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International Union of Pharmacology. LV. Nomenclature and Molecular Relationships of Two-P Potassium Channels

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Introduction

In less than a decade since their discovery, the study of K_{2P} channels has revealed that background leak of potassium ions via dedicated pathways is a highly regulated mechanism to control cellular excitability. Potassium leak pathways, active at rest, stabilize membrane potential below firing threshold and expedite repolarization. Although the existence of leak currents was proposed in 1952 by Hodgkin and Huxley, they remained a biophysical curiosity for more than 4 decades. Identification of the first molecular correlate of a potassium leak current was preceded by cloning of potassium channels in *Saccharomyces cerevisiae* and *Caenorhabditis elegans* with two pore-forming P loops in each subunit and four or eight transmembrane (TM¹) domains (Ketchum et al., 1995). Thereafter, $K_{2P}\emptyset$ was isolated by functional expression cloning from the neuromuscular tissue of *Drosophilia melanogaster* (Goldstein et al., 1996). Biophysical characterization revealed $K_{2P}\emptyset$ to be a potassium-selective channel with the predicted attributes of a background conductance, that is, a voltage-independent portal showing Goldman-Hodgkin-Katz (open) rectification. When the concentration of potassium is symmetrical across the membrane, $K_{2P}\emptyset$ currents change in a linear manner with voltage; under physiological conditions (high internal and low external potassium), $K_{2P}\emptyset$ passes greater outward than inward currents (Goldstein et al., 2001).

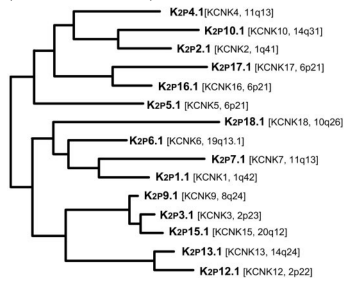


FIG. 1. Phylogenetic tree for K_{2P} channels. Amino acid sequence alignments and phylogenetic analysis for the 15 known members of the human K2P family were generated as described in the legend for Fig. 1 of "LIII. Nomenclature and Molecular Relationships of Voltage-Gated Potassium Channels." $K_{2P}18.1$ was added to the topology shown in the previous edition of this compendium by use of maximum parsimony and neighbor-joining algorithms. International Union of Pharmacology and HUGO Gene Nomenclature Committee names of the genes are shown together with their chromosomal localization.

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¹ Abbreviations: TM, transmembrane

A striking feature of K_{2P} channels is their subunit body plan: each has two P loops and four TM domains. This distinct 2P/4TM topology can be found in more than 70 predicted homologs in genome databases. Fifteen mammalian genes in the family are designated as KCNK genes encoding the K_{2P} channels (Fig. 1); most readily reveal ion channel function upon expression. As expected for regulators of excitability, K_{2P} channels are under tight control by a plethora of chemical and physical stimuli, including oxygen tension, pH, lipids, mechanical stretch, neurotransmitters, and G protein-coupled receptors; the channels are also the molecular targets for certain volatile and local anesthetics (Lesage and Lazdunski, 2000). Regulation of K_{2P} channels alters the attributes subject to change in any ion channel: number of pores at the site of operation, open probability, and unitary current (Plant et al., 2005). Nonetheless, some regulatory changes are striking; for example, phosphorylation of $K_{2P}2$ endows the open rectifier with sensitivity to voltage (Bockenhauer et al., 2001), and desumoylation of $K_{2P}1$ (removal of covalently-bound small ubiquitin-modifier protein) relieves chronic silencing of complexes that reside in the plasma membrane, thereby revealing that the protein can function as an ion channel and operates like $K_{2P}\emptyset$ as an open rectifier (Plant et al., 2005; Rajan et al., 2005). Tables 1 through 15 present the properties of $K_{2P}1.1$ through $K_{2P}18.1$ channels.

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All authors serve as the Subcommittee on K2P Channels of the Nomenclature Committee of the International Union of Pharmacology.

