Construction of a high-affinity receptor site for dihydropyridine agonists and antagonists by single amino acid substitutions in a non-L-type Ca\(^{2+}\) channel

GREGORY H. HOCKERMAN, BLAISE Z. PETERSON, ELIZABETH SHARP, TIMOTHY N. TANADA, TODD SCHEUER, AND WILLIAM A. CATTERALL

Department of Pharmacology, Box 357280, University of Washington, Seattle, WA 98195-7280

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ABSTRACT The activity of L-type Ca\(^{2+}\) channels is increased by dihydropyridine (DHP) agonists and inhibited by DHP antagonists, which are widely used in the therapy of cardiovascular disease. These drugs bind to the pore-forming \(\alpha_1\) subunits of L-type Ca\(^{2+}\) channels. To define the minimal requirements for DHP binding and action, we constructed a high-affinity DHP receptor site by substituting a total of nine amino acid residues from DHP-sensitive L-type \(\alpha_1\) subunits into the S5 and S6 transmembrane segments of domain III and the S6 transmembrane segment of domain IV of the DHP-insensitive P/Q-type \(\alpha_1\) subunit. The resulting chimeric \(\alpha_{1A/DHPS}\) subunit bound DHP antagonists with high affinity in radioligand binding assays and was inhibited by DHP antagonists with high affinity in voltage clamp experiments. Substitution of these nine amino acid residues yielded 86% of the binding energy of the L-type \(\alpha_{1A}\) subunit and 92% of the binding energy of the L-type \(\alpha_{1S}\) subunit for the high-affinity DHP antagonist PN200–110. The activity of chimeric Ca\(^{2+}\) channels containing \(\alpha_{1A/DHPS}\) was increased 3.5 \(\pm\) 0.7-fold by the DHP agonist (–)Bay K8644. The effect of this agonist was stereoselective as in L-type Ca\(^{2+}\) channels since (+) Bay K8644 inhibited the activity of \(\alpha_{1A/DHPS}\). The results show conclusively that DHP agonists and antagonists bind to a single receptor site at which they have opposite effects on Ca\(^{2+}\) channel activity. This site contains essential components from both domains III and IV, consistent with a domain interface model for binding and allosteric modulation of Ca\(^{2+}\) channel activity by DHPs.

Voltage-gated Ca\(^{2+}\) channels mediate Ca\(^{2+}\) influx in response to membrane depolarization and thereby initiate cellular activities such as secretion, contraction, and gene expression. Several types of voltage-gated Ca\(^{2+}\) channels have been distinguished by their physiological and pharmacological properties and have been designated L, N, P/Q, R, and T (reviewed in refs. 1 and 2). L-type Ca\(^{2+}\) channels are the molecular targets for the dihydropyridine (DHP) Ca\(^{2+}\) channel blockers that are widely used in the therapy of cardiovascular diseases; DHP modulation is the hallmark used to characterize these channels (reviewed in refs. 3 and 4).

The L-type Ca\(^{2+}\) channels consist of pore-forming \(\alpha_1\) subunits of 190 to 250 kDa in association with disulfide-linked \(\alpha_2\delta\) subunits of approximately 140 kDa, intracellular \(\beta\) subunits of 55 to 72 kDa, and, for the skeletal muscle L-type channel, an additional transmembrane \(\gamma\) subunit of 33 kDa (5). The \(\alpha_1\) subunits confer the characteristic pharmacologic and functional properties of Ca\(^{2+}\) channels, but their function is modulated by association with the auxiliary subunits. The pore-forming \(\alpha_1\) subunits can be divided into two distinct families, L-type and non-L-type, that share less than 40% amino acid identity. The L-type \(\alpha_1\) subunit family includes \(\alpha_{1L}\), which is expressed in skeletal muscle (6), \(\alpha_{1C}\), which is expressed in cardiac and smooth muscle, neurons, and many other cell types (7–9), and \(\alpha_{1D}\), which is expressed in endocrine and neuronal cells (10, 11). The non-L-type \(\alpha_1\) subunit family consists of at least three distinct gene products that are expressed primarily in neurons: \(\alpha_{1B}\) (N-type; refs. 12 and 13), \(\alpha_{1A}\) (P/Q-type; refs. 14 and 15), and \(\alpha_{1E}\) (R-type; ref. 16). The \(\alpha_1\) subunits contain four homologous domains (I through IV) that each contain six transmembrane segments (S1 through S6) (6–16).

