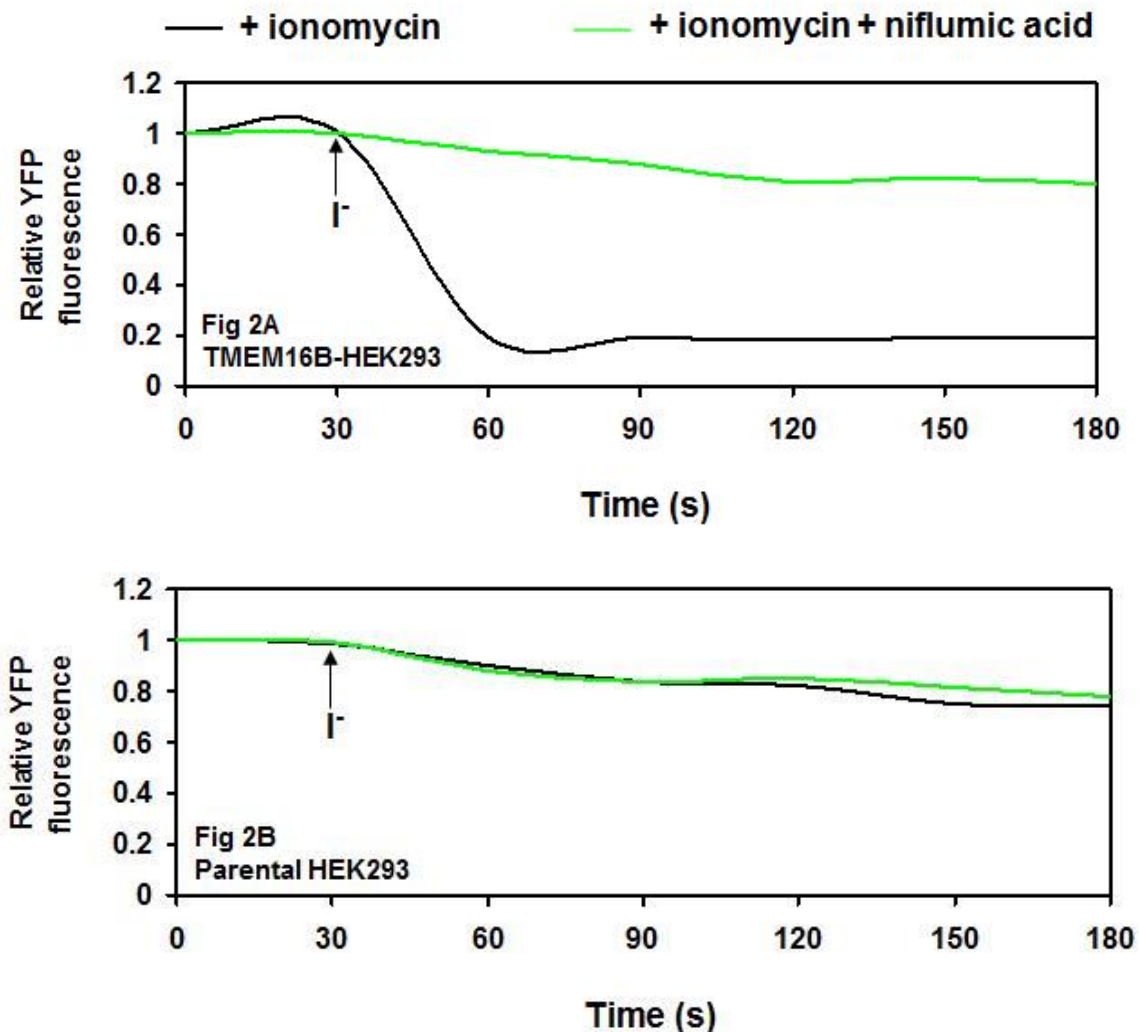
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**Figure 2. Ionomycin-induced  $I^-$  influx in TMEM16B-HEK293 cells was blocked by niflumic acid, a CaCC channel blocker. A) TMEM16B-HEK293; B) parental HEK293 cells.**

Cells were transiently transfected with halide-sensitive YFP-H148Q/I152L mutant, then treated with  $I^-$  (100 mM) saline solution plus ionomycin (1  $\mu$ M) (arrow) with (green) or without (black) pre-treatment of niflumic acid (100  $\mu$ M).  $I^-$  influx was measured by YFP fluorescence (excited at  $485 \pm 10$  nm and emission at  $528 \pm 10$  nm). Results showed that quenching of YFP fluorescence by ionomycin-induced  $I^-$  influx through TMEM16B was blocked by niflumic acid.



