Isoform-specific regulation of mood behavior and pancreatic β cell and cardiovascular function by L-type Ca\(^{2+}\) channels

Martina J. Sinneger-Brauns, Alfred Hetzenauer, Irene G. Huber, Erik Renström, Georg Wietzorredek, Stanislav Berjukov, Maurizio Cavalli, Doris Walter, Alexandra Koschak, Ralph Waldschütz, Steffen Hering, Sergio Bova, Patrik Rorsman, Olaf Pongs, Nicolas Singewald, and Jörg Striessnig

Introduction

Ca\(_{\alpha,1.2}\) and Ca\(_{\alpha,1.3}\) L-type Ca\(^{2+}\) channels (LTCCs) are believed to underlie Ca\(^{2+}\) currents in brain, pancreatic β cells, and the cardiovascular system. In the CNS, neuronal LTCCs control excitation-transcription coupling and neuronal plasticity. However, the pharmacotherapeutic implications of CNS LTCC modulation are difficult to study because LTCC modulators cause cardiovascular (activators and blockers) and neurotoxic (activators) effects. We selectively eliminated high dihydropyridine (DHP) sensitivity from Ca\(_{\alpha,1.2}\) \(\alpha_1\) subunits (Ca\(_{\alpha,1.2DHP-/-}\)) without affecting function and expression. This allowed separation of the DHP effects of Ca\(_{\alpha,1.2}\) from those of Ca\(_{\alpha,1.3}\) and other LTCCs. DHP effects on pancreatic β cell LTCC currents, insulin secretion, cardiac inotropy, and arterial smooth muscle contractility were lost in Ca\(_{\alpha,1.2DHP-/-}\) mice, which rules out a direct role of Ca\(_{\alpha,1.3}\) for these physiological processes. Using Ca\(_{\alpha,1.2DHP-/-}\) mice, we established DHPs as mood-modifying agents: LTCC activator–induced neurotoxicity was abolished and disclosed a depression-like behavioral effect without affecting spontaneous locomotor activity. LTCC activator BayK 8644 (BayK) activated only a specific set of brain areas. In the ventral striatum, BayK-induced release of glutamate and 5-HT, but not dopamine and noradrenaline, was abolished. This animal model provides a useful tool to elucidate whether Ca\(_{\alpha,1.3}\)-selective channel modulation represents a novel pharmacological approach to modify CNS function without major peripheral effects.

Ca\(^{2+}\) influx through L-type calcium channels (LTCCs) is an important modulator of neuronal excitability (1, 2, 3). LTCC activation increases intracellular free Ca\(^{2+}\) concentrations, which may eventually lead to activation of the transcription of genes associated with long-term changes in synaptic plasticity (for review see ref. 3). The coupling of neuronal excitation and transcription may critically depend on the activity of the dihydropyridine-sensitive (DHP-sensitive) LTCCs Ca\(_{\alpha,1.2}\) and Ca\(_{\alpha,1.3}\). Thus, these two channels represent very interesting and potentially important therapeutic targets in the CNS. Ca\(_{\alpha,1.2}\) and Ca\(_{\alpha,1.3}\) channel blockers, such as nifedipine, ameliorate age-related working memory deficits in rodents (4), have antidepressant-like actions (5), and can affect fear memory (6). On the other hand, LTCC activators, such as BayK 8644 (BayK), stimulate neurotransmitter release in vitro (7), but also induce a severe dystonic neurobehavioral syndrome in rodents, including self-biting (8). Ca\(_{\alpha,1.2}\) and Ca\(_{\alpha,1.3}\) have a broad and overlapping expression profile in the mammalian neuronal system, where Ca\(_{\alpha,1.2}\) appears to be more prominent than Ca\(_{\alpha,1.3}\) (1).

Ca\(_{\alpha,1.2}\) and Ca\(_{\alpha,1.3}\) have also been detected in many non-neuronal tissues, e.g., insulin-secretory β cells (9, 10), vascular smooth muscle cells (11), and heart atria (12). Therefore, it remains unclear whether the reported CNS effects of LTCC blockers are due to a direct or indirect action since they also may cause cardiodepression and vasodilation (13). Furthermore, the contribution of Ca\(_{\alpha,1.2}\) and/or Ca\(_{\alpha,1.3}\) channels to DHP effects is unknown, because presently available LTCC activators and blockers cannot distinguish between the two channel isoforms.

To allow us to address the relative roles of Ca\(_{\alpha,1.2}\) and Ca\(_{\alpha,1.3}\) in greater detail, we have created a mouse model in which the high DHP sensitivity of Ca\(_{\alpha,1.2}\) \(\alpha_1\) subunits was eliminated by replacement of Thr1066 in helix IIIS5 with a tyrosine residue (14). This essentially eliminates the contribution of this channel type to DHP effects. We used this mouse model to directly determine the contribution of Ca\(_{\alpha,1.3}\) for pancreatic β cell Ca\(^{2+}\) currents and insulin secretion as well as cardiac and arterial smooth muscle function. We were also able to determine its contribution for high-affinity (+)\({\text{[H]}}\)risadipine binding in brain and heart, BayK-indcued dystonia, BayK effects on regional brain neurotransmitter efflux, and mood-related behavioral DHP effects. Our data indicate that Ca\(_{\alpha,1.3}\)-selective activators can cause distinct alterations in CNS function without affecting LTCCs in the cardiovascular system and pancreatic β cells.

Nonstandard abbreviations used: BayK 8644 (BayK); bed nucleus of the stria terminals (BNST); dihydropyridine (DHP); inward Ba\(^{2+}\) current (\(I_{\text{Ba}}\)); inward Ca\(^{2+}\) current (\(I_{\text{Ca}}\)); L-type Ca\(^{2+}\) channel (LTCC); paraventricular hypothalamic nucleus (PVN); voltage for half-maximal activation (\(V_{\text{act}}\)); voltage for half-maximal inactivation (\(V_{\text{inact}}\)).

Conflict of interest: The authors have declared that no conflict of interest exists.

Citation for this article: J Clin Invest. 2004;113:1430-1439 (2004).

doi:10.1172/JCI200420208.
Methods

All animal experiments were approved by the Austrian Bundesministerium für Bildung, Wissenschaft, und Kultur and the local ethical committee at Lund University.