TABLE 1 $K_{2P}1.1$ channels

Channel name $K_{2P}1.1$

Description Two-pore domain potassium channel subunit¹

Other names KCNK1, TWIK-1, hOHO

Molecular information Human: 336aa, NM_002245, chr. 1q42-43, KCNK1,^{2, 3} GeneID: 3775, PMID: 8605869¹

Rat: 336aa, AF022819

Mouse: 336aa, NM_008430, chr. 8, kcnk14

Associated subunits Small ubiquitin-related modifier protein (SUMO-1) is covalently attached at lysine 2746;

exchange factor (EFA6) for small G protein ADP-ribosylation factor 6 (ARF6) (see

"Comments")7

Electrophysiological Functional assays Current Open rectifier

Conductance 32pS

Ion selectivity Not established Activation See "Comments" Inactivation See "Comments" Activators Not established

Gating inhibitors None

Blockers External pH (6.7)⁶

Radioligands

Brain, heart, lung, kidney, liver, placenta^{4,5} Channel distribution

Not established

Physiological functions Not established Mutations and Not established

pathophysiology

Pharmacological significance

Covalent attachment of SUMO to lysine 274 silences $K_{2P}1$; mutation of lysine 274 or Comments

desumoylation of $K_{2P}1$ by a SUMO-specific protease (SENP) reveals an open rectifier; like $K_{2P}3$ and $K_{2P}9$, $K_{2P}1$ is blocked by extracellular acidification due to titration of a histidine residue in the first pore loop; EFA6 interacts with the C-terminal part of $K_{\rm 2P}1$ —this interaction may be

important for channel internalization and recycling⁷

aa, amino acid; chr., chromosome; SUMO, small ubiquitin-related modifier protein.

^{1.} Lesage F, Guillemare E, Fink M, Duprat F, Lazdunski M, Romey G, and Barhanin J (1996) TWIK-1, a ubiquitous human weakly inward rectifying K+ channel with a novel structure. EMBO J 15:1004-1011.

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TABLE 2 $K_{op}2.1$ channels

Channel name K_{2p}2.1

Description Two-pore domain potassium channel subunit¹; open rectifier or voltage-dependent

Other names KCNK2, TREK-1, TPKC1

Molecular information Human: 426aa, NM_014217, chr. 1q41, KCNK2, GeneID: 3776, PMID: 90037613

Rat: 426aa, AF325671, chr. 5

Mouse: 411aa, XM_123605, chr. 1, kcnk23

Associated subunits Not established Functional assays Electrophysiological

Current Open or voltage-dependent^{4,5} (see "Comments")

Conductance 90pS (see "Comments")
Ion selectivity Not established
Activation See "Comments"
Inactivation See "Comments"

Activators Arachidonic acid (10 mM) and unsaturated fatty acids, 10 lysophospholipids, 7 volatile

anesthetics, 6,11 mechanical stress, 7,11 internal acidification 12

Gating inhibitors None

Blockers Ba²⁺ (1 mM), quinidine (100 mM), PKA, PKC

Radioligands None Channel distribution Brain,² heart

Channel distribution Brain, heart
Physiological functions Not established

Mutations and Characterization of $K_{2p}2$ knockout mice suggests a loss of sensitivity to general pathophysiology anesthetics and increased vulnerability to ischemia and reperfusion injury^{8,9}

Pharmacological significance Not establishe

Comments Phosphorylation of serine 348 regulates reversible interconversion between leak and voltage-dependent phenotypes⁵; "activation" and "deactivation" with voltage steps

voltage-dependent phenotypes⁵; "activation" and "deactivation" with voltage step seem to be instantaneous; the mouse variant may have a smaller conductance

aa, amino acids; chr., chromosome; PKA, protein kinase A; PKC, protein kinase C.

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Channel name $K_{2P}3.1$

Description Two-pore domain potassium channel subunit; open rectifier

Other names KCNK3, TASK-1, TBAK-1, OAT-1

Human: 394aa, NM_002246, chr. 2p24.1-23.3, KCNK3, GeneID: 3777, PMID: 931200520 Molecular information

Rat: 411aa, NP_203694, kcnk33

Mouse: 409aa, AF065162, chr. 5B,1 kcnk3

14-3-316, 17 and p11 (annexin II subunit), 18 see "Comments" Associated subunits

Electrophysiological Functional assays Current Open rectifier4

Conductance $10pS^5$

 $Rb^{+} > K^{+} > Cs^{+} > NH_{4}^{+} \gg Na^{+} > Li^{+}$ Ion selectivity

See "Comments" Activation Inactivation See "Comments"

Volatile anesthetics^{6,7}: halothane (1 mM),⁵ isofluorane (2 mM) Activators

Gating inhibitors

Blockers Ba²⁺ (500 mM), external pH (7.3),⁸⁻¹⁰ arachidonic acid (100 mM) (see "Comments"), and

anandamide $(3 \mu M)^{19}$

Radioligands

Brain, 11 heart, 12 lung, kidney, 13 small intestine, colon, pancreas, prostate, uterus, Channel distribution

placenta

Physiological functions Not established Not established Mutations and

pathophysiology

Pharmacological significance

Comments

Not established

Activation and deactivation with voltage steps seems to be instantaneous, but there is also a small, time-dependent change in Po; current is half-blocked at pH 7.3 at physiological external conditions—increasing external potassium decreases proton blockade; pharmacology studies of the rat variant reveal blockade also by zinc, TEA, and quinidine 14,15 ; K_{2P} 3-like currents are reported in cerebellar granular neurons and motor-neurons^{11,15}; interaction with 14-3-3 protein is essential for forward trafficking; K_{2p} 3 can form heterodimers with K_{2p} 9.1 in heterologous expression systems consistent with electrophysiological studies that suggest heterodimerzation; K₀P3 is also suggested to be a target for transmitter modulation of neuronal excitability 11,15

aa, amino acids; chr., chromosome; TEA, tetrylethylammonium.