The DHPs are allosteric modulators that act on L-type Ca\(^{2+}\) channels as either agonists or antagonists (reviewed in refs. 3, 4, and 17). Charged DHPs are thought to traverse an extracellular pathway to gain access to the DHP receptor site located within the lipid bilayer 11–14 Å from the extracellular surface of the cell membrane (18–21). Photoreactive DHPs specifically label the \(\alpha_1\) subunit of the Ca\(^{2+}\) channel (5, 22–27). The predominant sites of labeling correspond to transmembrane segment S6 in domain III (III S6) and transmembrane segment S6 in domain IV (IV S6; refs. 28–31). Analysis of chimeric Ca\(^{2+}\) channels implicated transmembrane segments III S5, III S6, and IV S6 (32–34) in DHP binding. Site-directed mutagenesis of single amino acid residues in segments III S6 and IV S6 that are conserved in all Ca\(^{2+}\) channel subtypes has large effects on DHP affinity (35, 36). In addition, mutations of residues that differ between L-type and non-L-type Ca\(^{2+}\) channels revealed multiple amino acids in transmembrane segments III S5, III S6, and IV S6 that are important determinants of high-affinity binding of DHP agonists and antagonists to L-type Ca\(^{2+}\) channels (34–38). In the experiments reported here, we have substituted nine key amino acid residues that are present in all L-type \(\alpha_1\) subunits into the non-L-type \(\alpha_{1A}\) subunit and measured both activation by DHP agonists and inhibition by DHP antagonists. The results show that these nine amino acid residues are sufficient to constitute a high-affinity receptor site for DHPs that responds appropriately to both DHP agonists and antagonists and is stereoselective like the native DHP receptor of L-type Ca\(^{2+}\) channels.

EXPERIMENTAL PROCEDURES

Construction of Mutant Ca\(^{2+}\) Channels. For the construction of \(\alpha_{1A/DHPS}\), the AscI–BsrGI fragment (nucleotides 2582–5408) from \(\alpha_{1A}\) (14) was ligated into pNeb193 (New England Biolabs) and used as template for mutagenic PCR reactions. The mutations in transmembrane segment III S5 were made in a single PCR reaction using primers that overlapped the BstXI (nucleotide 2950) and ApoI (nucleotide 4057) sites. The mutagenic mismatches were incorporated by the primer overlapping the ApoI site. The mutations in III S6 were made in a

Abbreviation: DHP, dihydropyridine.
single PCR reaction using primers that overlapped the XcmI (nucleotide 4316) and BspEI (nucleotide 5102) sites. The mutagenic mismatches were incorporated by the primer overlapping the XcmI site. The mutations in IVS6 were made using the splice overlap extension method (39). In the first reactions, mutagenic primers annealing within the cDNA encoding IVS6 were paired with primers that overlapped either the BspEI or the NorI (nucleotides 5102 and 5384, respectively) sites that flank IVS6. These overlapping fragments were coupled and amplified in a subsequent PCR reaction to give the full-length BspEI–NorI fragment containing the mutations in IVS6. The PCR products containing the desired mutations were assembled in AscI–BglII–pNBE193, and the AscI–BglII fragment was ligated into full-length α1A in the expression vector pMT2. The desired mutations were verified, and the integrity of the clone was confirmed by cDNA sequencing and extensive restriction digest analysis.

Expression of Ca2+ Channels. Human tsA-201 cells, a simian virus 40 (SV40) T-antigen expressing derivative of the human embryonic kidney cell line HEK293 (a gift of Robert DuBridge, Cell Genetics, Foster City, CA), were maintained in DMEM/F-12 (GIBCO/BRL) enriched with 10% fetal bovine serum. Human tsA-201 cells were cotransfected with α1A, α1ADHPS, or α1CII (9); β1a (40); α2δ (41); and CD8 antigen (EBO-pCD-Leu2, American Type Culture Collection) such that the molar ratio of the plasmids was 1:1:1:0.8. Cells were transfected by Ca2+ phosphate precipitation (42), and cells were replated at low density for electrophysiological recording 20–24 hours later. The α1CII cDNA was in the expression plasmid ZEM 229 (a gift of Eileen Mulvihill, Zymogenetics, Seattle). The α2δ cDNA was in the expression plasmid ZEM 228 (a gift of Eileen Mulvihill). The α1IA, α1ADHPS, and β1b cDNA were in the expression plasmid pMT2 (Genetics Institute, Boston).