Generation of Ca,1.2DHP–/– mice. Mouse genomic DNA clones (M24166Q3 and B17149Q3) containing IilSS (exon 24) encoding regions of the Ca,1.2 α1 subunit were isolated from a LAWRIST7 129 mouse strain genomic library (Deutsches Ressourcenzentrum für Genomforschung GmbH, Berlin, Germany). The targeting construct consisted of a 10.3-kb genomic Csp45I-SpeI fragment including the T1066Y mutation (exon 24), a neoymycin resistance gene (neo), and the plasmid pBLYESSCRIPT (Stratagene, La Jolla, California, USA). T1066Y was generated by overlap-extension PCR (15). The neo gene, flanked by two loxP elements in parallel orientation and driven by the phosphoglycerate kinase promoter, was inserted into the intron sequence 280-bp upstream of exon 24 by ligating into XhoI and SmaI recognition sites that were generated by overlap extension. The NotI-linearized targeting construct was electroporated into 129 (R1) ES cells, which were subjected to selection by genetin (G418; Invitrogen Corp., San Diego, California, USA). Two hundred and ninety-six transfected G418-resistant ES cell colonies were screened for homologous recombination events by Southern blot analysis of BamHI-digested genomic DNA with a 5T external probe (350 bp). The PCR-generated probe detected a 7.5-kb band for the WT and a 9.5-kb band for the mutated allele. The T1066Y mutation was detected by PCR using primers amplifying either the WT (fragment size 841 bp) or the mutated allele fragment (616 bp).

From 16 positive ES cell colonies, two were selected for blastocyst injection to create chimeric mice. Transmission of the mutant Ca,1.2 α1 allele from both clones was confirmed by PCR and Southern blot analysis of genomic DNA. The positive F1 progeny were crossed with C57BL/6J mice, and their F2 heterozygous offspring were genotyped by PCR, yielding fragment sizes of 390 bp and 475 bp for the WT (C57BL/6J) and a 9.5-kb band for the mutated allele. Heterozygous offspring were backcrossed into C57BL/6J mice for at least five generations by repetitive backcrossing to C57BL/6J mice. The heterozygous and WT littermates were used for all experiments.

Neurotransmitter efflux. Neurotransmitter efflux in the ventral striatum was determined using a micro push-pull superfusion technique. WT or Ca,1.2DHP–/– mice were anesthetized with urethane (1.2 g/kg) and mounted in a stereotaxic frame (Trent Wells, South Gate, California, USA), and the push-pull cannula (outer diameter, 0.5 mm; inner diameter, 0.3 mm) was inserted unilaterally into the ventral striatum. The coordinates from bregma (19) were as follows (in mm): AP, +1.0; L, 1.2; V, –4.5. The ventral striatum was superfused with artificial CSF (in mM: NaCl, 140; KCl, 3; glucose, 3; CaCl2, 1.5; MgCl2, 1; NaHOPO4, 1; and Na2HPO4, 1; pH 7.2) using a CMA 100 microinjection pump (CMA Microdialysis AB, Solna, Sweden) at a rate of 14 μl/min. Steady-state levels of neurotransmitters were reached 80 minutes after starting the superfusion. Four samples (20-minute sampling time each) were collected thereafter to determine basal extracellular neurotransmitter levels. Subsequently, BayK was applied either locally (0.1 μM added to the CSF) or systemically (2 mg/kg given intraperitoneally), and samples were collected for an additional 60 minutes and 100 minutes, respectively. Samples were immediately stored at –80°C and neurotransmitter concentrations were determined as described (20–22). At the end of the experiment mice were sacrificed with an overdose of sodium pentobarbital, the brain was removed, and the localization of the cannula was verified histologically in 50-μm sections.

Accelerating rotarod test. The accelerating rotarod test was performed as described (23). On days 1–3, mice were given three practice trials separated by 1 minute. On days 4 and 5, mice were given two practice trials, and then a third trial was carried out 20 minutes after injection of vehicle to habituate the mice to the injection procedures. On day 6, mice were given two practice trials, followed by injection with either vehicle or BayK. Twenty minutes after the injection, the test trial was performed.

Open-field test. The open field consisted of a plastic box (41 × 41 × 41 cm) equipped with an automated activity monitoring system (TruScan; Coulbourn Instruments, Allentown, Pennsylvania, USA). Illumination at floor level was 500 lux. The test was carried out as described (24). Center entries, center time, center distance, vertical plane entries (rearing), and total distance traveled were recorded.

Forced-swim test. The forced-swim test based on the test established by Porsolt et al. in 1977 was performed as described elsewhere (24). Mice were individually placed in a glass cylinder (diameter, 11.5 cm; height, 24 cm) containing 15 cm fresh tap water maintained at 24–25°C. Their activity was videotaped over a period of 6 minutes. The duration of immobility throughout the final 4 minutes of the test was assessed. Four mg/kg BayK (or vehicle) was injected intraperitoneally 20 minutes before the test. Twenty-five mg/kg nifedipine (or vehicle) was administered 23 hours after a 15-minute pretest in the absence of drugs and 60 minutes prior to the 6-minute testing period.

Measurement of [3H] in neonatal cardiomyocytes. Inward Ba2+ currents ([3H]I) through voltage-gated Ca2+ channels were recorded at 22–25°C using the patch-clamp technique as previously described (25). Patch pipettes (resistance, 1–4 MΩ) were made from borosilicate glass and filled with pipette solution containing (in mM): CsCl, 60; CsOH, 60; aspartate, 60; MgCl2, 2; HEPES, 10; and EGTA, 10 (adjusted to pH 7.25 with CsOH). The bath solution contained (in mM): BaCl2, 10; N-methyl-d-glucamine, 190; HEPES, 10; glucose, 20; 4-aminopyridine, 4; tetraethylammonium chloride, 27; and MgCl2, 3; buffered to pH 7.4 with methanesulfonic acid. All data were digitized and filtered at 2 kHz with a four-pole Bessel filter. Leak currents were subtracted digitally (using average values of scaled leak-
age currents elicited by a 10-mV hyperpolarizing pulse) or electronically by means of an analogue circuit.

Pancreatic islet experiments. Pancreatic islets were isolated and insulin release was measured in static batch incubations by RIA, as described (26). Voltage-gated Ca$^{2+}$ currents were recorded using the perforated-patch whole-cell approach. To block a prominent outward delayed rectifying K$^{+}$ current and to resolve the inward Ca$^{2+}$ current component, the extracellular buffer was supplemented with 20 mM tetraethylammonium chloride (TEA-Cl), and the pipette solution was made using Cs$^{+}$. The extracellular Ca$^{2+}$ concentration was 2.6 mM. Effects of isradipine and BayK were determined in the steady state.