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Channel name

Description Two-pore domain potassium channel subunit; open rectifier

Other names KCNK4, TRAAK

Human: 393aa, NM 016611, chr. 11q13, KCNK4, GeneID: 50801, PMID: 107674091 Molecular information

Rat: 397aa, NM_053804, kcnk4

Mouse: 398aa, NM_008431, chr. 19, kcnk4

Not established Associated subunits Functional assays Electrophysiological² Current Open rectifier

Conductance 46pS

Ion selectivity Not established Activation See "Comments" See "Comments" Inactivation

Arachidonic acid (10 mM),3 mechanical stress,4 heat6 (see "Comments"), unsaturated Activators

fatty acids,3 lysopholipids,7 riluzole8

Gating inhibitors None Blockers Gd^+ Radioligands None

Brain,⁵ kidney, small intestine, placenta, prostate Channel distribution

Physiological functions Not established Mutations and Not established

pathophysiology Pharmacological significance Not established

Comments Activation and deactivation with voltage steps seem to be instantaneous; knockout

mice have no obvious phenotype9; the open probability of K2P4 increases with

temperature with an activation threshold of 31°C in COS-7 cells

aa, amino acids; chr., chromosome

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TABLE 5 $K_{op}5.1$ channels

Channel name $K_{2P}5.1$

Description Two-pore domain potassium channel subunit; open rectifier

Other names KCNK5, TASK-2

Molecular information Human: 499aa, NM_003740, chr. 6p21, KCNK5, GeneID: 8645, PMID: 98129781

Rat: not cloned

Mouse: 502aa, NM_021542, kcnk5

Associated subunits

Functional assays

Current

Conductance

Ion selectivity

Activation

Activation

Activation

Activation

Not established

See "Comments"

See "Comments"

See "Comments"

Activators Volatile anaesthetics²: halothane (~570 mM)

Gating inhibitors None

Blockers Quinidine (22 mM), external pH (6.5), local anesthetics: lidocaine (1 mM), bupivacaine (1

mM), clofilium $(25 \mu M)^7$

Radioligands None

Channel distribution Brain, kidney, liver, small intestine, pancreas, placenta

Physiological functions A role in cell volume regulation^{7,8} (see "Comments") and sensing external basolateral pH

changes associated with HCO3 transport in primary-cultured proximal tubular cells4

Not established

pathophysiology

Pharmacological significance

Comments

Mutations and

Not established

Activation and deactivation with voltage steps seem instantaneous; the conductance of $\rm K_{2P}5$ depends on the ionic conditions; the slope conductance was reported as 15pS with 5 mM external potassium and as high as 60pS when external potassium is high (155 mM)¹— this may reflect an Na⁺-dependent inward rectification that becomes progressively less pronounced with time⁵; like $\rm K_{2P}16$ and 17, current through $\rm K_{2P}5$ channels is diminished at physiological pH; channel open probability increases with external pH; formation of an intersubunit disulfide bridge in $\rm K_{2P}5$ does not affect channel activity⁹; exposure to hypotonicity (change from 300–200 mOsm in external solution) enhanced m $\rm K_{2P}5$ currents when this channel was heterologously expressed in HEK293 cells, and osmotic cell shrinkage led to inhibition (change from 300–400 mOsm in external solution)

aa, amino acids; chr., chromosome.