Electrophysiology. Transfected cells were recognized by labeling with anti-CD8 antibody-coated beads (M450 CD8 Dynabeads, Dynal). Barium currents through Ca2+ channels were recorded using the whole-cell configuration of the patch clamp technique. Patch electrodes were pulled from VWR micropipettes and fire-polished to produce an inner tip diameter of 4–6 μm. Currents were recorded using an Axon Instruments Axopatch 200B patch clamp amplifier and filtered at 1 or 2 kHz (8-pole Bessel filter, ~3 dB). Voltage pulses were applied and data were acquired using pClamp6 software (Axon Instruments). Linear leak and capacitance currents have been subtracted using an on-line P/4 subtraction paradigm. (+)PN200–110 was applied to cells using a fast perfusion system with background perfusion. (~)Bay K 8644 was added to the bath, without background perfusion, as a 10× stock. The bath saline contained 150 mM of Tris, 2 mM of MgCl2, and 10 mM of BaCl2. The intracellular saline contained 130 mM of N-methyl-D-glucamine, 10 mM of EGTA, 60 mM of Hesper, 2 mM of MgATP, and 1 mM of MgCl2. The pH of both solutions was adjusted to 7.3 with methanesulfonic acid. All experiments were performed at room temperature (20–23°C).

Preparation of Membranes. Transfected tsA-201 cells were washed twice, scraped from the cell culture dish, and homogenized using a glass-teflon homogenizer in Buffer A (50 mM of Tris/100 μM of phenylmethylsulfonyl fluoride/100 μM of benzamidine/1.0 μM of pepstatin A/1.0 μg/ml of leupeptin/2.0 μg/ml of aprotinin, pH 8.0). The homogenate was centrifuged at 700 × g for 5 min. The resulting pellet was discarded and the supernatant was centrifuged 30 min at 100,000 × g. The supernatant was discarded and the membrane pellet was washed and homogenized in Buffer A. The resulting membrane homogenate was divided into aliquots and stored at −80°C for up to 3 months without detectable loss of (+)[3H]PN200–110 binding activity.

Radioligand Binding. Equilibrium binding assays were performed in Buffer A with 20–200 μg of membrane protein, 0.01–5 nM of (+)[3H]PN200–110, and 1 mM of Ca2+ at 32°C for 180–210 min. Nonspecific binding was determined in the presence of 1 μM (+)-PN200–110, and bound and free ligands were separated by vacuum filtration over GF/C glass fiber filters. Filters were washed using ice-cold wash buffer (10 mM of Tris/1% polyethylene glycol 8000/0.1% BSA/0.01% Triton X-100, pH 8.0), and bound radioactivity was detected by liquid scintillation counting. Dissociation constants (Kd) were determined using the radioligand data analysis program LIGAND (Biosoft, Cambridge, U.K.). All data are means ± SEM.

RESULTS

Construction of a High Affinity DHP Site in α1A. The key amino acid residues required for high affinity DHP binding as determined in previous studies are illustrated in Fig. 1. Photoaffinity labeling followed by antibody mapping of the labeled peptide fragments showed that transmembrane segments IIIS6 and IVS6 are components of the DHP receptor site in L-type Ca2+ channels (refs. 28 and 29; Fig. 1A, shaded segments). Construction and analysis of chimeric Ca2+ channels confirmed the importance of transmembrane segments IIIS6 (32) and IVS6 (32–34) and further demonstrated an important role of transmembrane segment IIIS5 (ref. 32; Fig. 1A, diagonally hatched segment). Mutation of single amino acid residues identified three conserved amino acid residues in transmem-
brane segment IIIS6 and two in IVS6 that are important for DHP binding and are present in all Ca\(^{2+}\) channels (refs. 35 and 36; Fig. 1B, white circles, lowercase letters). In addition, two L-type-specific residues in segment IISS5, three in segment IIIS6, and four in segment IVS6 were found to be important for DHP binding (refs. 34–38; Fig. 1B, dark circles). The success of experiments with chimeric Ca\(^{2+}\) channels in which segments of L-type Ca\(^{2+}\) channels are transferred into non-L-type Ca\(^{2+}\) channels suggests that the basic structure of non-L-type channels is appropriate to support DHP binding, even though the amino acid sequence is less than 40% identical. Similarly, the requirement for amino acid residues that are conserved between L-type and non-L-type Ca\(^{2+}\) channels for high-affinity DHP binding suggests that the DHP binding site contains structural features that are common to all Ca\(^{2+}\) channels. If all of the L-type-specific amino acid residues that are required for high affinity DHP binding have been identified, substitution of the nine amino acid residues highlighted as dark circles in Fig. 1B into a non-L-type Ca\(^{2+}\) channel should be sufficient to construct a high-affinity DHP binding site. To test this idea, we substituted these nine amino acid residues for their counterparts in the DHP-insensitive rba isoform of the \(\alpha_{1A}\) subunit of P/Q-type Ca\(^{2+}\) channels using site-directed mutagenesis methods as described in Experimental Procedures.

