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# International Union of Pharmacology. XLVIII. Nomenclature and Structure-Function Relationships of Voltage-Gated Calcium Channels

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Abstract—The family of voltage-gated calcium channels serves as the key transducers of cell surface membrane potential changes into local intracellular calcium transients that initiate many different physiological events. There are 10 members of the voltage-gated calcium channel family that have been characterized in

mammals, and they serve distinct roles in cellular signal transduction. This article presents the molecular relationships and physiological functions of these calcium channel proteins and provides comprehensive information on their molecular, genetic, physiological, and pharmacological properties.

# Introduction

Voltage-gated calcium channels mediate calcium influx in response to membrane depolarization and regulate intracellular processes such as contraction, secretion, neurotransmission, and gene expression in many different cell types. Their activity is essential to couple electrical signals in the cell surface to physiological events in cells. They are members of a gene superfamily of transmembrane ion channel proteins that includes voltage-gated potassium and sodium channels (Yu and Catterall, 2004). This compendium presents an introduction to their biochemical, molecular, and genetic properties, their physiological roles, and their pharmacological significance. Table 1 and the summary tables that follow the text of this article give comprehensive information on each member of the calcium channel family.

# **Calcium Channel Subunits**

The calcium channels that have been characterized biochemically are complex proteins composed of four or five distinct subunits that are encoded by multiple genes (Fig. 1; Catterall, 2000). The  $\alpha_1$  subunit of 190 to 250 kDa is the largest subunit, and it incorporates the conduction pore, the voltage sensor and gating apparatus, and most of the known sites of channel regulation by second messengers, drugs, and toxins. Like the  $\alpha$  sub-

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The authors serve as the Subcommittee on Calcium Channels of the Nomenclature Committee of the International Union of Pharmacology. Article, publication date, and citation information can be found at

http://pharmrev.aspetjournals.org. doi:10.1124/pr.57.4.5. units of sodium channels, the  $\alpha_1$  subunit of voltagegated calcium channels is organized in four homologous domains (I–IV), with six transmembrane segments (S1– S6) in each. The S4 segment serves as the voltage sensor. The pore loop between transmembrane segments S5 and S6 in each domain determines ion conductance and selectivity, and changes of only three amino acids in the pore loops in domains I, III, and IV will convert a sodium channel to calcium selectivity. An intracellular  $\beta$  subunit and a transmembrane, disulfide-linked  $\alpha_2\delta$  subunit complex are components of most types of calcium channels. A γ subunit has also been found in skeletal muscle calcium channels, and related subunits are expressed in heart and brain. Although these auxiliary subunits modulate the properties of the channel complex, the pharmacological and electrophysiological diversity of calcium channels arises primarily from the existence of multiple  $\alpha_1$  subunits (Hofmann et al., 1994).

# **Calcium Currents**

Calcium currents recorded in different cell types have diverse physiological and pharmacological properties, and an alphabetical nomenclature has evolved for the distinct classes of calcium currents (Tsien et al., 1995). L-type calcium currents typically require a strong depolarization for activation, are long-lasting, and are blocked by the organic L-type calcium channel antagonists, including dihydropyridines, phenylalkylamines, and benzothiazepines. They are the main calcium currents recorded in muscle and endocrine cells, where they initiate contraction and secretion. L-type currents activating at lower voltages also exist predominantly in neurons and cardiac pacemaker cells. N-type, P/Q-type, and R-type calcium currents

Ca<sub>v</sub>3.1

 $Ca_{V}3.2$ 

Cav3.3

Channel	Current	Localization	Specific Antagonists	Cellular Functions
$\mathrm{Ca_{V}}1.1$	L	Skeletal muscle; transverse tubules	Dihydropyridines; phenylalkylamines; benzothiazepines	Excitation-contraction coupling
$\mathrm{Ca_{V}1.2}$	L	Cardiac myocytes; smooth muscle myocytes; endocrine cells; neuronal cell bodies; proximal dendrites	Dihydropyridines; phenylalkylamines; benzothiazepines	Excitation-contraction coupling; hormone release; regulation of transcription; synaptic integration
$\mathrm{Ca_{v}}1.3$	L	Endocrine cells; neuronal cell bodies and dendrites; cardiac atrial myocytes and pacemaker cells; cochlear hair cells	Dihydropyridines; phenylalkylamines; benzothiazepines	Hormone release; regulation of transcription; synaptic regulation; cardiac pacemaking; hearing; neurotransmitter release from sensory cells
$\mathrm{Ca_{V}}1.4$	L	Retinal rod and bipolar cells; spinal cord; adrenal gland; mast cells	Dihydropyridines; phenylalkylamines; benzothiazepines	Neurotransmitter release from photoreceptors
$\mathrm{Ca_{V}}2.1$	P/Q	Nerve terminals and dendrites; neuroendocrine cells	$\omega$ -Agatoxin IVA	Neurotransmitter release; dendritic Ca <sup>2+</sup> transients; hormone release
$\mathrm{Ca_{V}}2.2$	N	Nerve terminals and dendrites; neuroendocrine cells	$\omega$ -Conotoxin-GVIA	Neurotransmitter release; dendritic Ca <sup>2+</sup> transients; hormone release
$\mathrm{Ca_{V}2.3}$	R	Neuronal cell bodies and dendrites	SNX-482	Repetitive firing; dendritic calcium

None

None

None

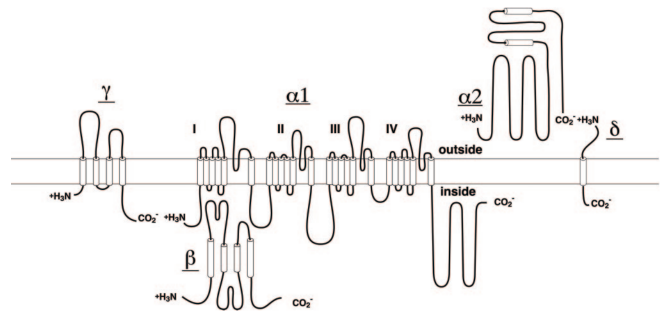
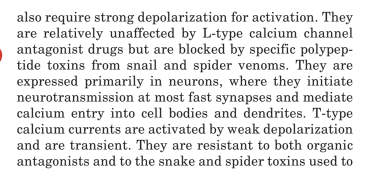


Fig. 1. Subunit structure of Ca<sub>V</sub>1 channels. The subunit composition and structure of calcium channels purified from skeletal muscle are illustrated. The model is updated from the original description of the subunit structure of skeletal muscle calcium channels. This model fits available biochemical and molecular biological results for other  $Ca_V1$  channels and for  $Ca_V2$  channels. Predicted  $\alpha$  helices are depicted as cylinders. The lengths of lines correspond approximately to the lengths of the polypeptide segments represented.



Neuronal cell bodies and dendrites;

Neuronal cell bodies and dendrites;

Neuronal cell bodies and dendrites

cardiac and smooth muscle myocytes

cardiac and smooth muscle myocytes

define the N- and P/Q-type calcium currents. They are expressed in a wide variety of cell types, where they are involved in shaping the action potential and controlling patterns of repetitive firing.