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Channel name $K_{2p}6.1$

Description Two-pore domain potassium channel subunit; open rectifier

Other names KCNK6, TWIK-2, TOSS

Human: 313aa, 1 NM_004823, chr0.19q13-1, 2 KCNK6, GeneID: 9424, PMID: 103590731 Molecular information

Rat: 313aa, NM 053806, kcnk6

Mouse: not cloned Associated subunits Not established Electrophysiological Functional assays Current Open rectifier^{3,4}

Conductance <5pS

Ion selectivity Not established Activation See "Comments" Inactivation See "Comments" Activators Arachidonic acid

Gating inhibitors None

 Ba^{2+} (100 μM), quinidine (100 mM), volatile anesthetics Blockers

Radioligands None

Channel distribution Pancreas, placenta, heart (see "Comments")

Physiological functions Not established Mutations and Not established

pathophysiology

Pharmacological significance Not established

Comments Activation and deactivation with voltage steps seem to be instantaneous; displays

time-dependent inactivation at depolarized potentials4; the rat variant has been reported to be widely expressed (including brain, lung, kidney, liver, spleen, heart,

esophagus, stomach, colon, and skeletal muscle)

2. Gray AT, Kindler CH, Sampson ER, and Yost CS (1999) Assignment of KCNK6 encoding the human weak inward rectifier potassium channel TWIK-2 to chromosome band 19q13.1 by radiation hybrid mapping. Cytogenet Cell Genet 84:190-191.

TABLE 7 $K_{op}7.1$ channels

Channel	name	$K_{op}7.1$
Chamie	name	TYODILL

Description Two-pore domain potassium channel subunit

Other names KCNK7, kcnk8 (see "Comments")

Human: 307aa (see "Comments"), NM_033347, chr0.11q13, KCNK7, GeneID:10089, PMID: Molecular information

10206991¹ Rat: not cloned

Mouse: 335aa, NM_010609, chr. 19,2B, kcnk8 (see "Comments")1,2

Associated subunits Not established Functional assays Electrophysiological

Current Not established (see "Comments")

Not established Conductance Not established Ion selectivity Not established Activation Not established Inactivation

Activators None Gating inhibitors None Blockers None Radioligands None

Brain (human), retina (mouse) Channel distribution

Not established Physiological functions Mutations and Not established

pathophysiology

Pharmacological significance Not established

Comments The product of this gene has not vet been shown to form a functional channel; five splice

variants have been identified in human; the mouse isolate was cited as kcnk6 and then kcnk8 but is now called K_{2P} 7 due to its homology and syntenic location to human KCNK7 2

aa, amino acids; chr., chromosome; TOSS, TWIK-originated similarity sequence.

^{1.} Pountney DJ, Gulkarov I, Vega-Saenz de Miera E, Holmes D, Saganich M, Rudy B, Artman M, and Coetzee WA (1999) Identification and cloning of TWIK-originated similarity sequence (TOSS): a novel human 2-pore K+ channel principal subunit. FEBS Lett 450:191-196.

^{3.} Chavez RA, Gray AT, Zhao BB, Kindler CH, Mazurek MJ, Mehta Y, Forsayeth JR, and Yost CS (1999) TWIK-2, a new weak inward rectifying member of the tandem pore domain potassium channel family. J Biol Chem 274:7887–7892.

4. Patel AJ, Maingret F, Magnone V, Fosset M, Lazdunski M, and Honoré E (2000) TWIK-2, an inactivating 2P domain K⁺ channel. J Biol Chem 275:28722–28730.

aa, amino acids; chr., chromosome

^{1.} Salinas M, Reyes R, Lesage F, Fosset M, Heurteaux C, Romey G, and Lazdunski M (1999) Cloning of a new mouse two-P domain channel subunit and a human homologue with a unique pore structure. J Biol Chem 274:11751-11760.

^{2.} Bockenhauer D, Nimmakayalu MA, Ward DC, Goldstein SAN, and Gallagher PG (2000) Genomic organization and chromosomal localization of the murine 2 P domain potassium channel gene Kcnk8: conservation of gene structure in 2P domain potassium channels. Gene 261:365-372.

TABLE 8 K_{2D} 9.1 channels

Channel name $K_{2P}9.1$

Description Two-pore domain potassium channel subunit; open rectifier

Other names KCNK9, TASK-3

Molecular information Human: 374aa, NM_016601, chr0.8q24-3, KCNK9, 1-3 GeneID: 51305, PMID:10734076

Rat: 395aa, NM_053405, kcnk9

Mouse: not cloned

Associated subunits 14-3-3 (see "Comments") 7,8 Functional assays Electrophysiological $^{1-3,\delta-11}$

Current Not established (see "Comments")

Conductance 27pS (see "Comments")
Ion selectivity Not established
Activation See "Comments"
Inactivation See "Comments"

Activators None Gating inhibitors None

Blockers External pH (6.5), ruthenium red (700 nM)⁸

Radioligands None

Channel distribution Brain (see "Comments")¹
Physiological functions See "Comments"⁴⁻⁶
Mutations and pathophysiology
Pharmacological significance Not established

Comments Activation and deactivation with voltage steps seem to be instantaneous; the guinea pig variant is

reported to have the same conductance and distribution as human and a conductance of 60pS; Northern blot analysis suggests that rat $K_{\rm 2P}9.1$ expression outside the CNS is extremely low, as is noted for the human and guinea pig gene; $K_{\rm 2P}9$ gene is amplified in several human carcinomas, and overexpression of $K_{\rm 2P}9$ protein in cell lines promotes tumor formation^{4,5}; like $K_{\rm 2P}3$, surface expression of $K_{\rm 2P}9$ depends on its association with 14-3-3 to release it from the endoplasmic

reticulum^{7,8}; potential heterodimerization of $K_{2P}9$ is discussed under $K_{2P}3^9$

aa, amino acids; chr., chromosome; CNS, central nervous system.