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## Culture Conditions

**Thaw Medium 1 (BPS Cat. #60187):** MEM medium (Hyclone #SH30024.01) supplemented with 10% FBS (Invitrogen #26140-079), 1% non-essential amino acids (Hyclone #SH30238.01), 1 mM Na pyruvate (Hyclone #SH30239.01), 1% Penicillin/Streptomycin (Hyclone SV30010.01)

**Growth Medium 1F (BPS Cat. #79540):** Thaw Medium 1 (BPS Cat. #60187) plus 100 µg/ml of Hygromycin B (Hyclone #SV30070.01) to ensure the recombinant expression is maintained. It may be necessary to adjust the percentage of CO<sub>2</sub> in the incubator, depending on the NaHCO<sub>3</sub> level in the basal medium. TMEM16B-HEK293 cells should exhibit a typical cell division time of approximately 24 hours.

It is recommended to quickly thaw the frozen cells from liquid nitrogen in a 37°C water-bath, transfer to a tube containing 10 ml of Thaw Medium 1 (**no Hygromycin B**), spin down cells, and resuspend cells in pre-warmed Thaw Medium 1 (**no Hygromycin B**). Transfer resuspended cells to T25 flask and culture at 37°C in a CO<sub>2</sub> incubator overnight. The next day, replace the medium with fresh Thaw Medium 1 (**no Hygromycin B**), and continue growing the culture in a CO<sub>2</sub> incubator at 37°C until the cells are ready to be split. Cells should reach ~60-80% confluence two days after being thawed. Cells should be split before they reach complete confluence. At first passage, switch to Growth Medium 1F (**contains Hygromycin B**).

To passage the cells, rinse cells with phosphate buffered saline (PBS), and detach cells from the culture vessel with 0.05% Trypsin/EDTA. Add Growth Medium 1F and transfer to a tube, spin down cells, resuspend cells and seed appropriate aliquots of cell suspension into new culture vessels. Subcultivation ration: 1:10 to 1:20 weekly or twice a week.

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### Vector and sequence

Human TMEM16B was cloned into pIRES-hyg vector (Clontech).

Polylinker: CMV-BsrGI-StuI-AflIII-NaeI-BssHII-NheI-**TMEM16B**-BamHI-EcoRV-BsiWI-BstXI-IRES-hygromycin<sup>R</sup>

hTMEM16B sequence (accession #NM\_001278596, UniProtKB/Swiss-Prot #Q9NQ90-1)

MATPGPRDIPLLPGSPRRLSPQAGSRGGQGPKHGQQCLKMPGPRAPGLQGGSNRDPGQP  
CGGESTRSSVINNYLDANEPVSLERLSRMHFHDSQRKVDYVLAYHYRKRGVHLAGQF  
PGHSLAIVSNGETGKPEHAGGPGDIELGPLDALEEERKEQREEFHNLMEAGLELEKDL  
ENKSQGSIFVRIHAPWQVLAREAEFLKIKVPTKKEMYEIKAGGSI AKKFS AALQKLSSH  
LQPRVPEHSNNKMKNLSYPFSREKMYLYNIQEKDTFFDNATRSRIVHEILKRTACSRAN  
NTMGINSLIANNIYEAAAYPLHDGEYDSPEDDMNDRKLLYQEWARYGVFYKFQPIDLIRK  
YFGEKIGLYFAWLGLYTSFLIPSSVIGVIVFLYGCATIEEDIPSREMCQQNAFTMCPL  
CDKSCDYWNLSACGTAQASHLFDNPATVFFSIFMALWATMFLENWKRLQMR LGYFWDL  
TGIEEEEEERAQEH SRPEYETKVREKMLKESNQSAVQKLETNTTECGDEDEDKLTWKDR  
FPGYLMNFASILFMIALTF SIVFGVIVYRITTA AALS LNKATRSNVRVTVTATAVI INL  
VVILILDEIYGAVAKWLTKIEVPKTEQTFEERLILKAFLLKFVNAYSPIFYVAFFKGRF  
VGRPGSYVYVFDGYRMEECAPGGCLMELCIQLS IIMLGKQLIQNNIFEIGVPKLKKLFR  
KLKDETEAGETDSASHKHPEQWDL DYSLEPYTGLTPEYMEMI IQFGFVTLFVASFPLAP  
VFALLNNVIEVRLDAKKFVTELRPDAVRTKDIGIWF DILSGIGKFSVISNAFVIAITS  
DFIPRLVYQYSYSHNGTLHG FVNHTLSFFNVS QLKEGTQPENSQFDQEVQFCRFKDYRE  
PPWAPNPYEF SKQYWFILSARLAFV IIFQNLVMFLSVLVDWMI PDIPTDISDQIKKEKS  
LLVDFFLKEEHEK LKLMDEPALRSPGGGDRSR SRAASSAPSGQSQLGSMSSSGSQHTNV

### References

Verkman A.S. and Galiotta L.J.V. (2009) Cholide channels as drug targets. *Nature Reviews* **8**:153-171.

Scudieri P. *et al.* (2012) The anoctamin family: TMEM16A and TMEM16B as calcium-activated chloride channels. *Exp. Physiol.* **97(2)**:177-183.

Stöhr H. *et al.* (2009) TMEM16B, a novel protein with calcium-dependent chloride channel activity, associates with a presynaptic protein complex in photoreceptor terminals. *J. Neurosci.* **29(21)**:6809-6818.

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