DHP effects on contractility of isolated cardiac muscle and mouse aortic rings. Isometric contractions (resting tension of 0.5 g) of mouse aortic rings were recorded as described (27). After equilibration (>60 min) they were repeatedly stimulated with phenylephrine (1 μM) until reproducible contractile responses were obtained. The effect of isradipine was tested on rings that were exposed to 90 mM KCl and allowed to contract until the responses reached a plateau. Parallel experiments were done to exclude an effect of the solvent (DMSO). BayK effects were measured on rings partially contracted with 15 mM KCl.

Langendorff isolated perfused hearts were mounted as described (28) with minor modifications. Briefly, the hearts from previously heparinized mice were quickly explanted and perfused through the aorta at constant flow (2 ml/min), and the atria were removed. The perfusion solution (containing in mM: NaCl, 118; KCl, 4.7; CaCl$_2$, 2.5; MgSO$_4$, 1.2; NaCO$_3$, 25; KH$_2$PO$_4$, 1.2; glucose, 11.1; and Na pyruvate, 2) was bubbled with a 95% O$_2$, 5% CO$_2$ gas mixture (pH 7.4 ± 0.01) at 37°C. After a stabilization period of 30 minutes, the hearts were electrically stimulated at a frequency of 5.5 Hz by placing a platinum electrode on the upper part of the right ventricle. Contractility was measured by placing in the left ventricle a steel cannula connected to a pressure transducer (Ugo Basile, Comerio, Italy). The hearts were allowed to stabilize for at least 60 minutes before drug exposure.

Telemetric recordings. Telemetric recordings were carried out as described previously (29). Control ECGs were recorded continuously (sampling rate, 500 Hz) from freely moving mice at least 2 days after transmitter implantation. Mice were then injected with 1 mg/kg of atropine followed by 20 mg/kg of propranolol to eliminate sympathetic and parasympathetic control of heart rate. Increasing concentrations of DHPs were then injected and heart rate was analyzed during 60-minute periods 20 minutes after drug application.

Immunoblotting. Immunoblots of brain and heart microsomal membranes were carried out as described (29). Thirty micrograms of membrane protein separated on 5% SDS-polyacrylamide gels and immunoblots probed with affinity-purified rabbit antibody anti-Ca$_{1.2}$α1.2 LTCCs is absent in Ca$_{1.2}$DHP–/– mice. The gene-targeting strategy is illustrated in Figure 1A. Successful gene targeting and the generation of homozygous mutants were confirmed by Southern blot analysis of genomic DNA (Figure 1B). In immunoblots, no effects of the mutation on Ca$_{1.2}$α1 expression density were detected in brain and heart membranes (Figure 1C) and after heterologous expression in tsA-201 cells (n = 2, not shown).

Results

High DHP sensitivity of Ca$_{1.2}$ LTCCs is absent in Ca$_{1.2}$DHP–/– mice. The gene-targeting strategy is illustrated in Figure 1A. Successful gene targeting and the generation of homozygous mutants were confirmed by Southern blot analysis of genomic DNA (Figure 1B). In immunoblots, no effects of the mutation on Ca$_{1.2}$α1 expression density were detected in brain and heart membranes (Figure 1C) and after heterologous expression in tsA-201 cells (n = 2, not shown).
No detectable change of the expression pattern of mRNA of Ca\(_{\alpha,1,2}\) or other Ca\(^{2+}\) channel α1 subunits was found in the mutant mouse brains (n = 3, not shown). Selective labeling of LTCCs with 0.9–1.4 nM (+)-\(^3\)H]isradipine revealed a decrease of specific DHP binding to 55% ± 2% and 15% ± 2% (n = 8) of WT values in brain membranes of heterozygous and homozygous mutants, respectively (Figure 1D). No significant binding remained in the heart membranes of homozygous mutants (4% ± 5%; n = 6) (Figure 1D). The reduced isradipine binding in brain could be attributed to a lower maximal binding capacity, which was 55% of WT values (426 ± 21 fmol/mg of protein) in heterozygous (237 ± 17 fmol/mg) and 20% in homozygous mutants (88 ± 9 fmol/mg; n = 7). Residual binding was of high affinity (K\(_D\) in pM: WT, 78 ± 17; heterozygous, 96 ± 27; homozygous, 175 ± 42; n = 7) and stimulated by (+)-tetrandrine (to 318% ± 19% of control binding = 6) (Figure 1D). The reduced isradipine binding is associated with Ca\(_{\alpha,1,3}\) in brain membranes of Ca\(_{\alpha,1,2}\)-deficient mice (30), and no other neuronal LTCC α1 subunits are known to exist in the brain, residual binding must be associated with Ca\(_{\alpha,1,3}\) LTCCs.

Next we confirmed that the mutation eliminates high DHP sensitivity of Ca\(_{\alpha,1,2}\) without changing its functional properties. We therefore recorded whole-cell currents (I\(_{\text{Ba}}\)) from neonatal ventricular cardiomyocytes, which are believed to exclusively express Ca\(_{\alpha,1,2}\) (12). The biophysical characteristics of I\(_{\text{Ba}}\) measured from cultured neonatal Ca\(_{\alpha,1,2}\)DHP–/– cardiomyocytes were indistinguishable from those of WT cardiomyocytes. Inactivation during depolarizations from –80 mV to different test potentials were obtained by fitting the inactivation data to a monoexponential function. No statistically significant difference was found between myocytes from WT and mutant mice.

![Image](213x496 to 542x702)

**Figure 2**

Biophysical properties of Ca\(_{\alpha,1,2}\) currents in Ca\(_{\alpha,1,2}\)DHP–/– cardiomyocytes. (A and B) I\(_{\text{Ba}}\) through LTCCs was recorded from cardiomyocytes isolated from neonatal WT or homozygous mutant mice. Currents recorded with no treatment (CO) or 3 minutes after perfusion with 1 μM isradipine (ISR 1) or 0.1 μM BayK (BayK 0.1) and 1 μM BayK (BayK 1). Holding potential, –80 mV; test potential, +10 mV. One of more than three experiments yielding similar results is shown. (C) Concentration-dependent isradipine inhibition. Each curve was constructed from experiments (n = 5) as described in A. D) Time constants (τ) for current inactivation during 300-ms depolarizations from –80 mV to different test potentials were obtained by fitting the inactivation data to a monoexponential function. No statistically significant difference was found between myocytes from WT and mutant mice.