1. Chapman CG, Meadows HJ, Godden RJ, Campbell DA, Duckworth M, Kelsell RE, Murdock PR, Randall AD, Rennie GI, and Gloger IS (2000) Cloning, localisation and functional expression of a novel human, cerebellum specific, two pore domain potassium channel. *Mol Brain Res* 82:74–83.

2. Kim Y, Bang H, and Kim D (2000) TASK-3, a new member of the tandem pore K(+) channel family. J Biol Chem 275:9340-9347.

3. Rajan S, Wischmeyer E, Xin Liu G, Preisig-Muller R, Daut J, Karschin A, and Derst C (2000) TASK-3, a novel tandem pore domain acid-sensitive K⁺ channel—an extracellular histidine as pH sensor. J Biol Chem 275:16650–16657.

4. Mu D, Chen L, Zhang X, See LH, Koch CM, Yen C, Tong JJ, Spiegel L, Nguyen KC, Servoss A, et al. (2003) Genomic amplification and oncogenic properties of the KCNK9 potassium channel gene. Cancer Cell 3:297–302.

5. Pei L, Wiser O, Slavin A, Mu D, Powers S, Jan LY, and Hoey T (2003) Oncogenic potential of TASK3 (Kcnk9) depends on K+ channel function. *Proc Natl Acad Sci USA* 100:7803–7807.

6. Lauritzen I, Zanzouri M, Honoré E, Duprat F, Ehrengruber MU, Lazdunski M, and Patel AJ (2003) K⁺-dependent cerebellar granule neuron apoptosis: role of TASK leak K+ channels. J Biol Chem 278:32068–32076.

7. Rajan S, Preisig-Muller R, Wischmeyer E, Nehring R, Hanley PJ, Renigunta V, Musset B, Schlichthorl G, Derst C, Karschin A, et al. (2002) Interaction with 14-3-3 proteins promotes functional expression of the potassium channels TASK-1 and TASK-3. J Physiol 545:13-26.

8. O'Kelly I, Butler MH, Zilberberg N, and Goldstein SA (2002) Forward transport. 14-3-3binding overcomes retention in endoplasmic reticulum by dibasic signals. Cell 111:577-588.

9. Kang DW, Han JH, Talley EM, Bayliss DA, and Kim D (2004) Functional expression of TASK-1/TASK-3 heteromer in cerebellar granule neurons. *J Physiol* 554:64–77. 10. Czirjak G and Enyedi P (2003) Ruthenium red inhibits TASK-3 potassium channel by interconnecting glutamate 70 of the two subunits. *Mol Pharmacol* 63:646–652: 11. Vega-Saenz de Miera E, Lau DH, Zhadina M, Pountney D, Coetzee WA, and Rudy B (2001) KT3.2 and KT3.3, two novel human two-pore K(+) channels closely related to TASK-1. *J Neurophysiol* 86:130–142.

TABLE 9 $K_{op}10.1$ channels

 $K_{2P}10.1$ Channel name

Description Two-pore domain potassium channel subunit; open rectifier¹

Other names KCNK10, TREK-2

Molecular information Human: 538aa (see 'Comments'), NM_138317, chr0.14q31, KCNK10, GeneID: 54207, PMID:

10747911¹

Rat: 538aa, NM_023096, kcnk10

Mouse: not cloned Not established Electrophysiological Open rectifier²

Conductance 100pS (see "Comments")

Ion selectivity Not established Activation See "Comments" Inactivation See "Comments"

Arachidonic acid, docosahexaenoic acid, linoleic acid, lysophosphatidylcholine,3 intracellular Activators

acidification, volatile anesthetics: halothane (~1 mM), isoflurane (~1 mM); riluzole (~1 mM),

heat.6 mechanical stress3

Gating inhibitors None

Quinidine (100 mM), PKA, PKC Blockers

Radioligands None

Kidney, pancreas, prostate, thymus, liver, heart (see "Comments")2 Channel distribution

See "Comments"^{4, 5} Physiological functions Not established Mutations and

pathophysiology

Associated subunits

Functional assays

Current

Pharmacological significance Not established

Comments Activation and deactivation with voltage steps seem to be instantaneous; splice variants have

> been identified in human and rat; the rat variant is reported to have a conductance of 68pS and to be expressed in brain; K_{2P}10-like currents are observed in cerebellar granular neurons, magnocellular neurosecretory cells of rat supraoptic nucleus, 4,5 rat cortical astrocytes, 7 and

insulin-secreting MIN6 cells⁸

aa, amino acids; chr., chromosome; PKA, protein kinase A; protein kinase C.