Functional Characteristics of the DHP-Sensitive \(\alpha_{1A}\) Chimera. The resulting \(\alpha_{1A/DHPS}\) subunit was expressed in tsA-201 cells with \(\alpha_{2\delta}\) and \(\beta_{1b}\) subunits and analyzed by whole cell voltage clamp as described in Experimental Procedures. Barium currents through the \(\alpha_{1A/DHPS}\) subunit activated normally but inactivated more rapidly than wild-type \(\alpha_{1A}\) (Fig. 2A; \(\alpha_{1A}\), \(\tau_{n} = 279\) ms; \(\alpha_{1A/DHPS}\), \(\tau_{n} = 123\) ms). The two channels had similar current-voltage relationships with peak barium current near 0 mV (Fig. 2B). However, the voltage dependence of inactivation of \(\alpha_{1A/DHPS}\) was shifted 40 mV toward more negative membrane potentials compared with wild-type \(\alpha_{1A}\) (Fig. 2C; \(\alpha_{1A}\), \(V_{50/12} = -48\) mV; \(\alpha_{1A/DHPS}\), \(V_{50/12} = -88\) mV). Thus, the \(\alpha_{1A/DHPS}\) chimera is functional as a Ca\(^{2+}\) channel, but has altered inactivation properties.

Inhibition of Barium Currents Through \(\alpha_{1A/DHPS}\) by a DHP Antagonist. DHP antagonists bind with higher affinity to the inactivated state of L-type Ca\(^{2+}\) channels (43–45). To compare the DHP sensitivity of wild-type \(\alpha_{1A}\) and \(\alpha_{1A/DHPS}\), we compensated for the shift in the voltage dependence of inactivation by measuring the block of barium currents from holding potentials that differed by 40 mV: -80 mV for \(\alpha_{1A}\) and -120 mV for \(\alpha_{1A/DHPS}\). These conditions yield the same ratio of resting and inactivated channels for each \(\alpha_{1}\) subunit. The more negative membrane potential used for the \(\alpha_{1A/DHPS}\) chimera would reduce inhibition of L-type Ca\(^{2+}\) channels by DHPs, so it is unlikely that this membrane potential per se would increase inhibition of the \(\alpha_{1A/DHPS}\) chimera.

The high-affinity DHP antagonist PN200–110 (isradipine) had no effect on P/Q-type Ca\(^{2+}\) channels containing \(\alpha_{1A}\), even at the high concentration of 10 \(\mu\)M (Fig. 3A). In contrast, chimeric Ca\(^{2+}\) channels containing \(\alpha_{1A/DHPS}\) were substantially inhibited by 10 \(nM\) of PN200–110 and completely inhibited by 100 \(nM\) of PN200–110 (Fig. 3A). The IC\(_{50}\) value estimated from titration experiments was 13.8 \(nM\) \pm 3 \(nM\) (Fig. 3C; \(n = 9\)). This compares favorably with the IC\(_{50}\) of 6.8 \(nM\) \pm 1 \(nM\) for block of L-type Ca\(^{2+}\) channels containing \(\alpha_{1C}\) expressed with \(\alpha_{2\delta}\) and \(\beta_{1b}\) in the same cell line (Fig. 3B). Ca\(^{2+}\) channels containing \(\alpha_{1A}\) are unaffected by PN200–110 up to at least 10 \(\mu\)M. Based on this, the minimum estimate of the IC\(_{50}\) for inhibition of P/Q-type Ca\(^{2+}\) channels containing \(\alpha_{1A}\) is 250 \(\mu\)M, more than four orders of magnitude higher than for \(\alpha_{1A/DHPS}\). Thus, substitution of nine amino acid residues in \(\alpha_{1A}\) is sufficient to constitute a high affinity receptor site for a DHP antagonist with more than 10,000-fold higher affinity than wild-type \(\alpha_{1A}\).