No detectable change of the expression pattern of mRNA of Ca\(_{\alpha,1,2}\) or other Ca\(^{2+}\) channel α1 subunits was found in the mutant mouse brains (n = 3, not shown). Selective labeling of LTCCs with 0.9–1.4 nM (+)-\(^3\)H]isradipine revealed a decrease of specific DHP binding to 55% ± 2% and 15% ± 2% (n = 8) of WT values in brain membranes of heterozygous and homozygous mutants, respectively (Figure 1D). No significant binding remained in the heart membranes of homozygous mutants (4% ± 5%; n = 6) (Figure 1D). The reduced isradipine binding in brain could be attributed to a lower maximal binding capacity, which was 55% of WT values (426 ± 21 fmol/mg of protein) in heterozygous (237 ± 17 fmol/mg) and 20% in homozygous mutants (88 ± 9 fmol/mg; n = 7). Residual binding was of high affinity (K\(_D\) in pM: WT, 78 ± 17; heterozygous, 96 ± 27; homozygous, 175 ± 42; n = 7) and stimulated by (+)-tetrandrine (to 318% ± 19% of control binding = 6) (Figure 1D). The reduced isradipine binding is associated with Ca\(_{\alpha,1,3}\) in brain membranes of Ca\(_{\alpha,1,2}\)-deficient mice (30), and no other neuronal LTCC α1 subunits are known to exist in the brain, residual binding must be associated with Ca\(_{\alpha,1,3}\) LTCCs.

Next we confirmed that the mutation eliminates high DHP sensitivity of Ca\(_{\alpha,1,2}\) without changing its functional properties. We therefore recorded whole-cell currents (I\(_{\text{Ba}}\)) from neonatal ventricular cardiomyocytes, which are believed to exclusively express Ca\(_{\alpha,1,2}\) (12). The biophysical characteristics of I\(_{\text{Ba}}\) measured from cultured neonatal Ca\(_{\alpha,1,2}\)DHP–/– cardiomyocytes were indistinguishable from those of WT cardiomyocytes. Inactivation during depolarizations from –80 mV to different test potentials were obtained by fitting the inactivation data to a monoexponential function. No statistically significant difference was found between myocytes from WT and mutant mice.

To 581% ± 13% of control I\(_{\text{Ba}}\) at 1 μM; n = 4) was completely absent in cardiomyocytes isolated from mutant mice (87% ± 3% of control; n = 3). Given the known BayK sensitivity of all known L-type channels (12,31), we conclude from these data that Ca\(_{\alpha,1,2}\)DHP–/– mice should allow discrimination of DHP effects on Ca\(_{\alpha,1,2}\) from other LTCCs in vitro and in vivo. We previously found that isradipine blocks recombinant Ca\(_{\alpha,1,3}\) channels under similar experimental conditions with an IC\(_{50}\) of 300 nM (15). Therefore the in vivo mutation of Thr1066 converted the about tenfold selectivity of isradipine for Ca\(_{\alpha,1,2}\) (15) to an at least tenfold selectivity for Ca\(_{\alpha,1,3}\), making this DHP a selective Ca\(_{\alpha,1,3}\) channel blocker in Ca\(_{\alpha,1,2}\)DHP–/– mice.

**Lack of Ca\(_{\alpha,1,3}\) LTCC contribution to cardiac and arterial smooth muscle contraction.** As illustrated in Figure 3, smooth muscle relaxant and negative inotropic actions of isradipine were completely absent in Ca\(_{\alpha,1,2}\)DHP–/– mice. As expected, the mutation did not affect verapamil sensitivity. Verapamil at a concentration of 1 μM caused an almost complete block of cardiac contraction in WT (not shown) and mutant mouse hearts (11.9% ± 3.5% of control, n = 4). Surprisingly, we also found that isradipine was even able to slightly enhance cardiac inotropy. This revealed an unexpected pharmacological action of isradipine, which is normally masked by the loss of contractility through Ca\(_{\alpha,1,2}\) LTCC block. We telemetrically recorded DHP effects on intrinsic sinoatrial node activity in mice pretreated with atropine and propranolol to eliminate autonomic control. BayK caused only a slight increase in heart rate in WT mice that was absent in mutants (Figure 3E). The bradycardic effect of isradipine in WT mice (Figure 3E) was not affected by the mutation. These data significantly extend our previous in vitro data demonstrating the importance of Ca\(_{\alpha,1,3}\) for sinoatrial node function (29). We show that cardiac and smooth muscle inotropy are controlled by Ca\(_{\alpha,1,2}\) but not Ca\(_{\alpha,1,3}\) activity, despite Ca\(_{\alpha,1,3}\) mRNA expression in aortic smooth muscle (11).

Ca\(_{\alpha,1,3}\) activity does not contribute to pancreatic β cell Ca\(^{2+}\) currents and insulin secretion. Our animal model also allowed us to address the still-controversial issue concerning the contribution of Ca\(_{\alpha,1,3}\) to pancreatic β cell Ca\(^{2+}\) currents and insulin secretion. Three different mouse models as well as biochemical data have provided evidence both for a role of Ca\(_{\alpha,1,3}\) in β cells (9, 10) and against it (26, 32). In our mice, DHP-sensitive Ca\(^{2+}\) current components in β cells
Fig. 1. Neuronal activation to BayK- and isradipine-induced contractions. (A) Representative experiment illustrating drug effects on cardiac ventricle contractility in Langendorff hearts from WT or homozygous mutant Ca\(^+\)v1.2DHP–/– mice. Electrically stimulated hearts were superfused with isradipine (ISR) at concentrations of 0.5 μM, 1 μM, 2.5 μM, and 10 μM or with 1 μM verapamil (VER) (see Results). (B) Percent inhibition of contractility in six WT (black bars) and seven mutant (white bar) mice. Statistical significance; *P < 0.001 (compared with Mut; one-way ANOVA followed by the Bonferroni multiple-comparison test); †P < 0.05 (compared with contractility in the absence of isradipine, 100%; one-sample Student t test). Data from 1 μM and 2.5 μM isradipine (mean ± range of n = 2) were not included in the significance analysis. (C) BayK-induced (1 μM) contraction of aortic smooth muscle rings from WT (n = 10) or mutant mice (n = 10). *P < 0.001 (Ca\(^+\)v1.2DHP–/– vs. WT; unpaired Student’s t test). (D) Effect of isradipine (1 μM) on aortic smooth muscle rings isolated from WT (n = 10) or mutant mice (n = 10) precontracted with 90 mM KCl. 1P < 0.001 (Ca\(^+\)v1.2DHP–/– vs. WT; unpaired Student’s t test). (E) Telemetric recordings (see Methods). Effects of increasing doses of BayK (left) and isradipine (right) on sinoatrial activity in vivo. R, resting, no drug. At, injection with atropine (1 mg/kg); P, injection with atropine followed by propranolol (20 mg/kg) before DHP application. Data presented as mean SEM of 4–52 experiments. *P < 0.05; **P < 0.01 for heart rate inhibition in the presence of drug (one-sample Student’s t test).