1. Bang H, Kim Y, and Kim D (2000) TREK-2, a new member of the mechano-sensitive tandem-pore K⁺ channel family. J Biol Chem 275:17412–17419.

2. Gu W, Schlichthorl G, Hirsch JR, Engels H, Karschin C, Karschin A, Derst C, Steinlein OK, and Daut J (2002) Expression pattern and functional characteristics of two novel splice variants of the two-pore-domain potassium channel TREK-2. J Physiol 539:657-668.

3. Lesage F, Terrenoire C, Romey G, and Lazdunski M (2000) Human TREK2, a 2P domain mechano-sensitive K⁺ channel with multiple regulations by polyunsaturated fatty acids, lysophospholipids, and Gs, Gi, and Gq protein-coupled receptors. J Biol Chem 275:28398–28405.

4. Han J, Truell J, Gnatenco C, and Kim D (2002) Characterization of four types of background potassium channels in rat cerebellar granule neurons. J Physiol 542:431-444.

5. Han J, Gnatenco C, Sladek CD, and Kim D (2003) Background and tandem-pore potassium channels in magnocellular neurosecretory cells of the rat supraoptic nucleus. J Physiol 546:625-639.

Kang DW, Choe CY, and Kim D (2005) Thermosensitivity of the two-pore domain K⁺ channels TREK-2 and TRAAK. J Physiol 564:103-116.

7. Gnatenco C, Han JH, Snyder AK, and Kim D (2002) Functional expression of TREK-2 K+ channel in cultured astrocytes. Brain Res 931:56-67

8. Kang DW, Choe C, and Kim D (2004) Functional expression of TREK-2 in insulin-secreting MIN6 cells. Biochem Biophys Res Commun 323:323-331.

Channel name K_{2P}12.1

Description Two-pore domain potassium channel subunit¹

Other names KCNK12, THIK-2

Molecular information Human: 430aa, NM_022055, chr0.2p22-p21, KCNK12, GeneID: 56660, PMID: 11060316¹

Rat: 430aa, NM_022292, kcnk12

Associated subunits Not established
Functional assays Electrophysiological

Current Not established (see "Comments")^{1,2}

Conductance No function demonstrated

Ion selectivityNot establishedActivationNot establishedInactivationNot established

Activators None
Gating inhibitors None
Blockers None
Radioligands None

Channel distribution Brain, heart, lung, kidney, liver, small intestine, colon, pancreas, prostate, placenta,

spleen, thymus, ovary

Physiological functions Not established Mutations and Not established

pathophysiology

Pharmacological significance Not established

Comments The product of this gene has not yet been shown to be a functional channel

aa, amino acids; chr., chromosome.

TABLE 11 $K_{op}13.1$ channels

Channel name	$K_{2P}13.1$
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Description Two-pore domain potassium channel subunit; open rectifier¹

Other names KCNK13, THIK-1

Molecular information Human: 408aa, NM_022054, chr0.14q24.1-24.3, KCNK13, GeneID: 56659,

PMID: 110603161

Rat: 405aa, NM_022293, kcnk13

Associated subunits

Functional assays

Current

Conductance

Ion selectivity

Activation

Associated subunits

Mouse: not cloned

Not established

Plectrophysiological

Open rectifier

Not established

Not established

See "Comments"

See "Comments"

Activators Arachidonic acid (0.98 mM)

Gating inhibitors None

Blockers Ba²⁺, halothane (2.83 mM)

Radioligands None

Channel distribution Brain, heart, lung, kidney, liver, spleen

Physiological functions Not established Mutations and Not established

pathophysiology

Pharmacological significance Not established

Comments Activation and deactivation with voltage steps seem to be instantaneous; $K_{2P}13$ like channels are found in the central nervous system of $Aplysia\ californica^2$

^{1.} Rajan S, Wischmeyer E, Karschin C, Preisig-Muller R, Grzeschik KH, Daut J, Karschin A, and Derst C (2001) THIK-1 and THIK-2, a novel subfamily of tandem pore domain K⁺ channels. J Biol Chem 276:7302–7311.