transients

Pacemaking; repetitive firing

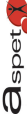
Pacemaking; repetitive firing

Pacemaking; repetitive firing

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# **Calcium Channel Genes**

Mammalian  $\alpha_1$  subunits are encoded by at least 10 distinct genes. Historically, various names have been given to the corresponding gene products, giving rise to



distinct and sometimes confusing nomenclatures. In 1994, a unified but arbitrary nomenclature was adopted in which  $\alpha_1$  subunits were referred to as  $\alpha_{1S}$  for the original skeletal muscle isoform and  $\alpha_{1A}$  through  $\alpha_{1E}$  for those discovered subsequently (Birnbaumer et al., 1994). In 2000, a rational nomenclature was adopted (Ertel et al., 2000) based on the well defined potassium channel nomenclature (Chandy and Gutman, 1993). Calcium channels were named using the chemical symbol of the principal permeating ion (Ca) with the principal physiological regulator (voltage) indicated as a subscript (Ca<sub>V</sub>). The numerical identifier corresponds to the  $Ca_V$  channel  $\alpha_1$  subunit gene subfamily (1 to 3 at present) and the order of discovery of the  $\alpha_1$  subunit within that subfamily (1 through n). According to this nomenclature, the Ca<sub>V</sub>1 subfamily (Ca<sub>V</sub>1.1-Ca<sub>V</sub>1.4) includes channels containing  $\alpha_{1S}$ ,  $\alpha_{1C}$ ,  $\alpha_{1D}$ , and  $\alpha_{1F}$ , which mediate L-type  $Ca^{2+}$  currents (Table 1). The  $Ca_V 2$  subfamily (Ca<sub>V</sub>2.1-Ca<sub>V</sub>2.3) includes channels containing  $\alpha_{1A}$ ,  $\alpha_{1B}$ , and  $\alpha_{1E}$ , which mediate P/Q-type, N-type, and R-type Ca<sup>2+</sup> currents, respectively (Table 1). The Ca<sub>v</sub>3 subfamily (Ca<sub>v</sub>3.1-Ca<sub>v</sub>3.3) includes channels containing  $\alpha_{1G}$ ,  $\alpha_{1H}$ , and  $\alpha_{1I}$ , which mediate T-type Ca<sup>2+</sup> currents.

The complete amino acid sequences of these  $\alpha_1$  subunits are more than 70% identical within a subfamily but less than 40% identical among the three subfamilies. These family relationships are illustrated for the more conserved transmembrane and pore domains in Fig. 2. The division of calcium channels into these three families is phylogenetically ancient, as representatives of each are found in the *Caenorhabditis elegans* genome. Consequently, the genes for the different  $\alpha_1$  subunits have become widely dispersed in the genome, and even the most closely related members of the family are not clustered on single chromosomes in mammals.

# Calcium Channel Molecular Pharmacology

The pharmacology of the three subfamilies of calcium channels is quite distinct. The Ca<sub>v</sub>1 channels are

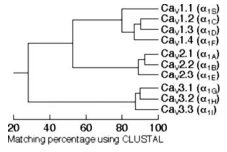


Fig. 2. Sequence similarity of voltage-gated calcium channel  $\alpha_1$  subunits. Phylogenetic representation of the primary sequences of the calcium channels. Only the membrane-spanning segments and pore loops (~350 amino acids) are compared. First, all sequence pairs were compared, which clearly defines three subfamilies with intrafamily sequence identities above 80% (Ca $_{\rm V}$ 1, Ca $_{\rm V}$ 2, and Ca $_{\rm V}$ 3). Then a consensus sequence was defined for each subfamily, and these three sequences were compared to one another, with intersubfamily sequence identities of  $\sim\!52\%$  (Ca $_{\rm V}$ 1 vs. Ca $_{\rm V}$ 2) and 28% (Ca $_{\rm V}$ 3 vs. Ca $_{\rm V}$ 1 or Ca $_{\rm V}$ 2).

the molecular targets of the organic calcium channel blockers used widely in the therapy of cardiovascular diseases. These drugs are thought to act at three separate, but allosterically coupled, receptor sites (Table 1; reviewed in Glossmann and Striessnig, 1990). Phenylalkylamines are intracellular pore blockers, which are thought to enter the pore from the cytoplasmic side of the channel and block it. Their receptor site is formed by amino acid residues in the S6 segments in domains III and IV, in close analogy to the local anesthetic receptor site on sodium channels (Hockerman et al., 1997; Hofmann et al., 1999; Striessnig, 1999). Dihydropyridines can be channel activators or inhibitors and therefore are thought to act allosterically to shift the channel toward the open or closed state rather than by occluding the pore. Their receptor site includes amino acid residues in the S6 segments of domains III and IV and the S5 segment of domain III. The dihydropyridine receptor site is closely apposed to the phenylalkylamine receptor site and shares some common amino acid residues. Diltiazem and related benzothiazepines are thought to bind to a third receptor site, but the amino acid residues that are required for their binding overlap extensively with those required for phenylalkylamine binding.

The  $\rm Ca_V 2$  subfamily of calcium channels is relatively insensitive to dihydropyridine calcium channel blockers, but these calcium channels are specifically blocked with high affinity by peptide toxins from spiders and marine snails (Miljanich and Ramachandran, 1995). The  $\rm Ca_V 2.1$  channels are blocked specifically by  $\omega$ -agatoxin IVA from funnel web spider venom. The  $\rm Ca_V 2.2$  channels are blocked specifically by  $\omega$ -conotoxin GVIA and related cone snail toxins. The  $\rm Ca_V 2.3$  channels are blocked specifically by the synthetic peptide toxin SNX-482 derived from tarantula venom. These peptide toxins are potent blockers of synaptic transmission because of their specific effects on the  $\rm Ca_V 2$  family of calcium channels.

The  $Ca_V3$  subfamily of calcium channels are insensitive to both the dihydropyridines that block  $Ca_V1$  channels and the spider and cone snail toxins that block the  $Ca_V2$  channels, and there are no widely useful pharmacological agents that block T-type calcium currents (Perez-Reyes, 2003). The organic calcium channel blocker mibefradil is somewhat selective for T-type versus L-type calcium currents (3- to 5-fold). The peptide kurtoxin inhibits the activation gating of  $Ca_V3.1$  and  $Ca_V3.2$  channels. Development of more specific and high-affinity blockers of the  $Ca_V3$  family of calcium channels would be useful for therapy and a more detailed analysis of the physiological roles of these channels.

Tables 2 through 11 summarize the major molecular, physiological, and pharmacological properties for each of the 10 calcium channels that have been functionally expressed. Quantitative data are included for

voltage dependence of activation and inactivation, single-channel conductance, and binding of drugs and neurotoxins, focusing on those agents that are widely used and diagnostic of channel identity and function.

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# TABLE 2

$ \begin{array}{c} \text{TABLE 2} \\ Ca_{\text{V}} 1.1 \ channels \end{array} $					
Channel name	Ca <sub>v</sub> 1.1				
Description	Voltage-gated calcium channel $\alpha_1$ -subunit				
Other names	$\alpha_{1s}$ , skeletal muscle L-type Ca <sup>2+</sup> channel, skeletal muscle dihydropyridine receptor				
Molecular information	Human: 1873aa, L33798 (PMID: 7713519), chr0.1q32, CACNA1S, LocusID: 779				
	Rat: 1146aa (partial sequence), L04684 (PMID: 1335956), chr. 13, Cacna1s, LocusID: 116652				
	Mouse: 1861aa, L06234 (PMID: 1281468), chr. 1, Cacna1s, LocusID: 12292 (see 'Comments')				
Associated subunits	$\alpha_2\delta,\beta,\gamma^{1,2}$				
Functional assays	Patch-clamp (whole-cell, single-channel), calcium imaging, gating charge movement, skeletal muscle contraction				
Current	$ m I_{Ca,L}$				
Conductance	13-17pS (in 90–110 mM Ba <sup>2+</sup> ) <sup>3,4</sup>				
Ion selectivity	$Ca^{2+} > Sr^{2+} > Mg^{2+} > Ba^{2+}$				
Activation	$V_{ m a}$ = 8–14 mV, $ au_{ m a}$ $>$ 50 ms at +10 mV (10 mM Ca <sup>2+</sup> ) <sup>4,6</sup>				
Inactivation	$V_{\rm h} = -8$ mV, $40\%$ current inactivation after 5 s $(-5$ mV) <sup>4</sup>				
Activators	BayK8644, dihydropyridine agonists, FPL64176 <sup>2,8,9</sup>				
Gating modifiers	Dihydropyridine antagonists (e.g., (+)- is radipine; $\rm IC_{50}=13~nM~at~-90~mV$ and 0.15 nM at $\rm -65~mV)^9$				
Blockers	Nonselective: cadmium (IC $_{50}$ < 0.5 mM) $^9$ ; selective for Ca $_{ m V}$ 1.x: verapamil, devapamil (IC $_{50}$ < 1 $\mu$ M) and other phenylalkylamines, (+)-cis-diltiazem (IC $_{50}$ < 80 $\mu$ M) $^9$				
Radioligands	(+)-[³H]is radipine ( $K_{\rm d}=0.2-0.7$ nM) and other dihydropyridines; (–)-[³H] devapamil ( $K_{\rm d}=2.5$ nM), (+)- $cis$ -[³H] diltiazem ( $K_{\rm d}=50$ nM)²				
Channel distribution	Skeletal muscle transverse tubules (tetramers) <sup>10</sup>				
Physiological functions	Excitation-contraction coupling and Ca <sup>2+</sup> homeostasis in skeletal muscle <sup>11</sup>				
Mutations and pathophysiology	Point mutations cause hypokalemic periodic paralysis and malignant hyperthermia susceptibility in humans and muscular dysgenesis in mice $(mdg/mdg)^{12,13}$				
Pharmacological significance	Not established				
Comments	The gene for Ca <sub>v</sub> 1.1 was first isolated and characterized in rabbit (1873aa, M23919, X05921);				
	several groups reported three-dimensional structures of the purified skeletal muscle calcium				
	channel complex determined using electron cryomicroscopy and single-particle averaging <sup>14</sup>				
on aming golds, the abromosome, Bay K8644, mathyl 1.4 dibudes 2.6 dimethyl 2 nitro 4.(2 triflyoromethylphonyl) pyriding 5 carboyylate. FDI 64176, mathyl 2.5					