BayK-induced neurotoxicity is Ca\(^+\)v1.2 dependent. In WT mice, injection of low doses (2 mg/kg) of BayK (but not of vehicle) produced mild to moderate motor disability (motor impairment score 2–3; see ref. 8), with limited ambulation, hypokinesia, and frequent abnormal postures (n = 5) (Figure 5). Higher doses (4–6 mg/kg, n = 3) led to severe behavioral impairment (score >3), including self-biting (not shown) (34). In contrast, no changes in motor function or abnormal behaviors were noticed in homozygous mutants given 2 mg/kg BayK subcutaneously (n = 5) (Figure 5). BayK at 4 mg/kg did not affect motor function in the rotarod treadmill test (time spent on rotarod: vehicle treatment, 263.8 ± 14.1 s, n = 9; BayK treatment, 254.3 ± 14.7 s, n = 10; P = 0.671 by Mann-Whitney U test). To test whether Ca\(^+\)v1.2DHP–/– mice tolerate even higher doses of BayK, mutants were injected with up to 10 mg/kg of BayK. Mice receiving 5 or 10 mg/kg subcutaneously showed no motor symptoms or abnormal behavior within 60 minutes (n > 8). This demonstrates that Ca\(^+\)v1.2 activation is required for BayK-induced neurotoxicity. The absence of toxic symptoms in Ca\(^+\)v1.2DHP–/– mice allowed us to further investigate the consequences of selective Ca\(^+\)v1.3 activation on brain function.

Ca\(^+\)v1.3 selectively contributes to BayK-induced Fos expression. Ca\(^+\)v1.2 and Ca\(^+\)v1.3 LTCCs have a broad and overlapping expression pattern in the brain (1). The selective Ca\(^+\)v1.3 activation in Ca\(^+\)v1.2DHP–/– mice by BayK allowed us to identify neurons activated after selective Ca\(^+\)v1.3 stimulation in vivo. Fos expression was used as a marker of neuronal activation (see ref. 18 and references therein).
Vehicle injection elicited only low levels of Fos-positive cells, which were similar in number in WT (not shown) and mutant mice (Figure 6A). BayK (2 mg/kg) evoked widespread Fos expression in brains of WT mice in most of the 65 brain areas investigated (n = 5). Induction was absent in only a few brain areas such as the ventral postomemorial thalamic nucleus or the mammillary nucleus (n = 3; not shown). The most prominent increase in Fos expression was observed in the infralimbic cortex (BayK, 32.7 ± 1.8 cells/0.01 mm²; vehicle, 8.8 ± 1.7 cells/0.01 mm²; P < 0.02), the pyramidal cell layer of the hippocampus (BayK, 33.3 ± 1.8 cells/0.01 mm²; vehicle, 9.4 ± 1.3 cells/0.01 mm²; P < 0.02), the retrospenial granular cortex (BayK, 49.0 ± 1.5 cells/0.01 mm²; vehicle, 15.0 ± 1.8 cells/0.01 mm²; P < 0.02), and the locus coeruleus (BayK, 40.3 ± 2.4 cells/0.01 mm²; vehicle, 3.6 ± 0.7 cells/0.01 mm²; P < 0.01). Figure 6A illustrates BayK-induced Fos expression in the cingulate cortex and dorsal striatum in WT animals. In the mutant mice (n = 8), BayK-induced Fos expression was greatly attenuated in most brain regions compared with WT mice. As illustrated in Figure 6A, BayK failed to enhance the number of Fos-positive cells in the cingulate cortex (BayK, 22.5 ± 1.3 cells/0.01 mm²; vehicle, 21.0 ± 1.5 cells/0.01 mm²; P = 0.45) and dorsal striatum. In striatal sections from mutants, no expression above the levels of vehicle controls was found in the caudate putamen (BayK, 8.6 ± 0.7 cells/0.01 mm²; vehicle, 7.9 ± 1.1 cells/0.01 mm²; P = 0.8). However, significant BayK-induced Fos expression in Ca,1.2DHP/–/– mice was detectable in a limited subset of brain regions. Figure 6 illustrates Fos induction in mutants by BayK in the nucleus accumbens (BayK, 19.0 ± 1.4 cells/0.01 mm²; vehicle, 12.9 ± 2.9 cells/0.01 mm²; P < 0.01; Figure 6B) and the bed nucleus of the stria terminalis (BNST): BayK, 27.5 ± 1.8; vehicle, 6.3 ± 0.9; P < 0.001; Figure 6C). High levels of BayK-induced Fos expression were also observed in the paraventricular hypothalamic nucleus (PVN) (BayK, 45.6 ± 2.6 cells/0.01 mm²; vehicle, 8.7 ± 2.1 cells/0.01 mm²) and the central amygdaloid nucleus (BayK, 22.6 ± 1.8 cells/0.01 mm²; vehicle, 6.5 ± 0.9 cells/0.01 mm²). These experiments revealed that most of the high BayK-induced neuronal activation in mouse brain in vivo is mediated by Ca,1.2 but that selective Ca,1.3 activation still stimulates Fos expression in a distinct set of brain areas without causing obvious toxic reactions.

Based on pharmacokinetic data of various DHPs in mice, micromolar BayK concentrations must be achieved in brain after intraperitoneal application of 4 mg/kg (see legend to Figure 6D). At these concentrations BayK stimulates the activity of Ca,1.3 but not mutant Ca,1.2 LTCCs (Figure 6D).