^{2.} Girard C, Duprat F, Terrenoire C, Tinel N, Fosset M, Romey G, Lazdunski M, and Lesage F (2001) Genomic and functional characteristics of novel human pancreatic 2P domain K⁺ channels. *Biochem Biophys Res Commun* 282:249–256.

aa, amino acids; chr., chromosome.

Rajan S, Wischmeyer E, Karschin C, Preisig-Muller R, Grzeschik KH, Daut J, Karschin A, and Derst C (2001) THIK-1 and THIK-2, a novel subfamily of tandem pore domain K⁺ channels. J Biol Chem 276:7302-7311.

Jezzini SH and Moroz LL (2004) Identification and distribution of a two-pore domain potassium channel in the CNS of Aplysia californica. Brain Res Mol Brain Res 27:7–38.

Channel name K_{2P}15.1

Description Two-pore domain potassium channel subunit

Other names KCNK15, TASK-5, 1,2 KT3.33

Molecular information Human: 330aa, NM_022358, chr0.20q12, KCNK15, GeneID: 60598, PMID: 11409881

Rat: 318aa, AF467250 Mouse: 324aa, XM_141526

Associated subunits Not established Functional assays Electrophysiological

Current Not established (see "Comments")

Conductance Not established Ion selectivity Not established Activation Not established Inactivation Not established

Activators None
Gating inhibitors None
Blockers None
Radioligands None

Channel distribution Brain, heart, lung, kidney, liver, pancreas, adrenal gland, thyroid, salivary gland, placenta

Physiological functions
Mutations and
pathophysiology
Pharmacological significance
Not established
Not established

Comments The product of this gene has not yet been shown to be a functional channel

aa, amino acids; chr., chromosome.

1. Kim D and Gnatenco C (2001) TASK-5, a new member of the tandem-pore K(+) channel family. Biochem Biophys Res Commun 284:923-930.

Ashmole I, Goodwin PA, and Stanfield PR (2001) TASK-5, a novel member of the tandem pore K⁺ channel family. Pflueg Arch Eur J Physiol 442:828-833.

3. Vega-Saenz de Miera E, Lau DH, Zhadina M, Pountney D, Coetzee WA, and Rudy B (2001) KT3.2 and KT3.3, two novel human two-pore K(+) channels closely related to TASK-1. J Neurophysiol 86:130–142.

TABLE 13 $K_{op}16.1$ channels

Channel name $K_{2P}16.1$

Description Two-pore domain potassium channel subunit; open rectifier

Other names KCNK16, TALK-1

Molecular information Human: 309aa, NM_032115, chr0.6p21.1-2, KCNK16, GeneID: 83795, PMID: 11263999¹

Rat: not cloned

Mouse: 337aa, XM_138942

Associated subunits Not established
Functional assays Electrophysiological¹
Current Open rectifier

Conductance $21pS \text{ at } -60 \text{ mV} \text{ and } 10 \text{ pS at } +60 \text{ mV}^2$

Ion selectivityNot establishedActivationSee "Comments"InactivationSee "Comments"

Activators Isoflurane (~800 mM), nitric oxide and reactive oxygen species³

Gating inhibitors None

Blockers Ba²⁺ (1 mM), quinidine (100 mM), chloroform (~800 mM), external pH (see "Comments")

Radioligands None

Channel distribution Heart, lung, liver, pancreas, and placenta

Physiological functions Not established Mutations and Not established

pathophysiology

Pharmacological significance Not established

Comments Activation and deactivation with voltage steps seem to be instantaneous; the open

probability of both $K_{\rm 2P}16$ and 17 increases with external pH—at present, it is unclear whether this represents proton block at physiological levels or activation of the channel by supraphysiological alkaline pHo; there are four splice variants of $K_{\rm 2P}16$, two of

which are functional4

aa, amino acids; chr., chromosome.

^{1.} Girard C, Duprat F, Terrenoire C, Tinel N, Fosset M, Romey G, Lazdunski M, and Lesage F (2001) Genomic and functional characteristics of novel human pancreatic 2P domain K(+) channels. Biochem Biophys Res Commun 282:249–256.

Kang D and Kim D (2004) Single-channel properties and pH sensitivity of two-pore domain K⁺ channels of the TALK family. Biochem Biophys Res Commun 315:836-844.

^{3.} Duprat F, Girard C, Jarretou G, and Lazdunski M (2004) Pancreatic two P domain K⁺ channels TALK-1 and TALK-2 are activated by nitric oxide and reactive oxygen species. J Physiol 562:235–244.