Increase of Barium Currents Through \(\alpha_{1A/DHPS}\) by a DHP Agonist. DHPs are allosteric modulators of L-type Ca\(^{2+}\) channels and can either inhibit or enhance channel activity (3, 4, 17). Activation of L-type Ca\(^{2+}\) channels is increased by DHP agonists like (-)Bay K8644 (17). Our previous results on mutations in the \(\alpha_{1S}\) subunit of L-type Ca\(^{2+}\) channels indicated that many of the same amino acid residues are required for binding of both DHP agonists and antagonists (35). To test whether DHP agonists can also bind and act at the minimal DHP receptor site constructed in \(\alpha_{1A/DHPS}\), we measured barium currents from a holding potential of -100 mV in the presence and absence of 10 \(\mu\)M (-) Bay K8644 (Fig. 4). Bay K8644 increased Ca\(^{2+}\) currents through L-type Ca\(^{2+}\) channels containing \(\alpha_{1C}\), but had no effect on P/Q-type channels containing \(\alpha_{1A}\) (Fig. 4 Left). In contrast, (-) Bay K8644 increased barium currents through chimeric Ca\(^{2+}\) channels containing \(\alpha_{1A/DHPS}\) by 3.53 \pm 0.66-fold (Fig. 4 Left; \(n = 3\)). The increased barium current was observed over a wide range of test pulse potentials, as for L-type Ca\(^{2+}\) channels containing \(\alpha_{1C}\) (Fig. 4 Right). In contrast, (-) Bay K8644 had no effect on barium currents through P/Q-type Ca\(^{2+}\) channels containing \(\alpha_{1A}\) at any test pulse potential (Fig. 4 Right). Thus, the minimal DHP receptor site of \(\alpha_{1A/DHPS}\) is sufficient for both binding and functional effect of a DHP agonist.
data were fit to the equation $y = 1.2 nM (36)$. The asterisk over the IC$_{50}$ value for the axes for $a$ analyzed the data by Scatchard plot (Fig. 5 $a$) from cells expressing $a$.

For Bay K8644, the $(+)$ current traces from cells expressing the indicated channel types in the absence of drug. Cells expressing $a$ or $a$ were depolarized to $100 mV$ for 100 ms every 20 seconds. After a control baseline had been established, PN200–110 was perfused onto the cells at the concentrations indicated. (B) IC$_{50}$ values for inhibition of $a$, $a$, and $a$ by PN200–110. The IC$_{50}$ for the $a$-channels was $13.8 \pm 3.0 nM (n = 9)$; for the $a$-channel, IC$_{50}$ was $6.8 \pm 1.2 nM (36)$. The asterisk over the IC$_{50}$ value for the $a$ channel (250 $\mu M$) is to indicate that this value is a lower limit since no block was detected at $10 \mu M$ PN200–110, the highest concentration studied. (C) Concentration-response curve of $a$ for PN200–110. The mean data were fit to the equation $1 - 1/(1+([IC_{50}]/[PN200-110]))$. Standard error ($n = 9$) was smaller than the diameter of the symbols in the figure. Current amplitudes at each drug concentration were expressed as a fraction of the current amplitude in the absence of drug.

The functional effects of many DHP agonists are stereospecific. For Bay K8644, the $(+)$ enantiomer is an agonist while the $(−)$ isomer is an antagonist (17). As for $L$-type $Ca^{2+}$ channels containing $a$, we found that chimeric $Ca^{2+}$ channels containing $a$ are inhibited by $(+)$ Bay K8644 (IC$_{50} = 303 \pm 102 nM$, $n = 8$; data not shown), and no activating effect of $(+)$ Bay K8644 was observed at any concentration tested. Thus, the stereoselectivity of DHP binding and action is also present in the minimal DHP receptor site constructed in $a$.