Ca,1.3 activation contributes to BayK-stimulated neurotransmitter efflux in the ventral striatum. We next determined to what extent Ca,1.3 activation alone contributes to the known BayK-induced neurotransmitter overflow in the CNS in vivo. The ventral striatum was selected to measure neurotransmitter efflux. In this region, efflux of various monoamines and amino acids has been successfully determined (35). LTCC isoforms are also

Figure 4
DHP sensitivity of L-type I_{Ca} and insulin secretion in isolated pancreatic β-cells. (A) Whole-cell I_{Ca,L} during 100-ms depolarizations from –70 mV to 0 mV in single pancreatic β-cells isolated from WT and Ca,1.2DHP/–/– mice. The L-type channel activator BayK (2 μM) or the L-type channel inhibitor isradipine (2 μM) were added as indicated. (B) Charge-voltage (Q-V) relations recorded in β-cells (100-ms depolarizations to voltages between –60 mV and +20 mV) from WT and Ca,1.2DHP/–/– mice in the absence (open circles) or presence of BayK (filled diamonds) or isradipine (filled circles). Data represent mean ± SEM (n = 4–5). In WT β-cells the effects of isradipine and BayK were significant (P < 0.05; Student t test) at depolarizations beyond –30 mV and –10 mV, respectively. (C) Insulin secretion measured in isolated islets from control (black bars) and Ca,1.2DHP/–/– (white bars) mice in the absence and presence of glucose and DHPs as indicated. Data are mean ± SEM (n = 6). ***P < 0.001 (Student t test) for isradipine inhibition; * P < 0.05 for BayK stimulation (WT vs. Ca,1.2DHP/–/–).

Figure 5
BayK-induced changes in motor disability grade in WT and Ca,1.2DHP/–/– mice. WT (closed circles, n = 5) or Ca,1.2DHP/–/– (open circles, n = 5) mice were injected subcutaneously with 2 mg/kg BayK and placed singly in 20 cm ×30 cm clear plastic cages. They were then observed for 1 minute each at 5-minute intervals. Motor disability was rated at each timepoint on a 4-point scale according to Jinnah et al. (8). No motor symptoms or abnormal behavior were observed after injection of vehicle alone.

BayK-induced Fos expression in WT and Ca,1.2DHP+/- mice. Mice were injected with vehicle or BayK solution (WT, 2 mg/kg; mutants, 4 mg/kg) and Fos expression was quantified by immunohistochemistry as described in Methods. (A) DS, dorsal striatum; Ci, cingulate cortex; LV, lateral ventricle. Magnification, ×40. Inset shows higher magnification of boxed areas. (B) Fos expression after BayK (right) or vehicle (left) application in the nucleus accumbens. ac, anterior commissure, anterior. Magnification, ×100. Inset magnification, ×800. (C) Fos expression after BayK (right) or vehicle (left) application in the BNST. Magnification, ×100. Inset magnification, ×800 (boxed area in the lateral division). acp, anterior commissure, posterior. (D) Stimulation of I Na through Ca,1.2 (Ca,1.2WT), mutant Ca,1.2 (Ca,1.2MUT), and Ca,1.3 by BayK after heterologous expression under identical conditions in tsA-201 cells as described (15). Based on DHP pharmacokinetic data in mice (53), we calculated BayK concentration in brain to reach concentrations between ~7 μM (peak concentration) and ~1 μM (after three elimination half-lives). All data were significantly different from 1 (control before drug application) (P < 0.05; one-sample Student’s t test) except Ca,1.2MUT, 1 μM and 5 μM BayK).

expressed and LTCC activators induce CREB phosphorylation and c-Fos expression in this area (see above) (36).

No difference in basal efflux of dopamine, noradrenaline, 5-HT, or glutamate was found between WT mice and homozygous mutants (n = 5–6, see legend to Figure 7). In WT mice, superfusion with 0.1 μM BayK increased the efflux of all studied neurotransmitters (Figure 7). BayK-induced neurotransmitter efflux was similar in WT and mutant mice for noradrenaline (fold increase: WT, 1.34 ± 0.05, n = 6; Ca,1.2DHP+/-, 1.39 ± 0.12, n = 5, P = 0.6) but was significantly attenuated in mutant mice for dopamine (fold increase: WT, 2.94 ± 0.35, n = 6; Ca,1.2DHP+/-, 1.53 ± 0.10, n = 5, P < 0.01) and absent for glutamate (fold increase: WT, 14.35 ± 7.26, n = 6; Ca,1.2DHP+/-, 1.76 ± 1.06, n = 5) and 5-HT efflux (fold increase: WT, 3.12 ± 1.27, n = 6; Ca,1.2DHP+/-, 1.05 ± 0.07; n = 4; all by Mann-Whitney U test) (Figure 7). Systemic application of BayK (2 mg/kg given intraperitoneally) in mutants also significantly enhanced noradrenaline (fold increase: 1.68 ± 0.21; n = 5, P = 0.03) and dopamine efflux (fold increase: 1.30 ± 0.08; n = 5, P = 0.04) but not glutamate (fold increase: 0.79 ± 0.26; n = 4, P > 0.7) and 5-HT (fold increase: 0.98 ± 0.35; n = 3, P = 1.00) within 10–20 minutes after drug application, suggesting that the stimulatory effect of BayK is not limited to intrastriatal application. We conclude that in the ventral striatum, BayK-induced noradrenaline efflux was mediated only through Ca,1.3, whereas enhanced glutamate and 5-HT efflux required Ca,1.2. Dopamine efflux stimulation was mediated by both isoforms.