- aa, amino acids; chr., chromosome; Bay K8644, methyl 1,4-dihydro-2,6-dimethyl-3-nitro-4-(2-trifluoromethylphenyl)-pyridine-5-carboxylate; FPL64176, methyl 2,5 dimethyl-4-[2-(phenylmethyl)benzoyl]-1H-pyrrole-3-carboxylate.
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# TABLE 3 Ca<sub>v</sub>1.2 channels

Channel name Description Other names

Molecular information

Associated subunits Functional assays

Current Conductance Ion selectivity Activation

Inactivation

Activators Gating modifiers

Blockers

Radioligands

Channel distribution

Physiological functions

Mutations and pathophysiology

Pharmacological significance

Comments

 $Ca_{y}1.2$ 

Voltage-gated calcium channel  $\alpha_1$  subunit

 $\alpha_{1C}$ , cardiac or smooth muscle L-type Ca<sup>2+</sup> channel, cardiac or smooth muscle dihydropyridine

Human: 2169aa, L29529 (cardiac; PMID: 8392192), 2138aa, Z34815 (fibroblast; PMID: 1316612); 2138aa, AF465484 (jejunum; PMID: 12176756); chr. 12p13.3, CACNA1C, LocusID: 775

Rat: 2169aa, M59786 (aortic smooth muscle; PMID: 2170396); 2140/2143aa, M67516/M67515 (brain; PMID: 1648941); chr. 4q42, Cacna1c, LocusID: 24239

Mouse: 2139aa, L01776 (brain; PMID: 1385406); chr. 6, Cacna1c, LocusID: 12288 (see 'Comments')  $\alpha_2\delta$ ,  $\beta$ ,  $\gamma^{1,2}$ 

Patch-clamp (whole-cell, single-channel), calcium imaging, cardiac or smooth muscle contraction hormone secretion

 $\begin{array}{l} I_{\rm Ca_{L}} \\ Ba^{2+} & (25 {\rm pS}) > Sr^{2+} = Ca^{2+} & (9 {\rm pS})^3 \\ Ca^{2+} > Sr^{2+} > Ba^{2+} \gg Mg^{2+} & {\rm from~permeability~ratios} \\ V_{\rm a} = -17~{\rm mV~(in~2~mM~Ca^{2+};~HEK~cells)^4;~-4~mV~(in~15~mM~Ba^{2+};~HEK~cells)~to~-18.8~mV~(in~5~mM~Ba^{2+};~HEK~cells~and~Xenopus~oocytes)^{5-7};~\tau_{\rm a} = 1~{\rm ms~at~+10~mV^5} \\ V_{\rm h} = -50~{\rm to~-60~mV~(in~2~mM~Ca^{2+};~HEK~cells),^4-18~to~-42~mV~(in~5-15~mM~Ba^{2+};~HEK~cells^8~(at~V_{\rm max}),^{2} \\ I_{\rm callg})^{5,7,8,9};~\tau_{\rm cat} = 150~{\rm ms},~\tau_{\rm slow} = 1100~{\rm ms};~61\%~{\rm inactivated~after~250~ms~in~HEK~cells^8~(at~V_{\rm max}),^{2} \\ I_{\rm callg})^{5,7,8,9};~\tau_{\rm cat} = 150~{\rm ms},~\tau_{\rm slow} = 1100~{\rm ms};~61\%~{\rm inactivated~after~250~ms~in~HEK~cells^8~(at~V_{\rm max}),^{2} \\ I_{\rm callg})^{5,7,8,9};~\tau_{\rm cat} = 150~{\rm ms},~\tau_{\rm slow} = 1100~{\rm ms};~61\%~{\rm inactivated~after~250~ms~in~HEK~cells^8~(at~V_{\rm max}),^{2} \\ I_{\rm callg})^{5,7,8,9};~\tau_{\rm cat} = 150~{\rm ms},~\tau_{\rm slow} = 1100~{\rm ms};~61\%~{\rm inactivated~after~250~ms~in~HEK~cells^{2}),^{2} \\ I_{\rm callg})^{5,7,8,9};~\tau_{\rm cat} = 150~{\rm ms},~\tau_{\rm slow} = 1100~{\rm ms};~61\%~{\rm inactivated~after~250~ms~in~HEK~cells^{2}),^{2} \\ I_{\rm callg})^{5,7,8,9};~\tau_{\rm cat} = 150~{\rm ms},~\tau_{\rm slow} = 1100~{\rm ms};~61\%~{\rm inactivated~after~250~ms~in~HEK~cells^{2}),^{2} \\ I_{\rm callg})^{5,7,8,9};~\tau_{\rm cat} = 150~{\rm ms},~\tau_{\rm slow} = 1100~{\rm ms};~61\%~{\rm inactivated~after~250~ms~in~HEK~cells^{2}),^{2} \\ I_{\rm callg})^{5,7,8,9};~\tau_{\rm cat} = 150~{\rm ms},~\tau_{\rm cat} = 150~{\rm ms},~\tau_{\rm cat} = 1100~{\rm ms};~\tau_{\rm cat} = 1100~{\rm ms},~\tau_{\rm cat} = 1100~{\rm ms},~\tau_{$ cells)<sup>5,7,8,9</sup>;  $\tau_{\rm fast}$  = 150 ms,  $\tau_{\rm slow}$  = 1100 ms; 61% inactivated after 250 ms in HEK cells<sup>8</sup> (at  $V_{\rm max}$  in 15 mM Ba<sup>2+</sup>)<sup>4</sup>; ~70% inactivation after 1 s (at  $V_{\rm max}$  in 2 mM Ca<sup>2+</sup>)<sup>4</sup>; inactivation is accelerated with Ca<sup>2+</sup> as charge carrier (calcium-dependent inactivation: 86% inactivated after 250 ms<sup>8,10</sup>) BayK8644, dihydropyridine agonists, FPL64176<sup>10,11</sup>

Dihydropyridine antagonists (e.g., isradipine,  $IC_{50} = 7$  nM at -60 mV; nimodipine,  $IC_{50} = 139$  nM at  $-80 \text{ mV})^{6,9}$ 

Nonselective:  $Cd^{2+12}$ ; selective for  $Ca_V1.x$ : devapamil ( $IC_{50}=50$  nM in 10 mM  $Ba^{2+}$  at -60 mV) and other phenylalkylamines; diltiazem ( $IC_{50}=33~\mu\mathrm{M}$  in 10 mM  $Ba^{2+}$  at -60 mV and  $0.05\mathrm{Hz})^{12}$ (+)-[ $^3$ H]isradipine ( $K_d < 0.1$  nM) and other dihydropyridines; (-)-[ $^3$ H]devapamil ( $K_d = 2.5$  nM), (+)cis-[<sup>3</sup>H]diltiazem ( $\ddot{K}_{d} = 50 \text{ nM}$ )<sup>11</sup>