**High Affinity Binding of PN200–110 to $a$ in Cell Membrane Preparations.** Inhibition of barium currents by PN200–110 measures primarily drug binding to the resting state of $Ca^{2+}$ channels that is predominant at the negative holding potential. Because DHP antagonists have higher affinity for the inactivated state of $Ca^{2+}$ channels (43–45), their binding affinity is higher in radioligand binding assays using membrane preparations whose membrane potential is near 0 mV where inactivation is complete. We first compared binding of $[^{3}H]$PN200–110 to a fixed concentration of 4.4 nM. $Ca^{2+}$ channels containing $a$ bound $[^{3}H]$PN200–110 specifically (63.7 fmol/mg cell protein) while those containing $a$ did not have significant specific binding. As expected from the electrophysiological results, $Ca^{2+}$ channels containing $a$ also bound $[^{3}H]$PN200–110 specifically (36.9 fmol/mg cell protein), but not as well as channels containing $a$. To compare the binding affinity of $a$ and $a$ quantitatively, we carried out a saturation binding study (Fig. 5$A$) and analyzed the data by Scatchard plot (Fig. 5$B$, note the different axes for $a$ and $a$). The results indicate a $K_d$ for binding to $a$ of 50 pM compared with a $K_d$ for $a$ of 1.48 nM, 27-fold higher. Thus, the $a$ chimeras have high affinity for PN200–110 in a radioligand binding assay as well as in electrophysiological experiments, but the difference in binding between $a$ and $a$ is greater in the ligand binding assay.

**Fig. 3.** Inhibition of $Ca^{2+}$ channels containing $a$, $a$, and $a$ subunits by the DHP antagonist PN200–110. (A) Representative current traces from cells expressing $a$ or $a$ in the presence and absence of the indicated concentrations of PN200–110. Cells expressing $a$ or $a$ were held at $−80$ or $−120$, respectively, and depolarized to 0 mV for 100 ms every 20 seconds. After a control baseline had been established, PN200–110 was perfused onto the cells at the concentrations indicated. (B) IC$_{50}$ values for inhibition of $a$, $a$, and $a$ by PN200–110. The IC$_{50}$ for the $a$-channels was $13.8 \pm 3.0 nM (n = 9)$; for the $a$-channel, IC$_{50}$ was $6.8 \pm 1.2 nM (36)$. The asterisk over the IC$_{50}$ value for the $a$ channel (250 $\mu M$) is to indicate that this value is a lower limit since no block was detected at $10 \mu M$ PN200–110, the highest concentration studied. (C) Concentration-response curve of $a$ for PN200–110. The mean data were fit to the equation $1 - 1/(1+([IC_{50}]/[PN200-110]))$. Standard error ($n = 9$) was smaller than the diameter of the symbols in the figure. Current amplitudes at each drug concentration were expressed as a fraction of the current amplitude in the absence of drug.

**Fig. 4.** Modulation of $Ca^{2+}$ channels containing $a$, $a$, or $a$ subunits by the DHP agonist $(+)$ Bay K 8644. (Left) Current traces from cells expressing the indicated channel types in the absence and presence of 10 $\mu M$ of $(+)$ Bay K 8644. Cells expressing $a$, $a$, or $a$ were depolarized from a holding potential of $−100 mV$ to 0 mV for 100 ms. Cells expressing $a$ or $a$ were held at $−60$ mV and depolarized to $+10 mV$ for 100 ms. For $Ca^{2+}$ channels containing $a$, $a$ or $a$, a series of ten 1 s depolarizations to 0 mV was used to bring the agonist effect to equilibrium. (Right) Current-voltage relationship before and after addition of 10 $\mu M$ of $(−)$ Bay K 8644 is shown for each channel type. The peak current amplitude of each I/V curve in the absence of $(−)$ Bay K 8644 was set to 1, and the current amplitudes in the presence of $(−)$ Bay K 8644 were expressed relative to the current amplitudes in the absence of drug. Cells expressing $a$, $a$, or $a$ were held at $−80$ mV or $−120$ mV. From these holding potentials, cells were depolarized to the indicated test pulse potentials from $−50 mV$ to $+50 mV$ in 10 mV increments for 100 ms, and barium currents were recorded.