Ca,1.3 activation induces depression-like behavioral effects. Some of the regions activated by BayK in Ca,1.2DHP+/- mice, such as the PVN, BNST, ventral striatum, or amygdala, are part of anatomical circuits implicated in the processing of depression-related behavior (37). In addition, antidepressant-like effects of DHP LTCC blockers have been reported in rodents (see Introduction). The preferred test for demonstrating antidepressant-like effects in rodents is the Porsolt behavioral despair (forced-swim) test (38). In this test the extent of immobility (passive floating) displayed by rodents during a swim challenge is taken as a measure of depression-like behavior, which is decreased by antidepressants. We therefore also used this experimental paradigm to investigate the role of Ca,1.3 for DHP effects on mood behavior. Administration of BayK to Ca,1.2DHP+/- mice caused a pronounced increase of immobility time (Figure 8C), indicating that BayK is able to induce a depression-like behavior in Ca,1.2DHP+/- mice. The neurotoxic effects of BayK precluded parallel behavioral tests in WT mice. BayK did not affect motor function in the rotarod treadmill test (see above) or spontaneous motor activity in the open-field test (distance traveled in 10 minutes: vehicle, 1,727.6 ± 87.7 mm, n = 7; BayK, 1,940.3 ± 117.9 mm, P = 0.142; center entries: vehicle, 13.3 ± 1.8, n = 7; BayK, 13.9 ± 3.0, n = 7, P = 1.00 by Mann-Whit-
Ca,1.1 α1 subunits (restricted expression in adult skeletal muscle, ref. 42), are not expressed at significant levels. The selectivity of DHP channel blocker action in Ca,1.2DHP−/− mice was clearly apparent from functional studies: 1 μM or 10 μM isradipine failed to reduce cardiac inotropy in isolated Langendorff hearts, caused no relaxation of vascular smooth muscle, and failed to inhibit pancreatic L-type currents in mutant mice at micromolar concentrations. An obvious advantage of Ca,1.2DHP−/− mice over Ca,1.2 α1−deficient mouse models is that compensatory expression of other Ca2+ channel subtypes cannot occur. A disadvantage is that, due to the remaining weak isradipine sensitivity of the mutant Ca,1.2 observed in Ca,1.2DHP−/− mice, any DHP antagonist effects observed in these mice need to be interpreted with caution as they could be mediated by weak block of the mutant Ca,1.2.

Peripheral actions of Ca,1.3-selective modulators. As LTCC activators typically cause a shift of voltage-dependent channel activation to more negative potentials and increase insulin secretion (15, 33), such drugs theoretically allow stimulation of insulin secretion in a glucose-sensitive manner (33). Our data clearly revealed that Ca,1.3 LTCCs are not suitable targets for insulin secretagogues because we found no evidence for the acute regulation of β cell Ca2+ currents or insulin secretion by DHPs in Ca,1.2DHP−/− mice. On the other hand, the absence of BayK effects on insulin secretion prevents adverse effects through insulin-induced metabolic disturbances, such as hypoglycemia, which

Figure 7
BayK-stimulated neurotransmitter efflux in the ventral striatum. Effect of intraatrial administration of BayK (0.1 μM) on neurotransmitter efflux in WT (filled circles) and Ca,1.2DHP−/− (open circles) mice. The bars indicate the application of 0.1 μM BayK. Data were normalized to the mean efflux rates in the two samples (control efflux) preceding drug application. Baseline efflux (fmol/min) in WT mice (n = 6) was: dopamine, 2.84 ± 0.44; noradrenaline, 0.77 ± 0.08; 5-HT, 1.10 ± 0.21; and glutamate, 974 ± 193. Baseline efflux (fmol/min) in Ca,1.2DHP−/− mice (n = 5) was: dopamine, 2.02 ± 0.27; noradrenaline, 0.58 ± 0.08; 5-HT, 0.96 ± 0.09; glutamate, 1,107 ± 161. Statistically significant differences from control efflux were determined using the Friedman test followed by the Wilcoxon signed-rank test (†P < 0.05). Differences between WT and mutant mice were calculated using the Mann-Whitney U test (‖P < 0.05).

Discussion
We report the successful generation of a novel mouse model that allows us to discriminate DHP effects on Ca,1.2 from other LTCC isoforms. As verified in ventricular myocytes, in vivo mutation of Thr1066 to tyrosine (T1066Y) completely removed BayK modulation of Ca,1.2. It also dramatically reduced DHP blocker sensitivity, rendering isradipine highly selective for Ca,1.3. This pharmacological phenotype was achieved without detectable alterations in channel gating or Ca,1.2 α1 expression. High-affinity DHP binding in brain was reduced to a level (20%) known to be associated with Ca,1.3 (1, 30). Any remaining BayK effects in nonretinal neurons must therefore be Ca,1.3-mediated. This must also hold true for most other tissues in which the other known LTCC isoforms, Ca,1.4 (restricted expression in retinal neurons; see refs. in ref. 41) and

Figure 8
Effect of nifedipine and BayK on immobility time in the forced-swim test. Immobility times determined during the last 4 minutes of a 6-minute test period after intraperitoneal injection of vehicle or DHPs. Due to its neurotoxicity, BayK was not tested in WT mice. The number of experiments is indicated in parentheses. (A and B) Immobility times of WT and Ca,1.2DHP−/− mice 60 minutes after application of nifedipine (NIF, 25 mg/kg) or vehicle (V). A pre-test was performed 24 hours before the test to increase immobility times. *P < 0.05 by Mann-Whitney U test. (C) Immobility times of Ca,1.2DHP−/− mice 20 minutes after application of BayK (4 mg/kg) or vehicle (V) without pre-test. **P < 0.01 by Mann-Whitney U test.
would complicate analysis of CNS effects. These data also resolve the conundrum regarding the molecular identity of the LTCC in β cells. Our data strongly suggest that in mouse β cells, contrary to previous reports (9, 10), Ca1.2 channels represent the most important conduit of Ca2+ entry involved in insulin secretion.

The bradycardiac effect of isradipine on intrinsic sinoatrial node activity observed in “pharmacologically denervated” hearts provides in vivo proof for our in vitro data demonstrating that Ca1.3 is the major LTCC controlling diastolic depolarization in resting animals (12, 29). Interestingly, BayK did not cause a significant increase in heart rate, not even at the high doses used in Ca1.2DHP−/− mice. This indicates that Ca1.3 channels stabilize normal heart rate (29) but that their selective pharmacological activation does not exert pronounced acceleration of intrinsic sinoatrial node automaticity in vivo.

We also show that DHP activator and blocker effects on the contractility of heart ventricular muscle and aortic smooth muscle are exclusively mediated by Ca1.2. The complete absence of BayK and isradipine effects on smooth muscle contraction is surprising because Ca1.3 α1 subunits are expressed in aortic muscle (11). In mammals, vasodilation induced by DHP channel blockers and vasoconstriction induced by BayK (13, 43) may alter cerebral blood flow, thus complicating the interpretation of CNS effects in vivo. The absence of these effects in Ca1.2DHP−/− mice provides us with a unique model to extend in vitro pharmacological research on neuronal LTCC function to the in vivo level.