Cardiac muscle, smooth muscle (including blood vessels, intestine, lung, uterus); endocrine cells (including pancreatic  $\beta$ -cells, pituitary); neurones<sup>13</sup>; subcellular localization: concentrated on granule-containing side of pancreatic  $\beta$ -cells<sup>14</sup>; neurons (preferentially somatodendritic)<sup>15</sup> Excitation-contraction coupling in cardiac or smooth muscle, action potential propagation in

sinoatrial and atrioventricular node, synaptic plasticity, hormone (e.g., insulin) secretion 10,13,16,17 Required for normal embryonic development (mouse, zebrafish) ed novo G406R mutation in alternative exon 8A in 1 allele causes Timothy syndrome<sup>20</sup>

Mediates cardiovascular effects of clinically used Ca<sup>2+</sup> antagonists<sup>17</sup>; high concentrations of dihydropyridines exert antidepressant effects through Ca<sub>v</sub>1.2 inhibition<sup>1</sup>

Tissue-specific splice variants exist—in addition to cardiac channels, smooth muscle and brain channels have been cloned  $^{7,21,22}$ ; the gene for  $\mathrm{Ca_v}1.2$  was first isolated and characterized in rabbit heart (2171aa, P15381, X15539)

aa, amino acids; chr., chromosome; HEK, human embryonic kidney.

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# TABLE 4 Ca<sub>v</sub>1.3 channels

Channel name	$\mathrm{Ca_{V}1}.$	3
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Description Voltage-gated calcium channel  $\alpha_1$  subunit Other names  $\alpha_{1D}$ , "neuroendocrine" L-type Ca<sup>2+</sup> channel

Human: 2161aa, M76558 (brain; PMID: 1309651); 2181aa, M83566 (pancreatic  $\beta$ -cells; PMID: Molecular information 1309948); chr. 3p14.3, CACNA1D, LocusID: 776

Rat: 1646aa, M57682 (brain; PMID: 1648940); 2203aa, D38101 (pancreatic  $\beta$ -cells; PMID: 7760845);

chr. 16p16, Cacna1d, LocusID: 29716 Mouse: 2144aa, AJ437291 (embryonic heart; PMID: 12900400); chr. 14, Cacna1d, LocusID: 12289

(see "Comments")

Associated subunits Most likely at least  $\alpha_2$ ,  $\beta$ , and  $\delta$  subunits

Functional assays Patch-clamp (whole-cell, single-channel), calcium imaging

Current  $I_{Ca,L}$ 

Pharmacological significance

Comments

Conductance Not established Ion selectivity Not established

Activation  $V_a = -15$  to -20 mV (mouse cochlear hair cells; 10 mM  $Ba^{2+}$ )<sup>1,2</sup>; -18 mV (in 15 mM  $Ba^{2+}$ ; HEK cells) to -37 mV (5 mM Ba $^{2+}$ ; 2 mM Ca $^{2+}$  HEK cells or *Xenopus* oocytes) $^{3,4}$ ;  $au_a < 1$  ms at +10

 $V_{\rm h}=-36$  to -43 mV<sup>3,5</sup>;  $au_{\rm fast}=190$  ms,  $au_{\rm slow}=1700$  ms (at  $V_{\rm max}$  in HEK cells)<sup>3</sup>; calcium-induced Inactivation inactivation is observed after expression in HEK cells<sup>3</sup> and in cochlear outer hair cells but not in

inner hair cells<sup>2</sup>

BayK86441-5 Activators

Gating modifiers Dihydropyridine antagonists (e.g., isradipine,  $IC_{50} = 30$  nM at -50 mV and 300 nM at -90 mV;

nimodipine, IC<sub>50</sub> = 3  $\mu$ M at -80 mV)<sup>3,4</sup>

Nonselective: Cd<sup>2+</sup>

Blockers (+)-[ $^{3}$ H]isradipine ( $K_{\rm d} < 0.5 \text{ nM}$ ) $^{3}$ ; in radioreceptor assays, HEK cell-expressed Ca<sub>v</sub>1.2 and Ca<sub>v</sub>1.3 Radioligands

> channels bind (+)- $[^{3}H]$  is radipine with indistinguishable  $K_{D}^{3}$ ; in functional experiments, however, Ca<sub>v</sub>1.2 channels show higher DHP sensitivity—this discrepancy is explained by the slower inactivation of Ca<sub>v</sub>1.3 decreasing the availability of inactivated channels for state-dependent DHP

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block

Sensory cells (photoreceptors, cochlear hair cells  $^{1,2}$ ), endocrine cells (including pancreatic  $\beta$ -cells, Channel distribution pituitary, adrenal chromaffin cells, pinealocytes), 7-9 low density in heart (atrial muscle, sinoatrial and atrioventricular node)<sup>1,7,10</sup> and vascular smooth muscle<sup>7</sup>; neurones<sup>6</sup>; subcellular localization:

on neurones preferentially located on proximal dendrites and cell bodies<sup>6</sup>

Neurotransmitter release in sensory cells, control of cardiac rhythm and atrioventricular node

Physiological functions conductance at rest, 1,10,12 mood behavior, 12 hormone secretion

Deafness, sinoatrial and atrioventricular node dysfunction, 1,10,12 no convincing evidence for Mutations and pathophysiology contribution to pancreatic  $\beta$ -cell L-type currents and insulin secretion in mouse models<sup>1,12,13</sup>

Hypothetical drug targets for modulators of heart rate, 1 antidepressant drugs 10 and drugs for hearing disorders<sup>1</sup>

Tissue-specific and developmental (exon 1b) splice variants exist—in addition to brain, pancreatic β-cell and cochlear variants have been cloned; it is likely that Ca<sub>v</sub>1.3 channels form most of the so-called 'low-voltage-activated' L-type currents found in the brain and sinoatrial node, although some splice variants of Ca<sub>v</sub>1.2 can also activate at more negative potentials

aa, amino acids; chr., chromosome; HEK, human embryonic kidney; DHP, dihydropyridine.

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# TABLE 5 Ca<sub>V</sub>1.4 channels

 $Ca_{v}1.4$ Channel name

Description Voltage-gated calcium channel  $\alpha_1$  subunit

Other names

Human: 1966aa, AJ224874 (PMID: 9662399); chr. Xp11.23, CACNA1F, LocusID: 778 Molecular information

Rat: 1981aa, AF365975 (PMID: 11526344); chr. Xq22, Cacna1f, LocusID: 114493 Mouse: 1985aa, AF192497 (PMID: 10873387); chr. X, Cacna1f, LocusID: 54652

Associated subunits Not established; preliminary functional evidence for  $\beta_2$  association in retinal neurons<sup>1</sup>

Functional assays Patch-clamp (whole-cell, single-channel), calcium imaging

Current

Preliminary evidence for very small single channel conductance (less than half of Ca<sub>v</sub>1.2); Ba<sup>2+</sup> > Conductance

 $Ca^{2+2,4,6}$ 

Not established Ion selectivity

 $V_{\rm a}=-2.5$  to -12 mV (2–20 mM Ca<sup>2+</sup> or 15–20 mM Ba<sup>2+</sup>; HEK cells)<sup>3–6</sup>;  $au_{\rm a}<1$  ms at  $V_{\rm max}$  (but Activation

slower components were also observed)3,6

Inactivation  $V_{\rm h} = -9$  to -27 mV (10–20 mM Ba $^{2+}$ , HEK cells) $^{4,6}$ ; inactivation kinetics even slower than those of

 $\mathrm{Ca_v}1.3$  with incomplete inactivation during 10-s depolarizations to  $V_{\mathrm{max}}{}^3$ ; calcium-induced inactivation is not observed for  $Ca_v1.4$  channels expressed in HEK cells<sup>3,4,6</sup> but after expression in