**DISCUSSION**

**Molecular Requirements for a High Affinity DHP Receptor Site.** Substitution of nine amino acid residues previously shown to be required for DHP binding (34–38) into the sequence of the non-$L$-type $a$ subunit is sufficient to construct a high-affinity DHP receptor site. This site was constructed in the structural context of the $a$ subunit that contains at least five conserved residues that are necessary for high-affinity DHP binding (35, 36). The resulting chimera $a$ has many of the pharmacological properties of an $L$-type $Ca^{2+}$ channel with respect to DHPs. The IC$_{50}$ for inhibition of $Ca^{2+}$ current by PN200–110 is 13.8 nM, only twofold higher than the IC$_{50}$ value for inhibition of $Ca^{2+}$ channels containing $a$ expressed under the same conditions. Inhibition of barium currents measures primarily the affinity of the resting state for DHPs because of the negative holding potential used in these experiments. In contrast, the binding experiments on membranes from transfected cells measure primarily the higher affinity of the inactivated state because of the prolonged depolarization of the membranes in the binding assay. The $K_d$ for binding of $[^{3}H]$PN200–110 to $Ca^{2+}$ channels containing $a$ is 1.48 nM, compared with 55 pM for $L$-type $Ca^{2+}$ channels containing $a$ (36) and 250 pM $L$-type $Ca^{2+}$ channels containing $a$ (35) expressed under the same conditions. Thus, the amino acid substitutions in $a$ are more
Fig. 5. Binding of [3H]PN200–110 to membrane preparations from cells expressing α1C or α1ADHPS channels. (A) Equilibrium binding of [3H]PN200–110 to membrane preparations from cells expressing α1C (open circles) or α1ADHPS (solid circles) subunits. Specific binding was determined by subtracting nonspecific binding of [3H]PN200–110 in the presence of 1 μM of unlabeled PN200–110 from total binding in the absence of the unlabeled drug at each [3H]PN200–110 concentration. Means of the data at each concentration are plotted ± SEM (n = 3). (B) Scatchard analysis of equilibrium binding of [3H]PN200–110 to α1C (open circles; dashed line) and α1ADHPS channels (solid circles; solid line). The ordinate axes are different for each subunit type to normalize the data. The Kd values in this experiment were 79 pM for α1C and 1.48 nM for α1DHPS.

A Single Receptor Site for DHP Agonists and Antagonists.

DHP agonists inhibit high-affinity binding of DHP antagonists, consistent with the idea that they share the same receptor site (45–47). However, binding studies on intact cells reveal complex interactions between binding of agonist and antagonist DHPs, and it has been hypothesized that agonists and antagonists occupy different sites when causing their opposite effects on Ca2+ channel function (48). In contrast to this hypothesis, our results show conclusively that DHP agonists and antagonists occupy the same receptor site and cause opposite effects on Ca2+ channel activity when bound there. Each of the nine single amino acids substituted in α1ADHPS is required for high-affinity antagonist binding in 1-type Ca2+ channels, and mutations of many of these residues also reduce affinity for binding of agonists to 1-type channels. Construction of a minimal DHP receptor site for high-affinity antagonist binding also is sufficient to confer stereoselective agonist binding and action. Molecular differences in the interactions of these two classes of drugs with the same receptor site must be responsible for the dramatic differences in their pharmacological effects.

A Domain-Interface Model for DHP Binding and Action.

Based on the original photoaffinity labeling of the DHP receptor site, it was proposed that DHPs bind to a single site at the interface of domains III and IV (30). This model is confirmed by the results presented here, which show that the minimal receptor site sufficient for high-affinity binding of DHP antagonists and for stereoselective binding and action of DHP agonists includes amino acid residues in transmembrane segments S5 and S6 in domain III and transmembrane S6 in domain IV. Allosteric effectors of enzymes most often bind to sites at the interfaces between subunits or domains (e.g., refs. 49 and 50), and current evidence indicates that the agonist binding site of nicotinic acetylcholine receptors is at a subunit interface (51). Evidently, these sites are particularly sensitive to binding of small ligands, which can alter interactions between domains or subunits when bound at their interface. Based on these analogies, it is likely that DHPs act as allosteric effectors that alter domain-domain interactions within the α1 subunits of Ca2+ channels. Determination of the structural basis for the opposite, stereoselective effects of DHP agonists and antagonists on Ca2+ channel function when bound at the DHP receptor site between domains III and IV will provide further information on the mechanism of action of these drugs and on the interactions between these Ca2+ channel domains in channel gating.