CNS effects of Ca1.3-selective modulators. Another important prerequisite to analyze the consequences of selective Ca1.3 activation is the absence of neurotoxic effects. Although we found that 20% of the total LTCC activity in the CNS is associated with Ca1.3, all the neurotoxic symptoms observed after application of BayK in WT animals were completely absent even at very high doses (10 mg/kg given intraperitoneally). Dystonia and neurobehavioral abnormalities, like self-biting (34), therefore require enhanced Ca2+ influx through Ca1.2. This finding demonstrates a role of Ca1.2 hyperactivity in dystonic symptoms. It is also supported by the previous observation that upregulation of Ca1.2 expression in cerebellar Purkinje cells may be responsible for dystonic episodes in tottering mutant mice, which suffer from a complex neurological syndrome caused by mutations in a non-L-type Ca2+ channel (Ca2.1) (44). Our study should therefore prompt further analysis of the role of LTCC dysfunction in dystonic syndromes.

The absence of neurotoxicity also allowed us to demonstrate that Ca1.3-selective activation directly activates neuronal circuits. This could be shown both as a stimulation of neurotransmitter release as well as an increased Fos expression. Whereas BayK induced ubiquitous Fos expression in WT brains, the Fos response was greatly attenuated in Ca1.2DHP−/− brains. This suggests that this drug effect is mainly transduced by Ca1.2. BayK-induced Fos expression in Ca1.2DHP−/− mice was noted in only a few regions, including limbic and hypothalamic areas such as the BNST, nucleus accumbens, amygdala, and the PVN. The activation of this set of areas, implicated in functions such as the integration of emotion-related behavior, drug addiction, learning, and memory, should prompt further studies with Ca1.2DHP−/− mice to reveal the role of Ca1.3 in DHP effects on these brain functions (6, 45–47).

Elimination of Ca1.2 BayK sensitivity not only prevented neuronal activation of certain brain regions but also resulted in differential effects on neurotransmitter release in the ventral striatum. In anesthetized WT mice, spontaneous efflux of all four studied neurotransmitters (dopamine, noradrenaline, 5-HT, and glutamate) was significantly enhanced by intrastratial administration of BayK. Since no cell bodies of noradrenergic, dopaminergic, 5-HTergic, or glutamatergic neurons are present, these neurotransmitters are most likely released from projection neurons, originating for example from the ventral tegmental area (dopaminergic), cortical areas (glutamatergic), the dorsal raphe nucleus (5-HTergic), or locus coeruleus (noradrenergic). Therefore, locally applied BayK enhances neurotransmitter efflux either directly, by activation of presynaptic LTCCs (48, 49), or indirectly, by affecting striatal neuronal networks controlling neurotransmitter efflux from these afferents. In the ventral striatum, Ca1.2 and Ca1.3 contribute to a different extent to BayK-induced neurotransmitter efflux. Enhancement of extracellular glutamate and 5-HT was completely dependent on Ca1.2 channels. In contrast, the selective activation of Ca1.3 was sufficient to stimulate extraneuronal dopamine and noradrenaline activity. Thus, a specific neurotransmitter efflux pattern can be induced by acutely activating Ca1.3. It remains to be shown whether this can also be observed in other brain areas.

In Ca1.2DHP−/− mice, we can now show that this neuronal activation pattern resulting from selective Ca1.3 activation also affects brain function in a specific manner. BayK increased depression-like behavior without affecting locomotor activity, thus providing convincing evidence that Ca1.3 hyperactivity can alter mammalian mood-related behavior. We also confirmed that DHP channel blockers induce antidepressant-like effects, which we found to be Ca1.2 dependent. Based on our findings with BayK it seems likely that this is mainly due to modulation of neuronal LTCC activity rather than indirect cardiovascular effects.

It is assumed that many brain areas mediate the diverse symptoms of depression (50). Interestingly, BayK particularly increased Fos expression in subcortical brain areas thought to be abnormal in depression, including the nucleus accumbens, amygdala, and the PVN (50). Similarly, the BNST, which displayed enhanced BayK-induced Fos expression, has also been implicated in depression-like behavior (51). As only a limited number of regions are activated by BayK in Ca1.2DHP−/− brains, this animal model represents a useful and novel tool to further study neuronal circuits underlying depression-like behavior (37). Moreover, if the depression-like behavior induced by BayK is reversed by known antidepressant drugs, then Ca1.2DHP−/− mice could serve as a novel animal model for depression suitable for the in vivo screening of antidepressant drug actions.

Our finding of an acute depressant-like effect of BayK establishes a role for LTCC in mood behavior but does not suggest therapeutic benefit. However, final predictions about the pharmacotherapeutic potential of selective Ca1.3 activators must await experimental results from chronic treatment (weeks to months) with different doses of DHP channel activators. The differential effects on neurotransmitter release observed in Ca1.2DHP−/− brains could lead to adaptive neuronal phenomena resulting in CNS effects not observed upon acute administration. This is for example observed for selective serotonin reuptake inhibitors (SSRIs), where only chronic administration results in therapeutic antidepressant effects, whereas anxiogenic side effects are present at the beginning of therapy in animals and humans (see refs. in ref. 52).

Our new mouse model paves the way for the elucidation of the contribution of Ca1.3 to diverse physiological functions. It will be especially suitable to predict pharmacological effects of selective Ca1.3 activation on CNS function.
Received for publication October 3, 2003, and accepted in revised form February 17, 2004.
Address correspondence to: Jörg Striessnig, Abteilung Pharmakologie und Toxikologie, Institut für Pharmazie, Universität Innsbruck, Peter Mayr-Strasse 1/1, A-6020 Innsbruck, Austria. Phone: 43-512-507-5600; Fax: 43-512-507-2931; E-mail: joerg.striessnig@uibk.ac.at.

Acknowledgments
We thank G. Pelster, J. Aldrian, E. Margreiter, and P. Bauer for excellent technical assistance and M. Brennsteiner for animal care. This work was supported by grants from the Austrian Science Fund (P-14820 to J. Striessnig and T38 to M. Sinnegner-Brauns), the European Community (IHRP-CT-2000-00088), the University of Innsbruck, the Austrian National Bank, and the Swedish Research Council (8647, 12334, and 13509).


The Journal of Clinical Investigation
http://www.jci.org
Volume 113
Number 10
May 2004
1439

research article