Xenopus oocytes2

BayK8644<sup>2-4,6</sup> Activators Gating modifiers Dihydropyridine antagonists: nifedipine (IC $_{50}$  = 944 nM at -100 mV,  $\sim 300$  nM at -50 mV $^4$ ;

isradipine:  $\sim$ 80% inhibition by 100 nM at  $-50 \text{ mV}^{3,6}$  and 1  $\mu\text{M}$  at  $-90 \text{ mV})^3$ ; D-cis-diltiazem  $(IC_{50}=92 \mu M)$ ; verapamil: 69% inhibition at 100  $\mu M$  (0.2 Hz, holding potential =  $-80 \text{ mV})^6$ 

Blockers Nonselective: Cd2+2

Radioligands Unlike for Ca<sub>v</sub>1.2 and Ca<sub>v</sub>1.3, no high-affinity (+)-[<sup>3</sup>H]isradipine binding detectable (HEK cells)

(J. Striessnig, unpublished observations)

Channel distribution Retinal photoreceptors and bipolar cells, spinal cord, lymphoid tissue (plasma and mast cells)<sup>1,4,7-10</sup>

Physiological functions Neurotransmitter release in retinal cells

Mutations cause X-linked congenital stationary night blindness type  $2^{7,9,11,12}$ Mutations and pathophysiology

Pharmacological significance Not established

Comments The biophysical properties of heterologously expressed Ca, 1.4 channels resemble those recorded in retinal neurons, suggesting that this channel type underlies retinal I<sub>Ca,L</sub>—however, similar to Ca, 1.4, Ca, 1.3 channels also inactivate slowly and activate rapidly and may therefore also

contribute to retinal  $I_{Ca,L}$ 

aa, amino acids; chr., chromosome; HEK, human embryonic kidney.

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# Spet

TABLE 6  $Ca_{V}2.1$  channels

Channel name Ca<sub>V</sub>2.1

Description Voltage-gated calcium channel  $\alpha_1$  subunit

Other names  $\alpha_{1A}$ , P-type, Q-type, rbA-I (in rat)<sup>1</sup>; BI-1, BI-2 (in rabbit)<sup>2</sup>

Molecular information Human: 2510aa, AF004883, 2662aa, AF004884, chr. 19p13, CACNA1A

Rat: 2212aa, M64373

Mouse: 2165aa, NM007578, NP031604 Rabbit: 2273aa, X57476 (see "Comments")

Associated subunits  $\alpha_2 \delta$ ,  $\beta$ , possibly  $\gamma$ 

Functional assays Voltage-clamp, patch-clamp, calcium imaging, neurotransmitter release

Current  $I_{Ca,P}$ ,  $I_{Ca,Q}$ 

Conductance 9, 14, 19pS (P-type, cerebellar Purkinje neurones)<sup>4</sup>; 16–17pS (for  $\alpha_{1A}/\alpha_2\delta/\beta$  in Xenopus oocytes)<sup>2,5,6</sup>

Ion selectivity  $Ba^{2+} > Ca^{2+}$ 

Activation  $V_a = -5 \text{ mV}$  for native P-type,  $V_a = -11 \text{ mV}$  for native Q-type (with 5 mM Ba<sup>2+</sup> charge carrier)<sup>7</sup>

 $V_{\rm a} = -4.1~{\rm mV}$  for rat  $\alpha_{\rm 1A\text{-}a}\!/\alpha_{\rm 2}\delta\!/\beta_{\rm 4}$ 

 $V_{\rm a}=+2.1~{\rm mV}$  for rat  $\alpha_{\rm 1A-b}/\alpha_2\delta/\beta_4$  (with 5 mM Ba<sup>2+</sup> charge carrier)<sup>6</sup>

 $V_{\rm a}=+9.5$  mV;  $au_{\rm a}=2.2$  ms at +10 mV for human  $lpha_{\rm 1A-1}/lpha_2\delta/eta_{\rm 1b}$  in HEK293 cells (with 15 mM Ba<sup>2+</sup>

charge carrier)3

 $V_{\rm h} = -17.2 \text{ mV for } \alpha_{\rm 1A-a}/\alpha_2 \delta/\beta_4; V_{\rm h} = -1.6 \text{ mV for } \alpha_{\rm 1A-b}/\alpha_2 \delta/\beta_4 \text{ (with 5 mM Ba}^{2+} \text{ charge carrier)}; V_{\rm h} = -1.6 \text{ mV for } \alpha_{\rm 1A-b}/\alpha_2 \delta/\beta_4 \text{ (with 5 mM Ba}^{2+} \text{ charge carrier)}; V_{\rm h} = -1.6 \text{ mV for } \alpha_{\rm 1A-b}/\alpha_2 \delta/\beta_4 \text{ (with 5 mM Ba}^{2+} \text{ charge carrier)}; V_{\rm h} = -1.6 \text{ mV for } \alpha_{\rm 1A-b}/\alpha_2 \delta/\beta_4 \text{ (with 5 mM Ba}^{2+} \text{ charge carrier)}; V_{\rm h} = -1.6 \text{ mV for } \alpha_{\rm 1A-b}/\alpha_2 \delta/\beta_4 \text{ (with 5 mM Ba}^{2+} \text{ charge carrier)}; V_{\rm h} = -1.6 \text{ mV for } \alpha_{\rm 1A-b}/\alpha_2 \delta/\beta_4 \text{ (with 5 mM Ba}^{2+} \text{ charge carrier)}; V_{\rm h} = -1.6 \text{ mV for } \alpha_{\rm 1A-b}/\alpha_2 \delta/\beta_4 \text{ (with 5 mM Ba}^{2+} \text{ charge carrier)}; V_{\rm h} = -1.6 \text{ mV for } \alpha_{\rm 1A-b}/\alpha_2 \delta/\beta_4 \text{ (with 5 mM Ba}^{2+} \text{ charge carrier)}; V_{\rm h} = -1.6 \text{ mV for } \alpha_{\rm 1A-b}/\alpha_2 \delta/\beta_4 \text{ (with 5 mM Ba}^{2+} \text{ charge carrier)}; V_{\rm h} = -1.6 \text{ mV for } \alpha_{\rm 1A-b}/\alpha_2 \delta/\beta_4 \text{ (with 5 mM Ba}^{2+} \text{ charge carrier)}; V_{\rm h} = -1.6 \text{ mV for } \alpha_{\rm 1A-b}/\alpha_2 \delta/\beta_4 \text{ (with 5 mM Ba}^{2+} \text{ charge carrier)}; V_{\rm h} = -1.6 \text{ mV for } \alpha_{\rm 1A-b}/\alpha_2 \delta/\beta_4 \text{ (with 5 mM Ba}^{2+} \text{ charge carrier)}; V_{\rm h} = -1.6 \text{ mV for } \alpha_{\rm 1A-b}/\alpha_2 \delta/\beta_4 \text{ (with 5 mM Ba}^{2+} \text{ charge carrier)}; V_{\rm h} = -1.6 \text{ mV for } \alpha_{\rm 1A-b}/\alpha_2 \delta/\beta_4 \text{ (with 5 mM Ba}^{2+} \text{ charge carrier)}; V_{\rm h} = -1.6 \text{ mV for } \alpha_{\rm 1A-b}/\alpha_2 \delta/\beta_4 \text{ (with 5 mM Ba}^{2+} \text{ charge carrier)}; V_{\rm h} = -1.6 \text{ mV for } \alpha_{\rm 1A-b}/\alpha_2 \delta/\beta_4 \text{ (with 5 mM Ba}^{2+} \text{ charge carrier)}; V_{\rm h} = -1.6 \text{ mV for } \alpha_{\rm 1A-b}/\alpha_2 \delta/\beta_4 \text{ (with 5 mM Ba}^{2+} \text{ charge carrier)}; V_{\rm h} = -1.6 \text{ mV for } \alpha_{\rm 1A-b}/\alpha_2 \delta/\beta_4 \text{ (with 5 mM Ba}^{2+} \text{ charge carrier)}; V_{\rm h} = -1.6 \text{ mV for } \alpha_{\rm 1A-b}/\alpha_2 \delta/\beta_4 \text{ (with 5 mM Ba}^{2+} \text{ charge carrier)}; V_{\rm h} = -1.6 \text{ mV for } \alpha_{\rm 1A-b}/\alpha_2 \delta/\beta_4 \text{ (with 5 mM Ba}^{2+} \text{ charge carrier)}; V_{\rm h} = -1.6 \text{ mV for } \alpha_{\rm 1A-b}/\alpha_2 \delta/\beta_4 \text{ (with 5 mM Ba}^{2+} \text{ charge carrier)}; V_{\rm h} = -1.6 \text{ mV for } \alpha_{\rm 1A-b}/\alpha_2 \delta/\beta_4 \text{ (with 5 mM$ 

