

KCNQ1 抗原(重组蛋白)

中文名称: KCNQ1 抗原 (重组蛋白)

英文名称: KCNQ1 Antigen (Recombinant Protein)

别 名: potassium voltage-gated channel, KQT-like subfamily, member 1; LQT; RWS; WRS; LQT1; SQT2; ATFB1; ATFB3; JLNS1; KCNA8; KCNA9; Kv1.9;K

储存: 冷冻(-20℃)

相关类别: 抗原

概述

Fusion protein corresponding to C terminal 250 amino acids of human KCNQ1

技术规格

| Full name: | potassium voltage-gated channel, KQT-like subfamily, member 1 |
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| Synonyms: | LQT; RWS; WRS; LQT1; SQT2; ATFB1; ATFB3; JLNS1; KCNA8; KCNA9 ; Kv1.9; Kv7.1; KVLQT1 |
| Swissprot: | P51787 |
| Gene Accession: | NP_000209 |
| Purity: | >85%, as determined by Coomassie blue stained SDS-PAGE |
| Expression system: | Escherichia coli |
| Tags: | His tag C-Terminus, GST tag N-Terminus |
| Background: | This gene encodes a voltage-gated potassium channel required for repolarization phase of the cardiac action potential. This protein ca n form heteromultimers with two other potassium channel proteins , KCNE1 and KCNE3. Mutations in this gene are associated with he reditary long QT syndrome 1 (also known as Romano-Ward syndro |



me), Jervell and Lange-Nielsen syndrome, and familial atrial fibrillat ion. This gene exhibits tissue-specific imprinting, with preferential e xpression from the maternal allele in some tissues, and biallelic ex pression in others. This gene is located in a region of chromosom e 11 amongst other imprinted genes that are associated with Beck with-Wiedemann syndrome (BWS), and itself has been shown to b e disrupted by chromosomal rearrangements in patients with BWS. Alternatively spliced transcript variants have been found for this ge ne.