^{4.} Han JH, Kang D, and Kim D (2003) Functional properties of four splice variants of a human pancreatic tandem-pore K⁺ channel, TALK-1. Am J Physiol 285:C529-C538.

TABLE 14 K_{2p} 17.1 channels

Channel name $K_{2P}17.1$

Description Two-pore domain potassium channel subunit; open rectifier

Other names KCNK17, TASK-4, TALK-2

Human: 332aa, NM_031460, chr0.6p21.1-2, KCNK17, GeneID: 89822, PMID: 112639991 Molecular information

> Rat: not cloned Mouse: not cloned

Not established Associated subunits Electrophysiological1 Functional assays

Open rectifier Current Not established Conductance Ion selectivity Not established Activation See "Comments" Inactivation See "Comments"

Nitric oxide and reactive oxygen species³ Activators

Gating inhibitors None

 Ba^{2+} , external pH,^{2,3} chloroform (~800 mM) Blockers

Radioligands None

Channel distribution Heart, lung, liver, pancreas,1 placenta

Physiological functions Not established Mutations and Not established

pathophysiology

Pharmacological significance Not established

Comments Activation and deactivation with voltage steps seem to be instantaneous; the open probability of $K_{\rm 2P}17$ increases as pHo is raised above physiological levels (see $K_{\rm 2P}16$)

aa, amino acids; chr., chromosome.
1. Girard C, Duprat F, Terrenoire C, Tinel N, Fosset M, Romey G, Lazdunski M, and Lesage F (2001) Genomic and functional characteristics of novel human pancreatic 2P domain K(+) channels. Biochem Biophys Res Commun 282:249–256.
2. Decher N, Maier M, Dittrich W, Gassenhuber J, Bruggemann A, Busch AE, and Steinmeyer K (2001) Characterization of TASK-4, a novel member of the pH-sensitive, two-pore domain potassium channel family. FEBS Lett 492:84–89.

^{3.} Duprat F, Girard C, Jarretou G, and Lazdunski M (2004) Pancreatic two P domain K+ channels TALK-1 and TALK-2 are activated by nitric oxide and reactive oxygen species. J Physiol 562:235-244.

TABLE 15 $K_{op}18.1$ channels

Channel name $K_{2P}18.1$

Description Two-pore domain potassium channel subunit; open rectifier

Other names KCNK18, TRESK-1/TRESK-2 (see "Comments")

Molecular information Human: 384aa, NM_181840, chr. 10q26.11, GeneID: 338567, PMID: 12754259¹

Rat: 405aa, NM_001003820 Mouse: 394aa, NM_207261

Associated subunits Not established
Functional assays Electrophysiological¹
Current Open rectifier

Conductance 13pS at +60 mV and 16pS at -60 mV for mouse $K_{2p}18^3$

Ion selectivity Not established

Activation Rapid Inactivation Slow

Activators Cytoplasmic Ca²⁺ via calcineurin, volatile anesthetics^{2,4}

Gating inhibitors None

Blockers Ba^{2+} (3 mM), quinine (100 mM) quinidine (100 mM), free fatty acids, external acidic pH

Radioligands None

Channel distribution Cerebrum, cerebellum, brain stem, spinal cord, and testis

Physiological functions Not established Mutations and Not established

pathophysiology

Pharmacological significance Not established

Comments Activation is instantaneous; single channel currents are noninactivating and

time-dependent; TRESK2 was cloned from mouse testis and shares 65% homology with human $\rm K_{2P}18$; as study continues, it will become clear whether this is the true correlate of the human channel or a distinct gene ($\rm K_{2P}19$); personal communication indicates that distinct cDNAs for both TRESK-1 and TRESK-2 are present in human

tissues (D. Kim, personal communication)

aa, amino acids; chr., chromosome.

^{1.} Sano Y, Inamura K, Miyake A, Mochizuki S, Kitada C, Yokoi H, Nozawa K, Okada H, Matsushime H, and Furuichi K (2003) A novel two-pore domain K⁺ channel, TRESK, is localized in the spinal cord. *J Biol Chem* 278:27406–27412.

^{2.} Czirjak G, Toth ZE, and Enyedi P (2004) The two-pore-domain K⁺ channel, TRESK, is activated by the cytoplasmic calcium signal through calcineurin. J Biol Chem 279:18550–18558.

^{3.} Kang D, Mariash E, and Kim D (2004) Functional expression of TRESK-2, a new member of the tandem-pore K + channel family. *J Biol Chem* 279:28063–28070.
4. Liu C, Au JD, Zou HL, Cotton JF, and Yost CS (2004) Potent activation of the human tandem pore domain K channel TRESK with clinical concentrations of volatile anesthetics. *Anesth Analg* 99:1715–1722.