= -17 mV,  $\tau_{\rm h} = 690$  ms at +10 mV human  $\alpha_{1\text{A}-1}/\alpha_2 \delta/\beta_{1\text{b}}$  in HEK293 cells (with 15 mM Ba<sup>2+</sup> charge carrier)<sup>3</sup>;  $\tau_{\rm h} > 1$  s at 0 mV native P-type (with 5 mM Ba<sup>2+</sup> charge carrier)<sup>7</sup> (see

"Comments")

Activators

Gating modifiers  $\omega\text{-agatoxin IVA (P-type }K_{\mathrm{d}}=1\text{--3 nM}^{8};\text{ Q-type }K_{\mathrm{d}}\sim100\text{--}200\text{ nM}^{5,9}),\;\omega\text{-agatoxin IVB}^{6}$ 

Blockers  $\omega$ -conotoxin MVIIC<sup>8</sup>; other blockers include piperidines, substituted diphenylbutylpiperidines, piperazines, volatile anesthetics, gabapentin, mibefradil, and peptide toxins DW13.3 and  $\omega$ -

conotoxin SVIB<sup>21–26</sup> (see "Comments")

Radioligands  $[^{125}I]\omega$ -conotoxin MVIIC

Channel distribution Neurons (presynaptic terminals, dendrites, some cell bodies), heart, pancreas, pituitary

Physiological functions Neurotransmitter release in central neurons and neuromuscular junction; excitation-secretion

coupling in pancreatic  $\beta$ -cells

Mutations and pathophysiology Missense mutations in IS4-IS5, IIS4-IIS6, IIIS4-IIIS6, and IVS4-IVS6 cause FHM<sup>27</sup>; a common

feature among FHM mutations is an apparent gain-of-function phenotype as a result of a shift in  $V_{50\rm act}$  to more hyperpolarized potentials (an increased probability of opening at the single channel level)<sup>28,29</sup>; other effects include a decrease in maximal current density at the whole-cell level and alterations of synaptic transmission<sup>28–31</sup>; point mutations in IIS1, IIS6-IIIS2, IIIS5-IIIS6, and IVS1-IVS5 cause episodic ataxia type-2, a polyglutamine expansion in the carboxyl region causes spinocerebellar ataxia type-6, and mutation of IS5-IS6 and IVS6 causes episodic and progressive

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ataxia $^{10-12,27}$ 

Pharmacological significance

Peptide toxins that selectively inhibit  $\rm Ca_{v}2.1$  channel block a significant portion of neurotransmission in the mammalian CNS<sup>13</sup>; block of  $\rm Ca_{v}2.1$  channels inhibits the late-phase formalin response and inflammatory pain but has no significant effect on mechanical allodynia or thermal hyperalgesia<sup>14–17</sup>; mice lacking a functional  $\rm Ca_{v}2.1$  gene exhibit cerebellar atrophy, severe muscle spasms, and ataxia and usually die by 3 to 4 weeks postnatal<sup>18,19</sup>

Comments

Rates of inactivation and  $V_h$  are differentially affected by coexpression with  $\beta_{1b}$ ,  $\beta_{2a}$ ,  $\beta_3$ , or  $\beta_4$  subunits, as well as by alternative splicing of the  $\alpha_{1A}$  subunit; identified regions of alternative splicing include the domain I-II linker, domain II-III linker, IVS3-IVS4, and the carboxyl terminus<sup>1,2,6,32-34</sup>; whole-cell currents with P-type kinetics seem to be conducted by the  $\alpha_{1A-b}$  splice variant coexpressed with any of the  $\beta$  subunits or by the  $\alpha_{1A-a}$  splice variant coexpressed with the  $\beta_{2a}$  subunities, whole-cell currents with Q-type kinetics seem to be encoded by  $\alpha_{1A-a}$  coexpressed with any of the  $\beta_{1b}$ ,  $\beta_3$ , or  $\beta_4$  subunities, whole-cell currents with Q-type pharmacology seem to be encoded by  $\alpha_{1A}$  splice variants containing Asp Pro residues in the domain IV S3-S4 linker, whereas whole-cell currents with P-type pharmacology seem to be conducted by  $\alpha_{1A}$  splice variants missing Asp Pro residues in IV S3-S4 linker, alternative splicing also alters current density, current-voltage relations, calcium/calmodulin-dependent facilitation, sensitivity to mibefradil, and binding to intracellular synaptic proteins such as Mint1, CASK, syntaxin, and SNAP-25<sup>26,32,36</sup>

aa, amino acids; chr., chromosome; HEK, human embryonic kidney; FHM, familial hemiplegic migraine; CNS, central nervous system.

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Ca<sub>v</sub>2.2 channels

Comments

TABLE 7

 $Ca_{V}2.2$ Channel name

Description Voltage-gated calcium channel  $\alpha_1$  subunit

Other names N-type,  $\alpha_{1B}$ ; rbB-I, rbB-II (in rat), <sup>1,2</sup> BIII (in rabbit)<sup>3</sup>

Human: 2339aa, M94172, 2237aa, M94173, chr. 9q34, CACN1B Molecular information

Rat: 2336aa, M929051

Mouse: 2329aa, NM007579, NP031605

 $\alpha_2\delta/\beta_1$ ,  $\beta_3$ ,  $\beta_4$ , possibly  $\gamma$ Associated subunits

Functional assays Voltage-clamp, patch-clamp, calcium imaging, neurotransmitter release, <sup>45</sup>Ca uptake into

Current  $I_{\mathrm{Ca},N}$ 

Conductance 20pS (bullfrog sympathetic neurones)<sup>6</sup>; 14.3pS (rabbit BIII cDNA in skeletal muscle myotubes)<sup>3</sup>

 $Ba^{2+} > Ca^{2+}$ Ion selectivity

 $V_{\rm a}=+7.8$  mV,  $\tau_{\rm a}=3$  ms at +10 mV (human  $\alpha_{\rm 1B}/\alpha_{\rm 2}\delta/\beta_{\rm 1-3}$  in HEK293 cells, 15 mM Ba<sup>2+</sup> charge Activation

carrier)<sup>4,7</sup>;  $V_a = +9.7$  mV,  $\tau_a = 2.8$  ms at +20 mV (rat  $\alpha_{1B-II}/\beta_{1b}$ , in *Xenopus* oocytes, 40 mM Ba<sup>2+</sup>

charge  $carrier)^2$ 

 $V_{\rm h}=-61$  mV,  $\tau_{\rm h}\sim\!200$  ms at +10 mV (human  $\alpha_{\rm 1B}$  / $\alpha_{\rm 2}\delta$ / $\beta_{\rm 1-3}$  in HEK293 cells, 15 mM Ba<sup>2+</sup> charge Inactivation

carrier)<sup>4,7</sup>;  $V_h = -67.5$  mV;  $\tau_h = 112$  ms at +20 mV (rat  $\alpha_{1B-II}/\beta_{1b}$  in Xenopus oocytes, 40 mM

 $Ba^{2+})^2$ 

Activators None Gating modifiers None

Blockers ω-conotoxin GVIA (1–2 μM, irreversible block), ω-conotoxin MVIIA (SNX-111, Ziconotide/Prialt), ωconotoxin MVIIC8; other blockers include piperidines, substituted diphenylbutylpiperidines, long alkyl chain molecules, aliphatic monoamines, tetrandine, gabapentin, peptidylamines, volatile

> anesthetics, the peptide toxins SNX-325 and DW13.3, as well as the ω-conotoxins SVIA, SVIB, and CVID<sup>20-34</sup>

Radioligands [125]]ω-conotoxin GVIA ( $K_{\rm d}=55$  pM, human  $\alpha_{\rm 1B}$  / $\alpha_{\rm 2}$ δ/ $\beta_{\rm 1-3}$  in HEK293 cells)<sup>4</sup>

Channel distribution Neurons (presynaptic terminals, dendrites, cell bodies)<sup>9</sup>

Physiological functions Neurotransmitter release in central and sympathetic neurons 10; sympathetic regulation of the circulatory system<sup>11,35</sup>; activity and vigilance state control<sup>36</sup>; sensation and transmission of pain

(see "Pharmacological significance" and "Comments") Mutations and pathophysiology Differing reports exist: mice lacking a functional Cav2.2 gene exhibit a normal life span and no

> detectable behavioral modifications compared with wild type but possess an increase in basal mean atrial pressure and other functional alterations to the sympathetic nervous system<sup>11</sup>—however, in a different study, approximately 1/3 of the mice lacking a functional Ca<sub>V</sub>2.2 gene did not survive to weaning, but surviving animals were normal except for a decrease in

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anxiety-related behavior and a suppression of inflammatory and neuropathic pain responses 12; no point mutations in the native Ca<sub>V</sub>2.2. gene have been reported to date In rats, intrathecal administration of ω-conotoxin GVIA or ω-conotoxin MVIIA shows strong effects Pharmacological significance on inflammatory pain, postsurgical pain, thermal hyperalgesia, and mechanical allodynia 13-15; in

humans, intrathecal administration of SNX-111 (Ziconotide/Prialt, synthetic ω-conotoxin MVIIA) to patients unresponsive to intrathecal opiates significantly reduced pain scores and in a number of specific instances resulted in relief after many years of continuous pain 16 In case studies, Ziconotide/Prialt has been examined for usefulness in the management of

intractable spasticity following spinal cord injury in patients unresponsive to baclofen and morphine<sup>17</sup>; side effects of intrathecal administration of Ziconotide/Prialt include nystagmus, sedation, confusion, auditory and visual hallucinations, severe agitation, and unruly behavior<sup>18</sup>; intravenous administration of Ziconotide to humans results in significant orthostatic hypotension<sup>19</sup>; identified regions of alternative splicing include the domain I-II linker, domain II-III linker, IIIS3-IIIS4, IVS3-IVS4, and the carboxyl terminus<sup>1-4,37-39</sup>; splicing affects a number of channel properties, including current-voltage relations and kinetics, and is associated with cellspecific expression—in particular, expression of the e37a splice isoform in dorsal root ganglia correlates with a subset of nociceptive neurons<sup>40–42</sup>; alternative splicing also alters interactions with intracellular synaptic proteins such as Mint1, CASK, syntaxin, and SNAP-25<sup>43-45</sup>

aa, amino acid; chr., chromosome; HEK, human embryonic kidney.

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TABLE 8  $Ca_{V}2.3\ channels$ 

Channel name Ca<sub>V</sub>2.3

Description Voltage-gated calcium channel  $\alpha_1$  subunit

Other names R-type,  $\alpha_{1E}$ ; rbE-II (in rat)<sup>1</sup>; BII-1, BII-2 (in rabbit)<sup>2</sup>

Molecular information Human: 2251aa, L29384, 2270aa, L29385, chr0.1q25-q31, CACNA1E

Rat: 2222aa,¹ GenBank accession no. L15453

Mouse: 2272aa, Q61290

Associated subunits  $\alpha_2 \delta/\beta$ , possibly  $\gamma$ 

Functional assays Voltage-clamp, patch-clamp, calcium imaging, neurotransmitter release

Current  $I_{Ca}$ 

Conductance Not established

Ion selectivity  $Ba^{2+} \sim Ca^{2+} \ (rat)^4; \ Ba^{2+} > Ca^{2+} \ (human)^3$ 

Activation  $V_a = +3.5$  mV,  $\tau_a = 1.3$  ms at 0 mV (human  $\alpha_{1E}/\alpha_2 \delta/\beta_{1-3}$ , 15 mM Ba<sup>2+</sup> charge carrier in HEK293

cells)

 $V_{\rm a}=-29.1$  mV,  $\tau_{\rm a}=2.1$  ms at -10 mV (rat  $\alpha_{\rm 1E}/\alpha_{\rm 2}\delta/\beta_{\rm 1b}, 4$  mM Ba<sup>2+</sup> charge carrier in Xenopus

oocytes)1

Inactivation  $V_h = -71 \text{ mV}, \tau_h = 74 \text{ ms at } 0 \text{ mV} \text{ (human } \alpha_{1E}/\alpha_2 \delta/\beta_{1-3}, 15 \text{ mM Ba}^{2+} \text{ charge carrier in HEK293}$ 

cells)<sup>3</sup>;  $V_{\rm h}=-78.1$  mV,  $\tau_{\rm h}=100$  ms at -10 mV (rat  $\alpha_{\rm 1E}/\alpha_{\rm 2}\delta/\beta_{\rm 1b}$ , 4 mM Ba<sup>2+</sup> charge carrier in

Xenopus oocytes)<sup>1</sup>

Activators None Gating modifiers None

Blockers SNX-482, Ni  $^{2+}$  (IC  $_{50} = 27~\mu\text{M}$ ), Cd  $^{2+}$  (IC  $_{50} = 0.8~\mu\text{M}$ ), mibefradil (IC  $_{50} = 0.4~\mu\text{M}$ ),  $^{10}$  volatile

anesthetics<sup>11</sup>

Radioligands None

Channel distribution Neurons (cell bodies, dendrites, some presynaptic terminals), heart, testes, pituitary

Physiological functions Neurotransmitter release, repetitive firing, long-term potentiation, post-tetanic potentiation,

neurosecretion 12-14

Mutations and pathophysiology

No point mutations in the native Ca<sub>v</sub>2.3 gene have been reported; mice deficient for the Ca<sub>v</sub>2.3 gene retain a substantial cerebellar R-type current,<sup>5</sup> suggesting that R-type currents actually reflect a heterogeneous mixture of channels; homozygous Ca<sub>v</sub>2.3-null mice survive to adulthood, reproduce, and are apparently behaviorally normal<sup>5,6</sup>; mutant mice exhibit an increased resistance to formalin-induced pain, suggesting an involvement of the Ca<sub>v</sub>2.3 calcium channel in transmitting and/or the development of somatic inflammatory pain<sup>6</sup>

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Pharmacological significance

Comments

See "Comments"

Ca<sub>v</sub>2.3 has been variously reported to encode a novel type of calcium channel with properties shared between both low- and high-threshold calcium channels<sup>1,4</sup> or a type of high-threshold channel resistant to DHPs, ω-agatoxin-IVA, and ω-conotoxin-GVIA and called R-type (for "residual")<sup>7</sup>

The tarantula toxin SNX-482 blocks exogenously expressed  $Ca_v2.3$  currents<sup>8</sup> but is only partially effective on native cerebellar R-type currents,<sup>9</sup> suggesting that  $Ca_v2.3$  does not always conduct a significant portion of the R-type current as originally defined<sup>7</sup>; identified regions of alternative splicing include the domain II-III linker and carboxyl terminus and have been shown to affect channel kinetics and  $Ca^{2+}$ -dependent stimulation<sup>1-3,15,16</sup>

aa, amino acids; chr., chromosome; HEK, human embryonic kidney; DHP, dihydropyridine.

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# TABLE 9 $Ca_{v}3.1\ channels$

Channel name Ca<sub>v</sub>3.1

Description Voltage-gated calcium channel  $\alpha_1$  subunit

Molecular information Human: 2377aa, O43497, NM\_018896, chr. 17q22, CACNA1G<sup>1</sup>

Rat: 2254aa, O54898, AF027984

Mouse: 2288aa, CAI25956, NM\_009783 (see "Comments")

Associated subunits No biochemical evidence, small changes induced by  $\alpha_2 \delta_1^2$  and  $\alpha_2 \delta_2^{3,4}$ 

Functional assays Voltage-clamp, calcium imaging

 $\begin{array}{ll} Current & I_{Ca,T} \\ Conductance & 7.5pS^1 \end{array}$ 

Ion selectivity  $Sr^{2+} > Ba^{2+} = Ca^{2+}$ 

 $\begin{array}{ll} \mbox{Activation} & V_{\rm a} = -46 \ \mbox{mV}, \ \tau_{\rm a} = 1 \ \mbox{ms at } -10 \ \mbox{mV}^{5,6} \\ \mbox{Inactivation} & V_{\rm h} = -73 \ \mbox{mV}, \ \tau_{\rm h} = 11 \ \mbox{ms at } -10 \ \mbox{mV}^{5,6} \\ \end{array}$ 

Activators Not established

Gating modifiers Kurtoxin,  $IC_{50} = 15 \text{ nM}^7$ 

Blockers No subtype-specific blocker<sup>8</sup>; selective for Ca<sub>v</sub>3.x relative to Ca<sub>v</sub>1.x and Ca<sub>v</sub>2.x: mibefradil, <sup>9,10</sup>

U92032, 11 penfluridol and pimozide 12; nonselective: nickel (IC<sub>50</sub> = 250  $\mu$ M), 13 amiloride 14

Radioligands None

Channel distribution Brain, especially soma and dendrites of neurons in olfactory bulb, amygdala, cerebral cortex,

hippocampus, thalamus, hypothalamus, cerebellum, brain stem (human RNA blots, <sup>1,5</sup> rat in situ hybridization <sup>15</sup> and immunocytochemistry <sup>16</sup>); ovary, placenta, heart (especially sinoatrial node;

mouse in situ hybridization<sup>17</sup>)

Physiological functions Thalamic oscillations 18

Mutations and pathophysiology Not established

Pharmacological significance May mediate effect of absence antiepileptic drugs such as ethosuximide 19 and other thalamocortical

dysrhythmias<sup>20</sup>

Comments Splice variants that differ in their voltage dependence have been cloned<sup>5</sup>

aa, amino acids; chr., chromosome; U92032, 7-[[4-bis(fluorophenyl)methyl]-1-piperazinyl]methyl-2-[(2-hydroxyethyl)amino]4-(1-methylethyl)-2,4,6-cycloheptatrien-1-one.

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# TABLE 10 Ca<sub>v</sub>3.2 channels

Channel name Ca..3.2

Description Voltage-gated calcium channel  $\alpha_1$  subunit

T-type,  $\alpha_1 3.2$ ,  $\alpha_{1H}$ Other names

Human: 2353aa, O95180, AF051946, chr0.16p13.3, CACNA1H1 Molecular information

> Rat: 2359aa, AAG35187, AF290213 Mouse: 2365aa, NP\_067390, NM\_021415

Associated subunits Not established

Functional assays Voltage-clamp, calcium imaging

 $\begin{array}{c} I_{\mathrm{Ca,T}} \\ 9 p S^2 \end{array}$ Current Conductance  $Ba^{2+} = Ca^{2+}$ Ion selectivity

 $V_{\rm a}=-46$  mV,  $\tau_{\rm a}=2$  ms at -10 mV  $^3$   $V_{\rm h}=-72$  mV,  $\tau_{\rm h}=16$  ms at -10 mV  $^3$ Activation Inactivation

Activators None Kurtoxin<sup>4</sup> Gating modifiers

Blockers  $\text{Ca}_{\text{v}}3.2$  is more sensitive than  $\text{Ca}_{\text{v}}3.1$  to block by nickel (IC<sub>50</sub> = 12  $\mu\text{M}$ )<sup>5</sup> and possibly phenytoin<sup>6</sup> and

amiloride $^{2,7}$ ; selective for  $Ca_v3.x$  relative to  $Ca_v1.x$  and  $Ca_v2.x$ : mibefradil,  $^{8,9}$  U92032,  $^{10}$  penfluridol and pimozide,  $^{11}$  and amiloride $^{12}$ ; nonselective: nimodipine,  $^{2}$  anesthetics $^{5}$ 

Radioligands

Kidney (human Northern<sup>1</sup>), rat smooth muscle (RT-PCR<sup>13</sup>), liver (human Northern<sup>1</sup>), adrenal cortex Channel distribution

(rat, bovine; in situ hybridization and RT-PCR<sup>14</sup>), brain (especially in olfactory bulb, striatum, cerebral cortex, hippocampus, reticular thalamic nucleus; rat in situ hybridization<sup>15</sup>), and heart Downloaded from pharmrev.aspetjournals.org by guest on

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(especially sinoatrial node; mouse in situ hybridization<sup>16</sup>) Smooth muscle contraction,<sup>17</sup> smooth muscle proliferation,<sup>18</sup> aldosterone secretion,<sup>19</sup> cortisol Physiological functions

 $secretion^{20}$ 

Single nucleotide polymorphisms associated with childhood absence epilepsy patients in a Chinese Mutations and pathophysiology

May mediate effect of absence antiepileptic drugs such as ethosuximide  $^{22}$  and other thalamocortical Pharmacological significance

dysrhythmias<sup>23</sup>; potential drug target in hypertension and angina pectoris<sup>24</sup>

Splice variation found in the linker connecting repeat 3 and  $4^{25}$ 

aa, amino acids; chr., chromosome; RT-PCR, reverse-transcriptase-polymerase chain reaction. 1. Cribbs LL, Lee JH, Yang J, Satin J, Zhang Y, Daud A, Barclay J, Williamson MP, Fox M, Rees M, et al. (1998) Cloning and characterization of  $\alpha_{1H}$  from human heart, a member of the T-type Ca2+ channel gene family. Circ Res 83:103-109.

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# TABLE 11 Ca<sub>v</sub>3.3 channels

Ca<sub>v</sub>3.3 Channel name

Description Voltage-gated calcium channel α<sub>1</sub> subunit

T-type,  $\alpha_1 3.3$ ,  $\alpha_{1I}$ Other names

Human: 2251aa, AAM67414, AF393329, chr. 22q13.1, CACNA11 1 Molecular information

> Rat: 1835aa, AF086827, AAD17796 Mouse 2753aa: XP\_139476, XM\_139476

Associated subunits No biochemical evidence, small changes induced by  $\gamma_2^2$ 

Functional assays Voltage-clamp, calcium imaging

Current  $11pS^1$ Conductance  $Ba^{2+} = Ca^{2+}$ Ion selectivity

 $V_{\rm a} = -44$  mV,  $\tau_{\rm a} = 7$  ms at -10 mV<sup>4</sup> Activation  $V_{\rm h} = -72 \text{ mV}, \ \tau_{\rm h} = 69 \text{ ms at } -10 \text{ mV}^4$ Inactivation

Activators Not established Gating modifiers None

Blockers No subtype-specific blocker<sup>5</sup>; selective for Ca<sub>v</sub>3.x relative to Ca<sub>v</sub>1.x and Ca<sub>v</sub>2.x: mibefradil, <sup>6,7</sup>

U92032, penfluridol, pimozide; nonselective: nickel (IC<sub>50</sub> = 216  $\mu$ M)<sup>10</sup>

Radioligands

Channel distribution Brain, especially olfactory bulb, striatum, cerebral cortex, hippocampus, reticular nucleus, lateral

habenula, cerebellum (rat in situ hybridization, 11 human Northern 12)

Thalamic oscillations<sup>13</sup> Physiological functions Mutations and pathophysiology Not established

Pharmacological significance May mediate effect of absence antiepileptic drugs such as ethosuximide<sup>14</sup> and other thalamocortical

dysrhythmias<sup>15</sup>

Comments Splice variants have been reported<sup>16</sup>

aa, amino acids; chr., chromosome.

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