Assessing potentiation of the \((\alpha_4)^3(\beta_2)^2\) nicotinic acetylcholine receptor by the allosteric agonist CMPI

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The extracellular domain of the nicotinic acetylcholine receptor isoforms formed by three α4 and two β2 subunits ((α4)3(β2)2 nAChR) harbors two high-affinity “canonical” acetylcholine (ACh)-binding sites located in the two α4:β2 intersubunit interfaces and a low-affinity “noncanonical” ACh-binding site located in the α4:α4 intersubunit interface. In this study, we used ACh, cytisine, and nicotine (which bind at both the α4:α4 and α4:β2 interfaces), TC-2559 (which binds at the α4:β2 but not at the α4:α4 interface), and 3-(2-chlorophenyl)-5-(5-methyl-1-(piperidin-4-yl)-1H-pyrazol-4-yl)isoxazole (CMPI, which binds at the α4:α4 but not at the α4:β2 interface), to investigate the binding and gating properties of CMPI at the α4:α4 interface. We recorded whole-cell currents from Xenopus laevis oocytes expressing (α4)3(β2)2 nAChR in response to applications of these ligands, alone or in combination. The electrophysiological data were analyzed in the framework of a modified Monod–Wyman–Changeux allosteric activation model. We show that CMPI is a high-affinity, high-efficacy agonist at the α4:α4 binding site and that its weak direct activating effect is accounted for by its inability to productively interact with the α4:β2 sites. The data presented here enhance our understanding of the functional contributions of ligand binding at the α4:α4 subunit interface to (α4)3(β2)2 nAChR-channel gating. These findings support the potential use of α4:α4 specific ligands to increase the efficacy of the neurotransmitter ACh in conditions associated with decline in nAChRs activity in the brain.

Neuronal nicotinic acetylcholine receptors (nAChRs) are pentameric ligand-gated ion channels formed of identical or distinct but homologous subunits (α2–α10 and β2–β4). Homomeric α7 and heteromeric α4β2 nAChRs are the major subtypes in the brain (1–3). Postsynaptic nAChRs mediate fast synaptic transmission, whereas presynaptic nAChRs modulate the release of many neurotransmitters (4, 5). Thus, nicotinic receptors are involved in complex brain functions, including cognition, pain perception, and neuronal survival during aging (3, 5). Furthermore, nAChRs mediate the behavioral effects of nicotine, the major addictive component of tobacco smoking, and are considered a major molecular target for pharmacotherapeutic interventions to manage nicotine dependence (6, 7). Therapeutics targeting the nAChRs also have potential clinical relevance in reducing chronic pain and slow cognitive decline associated with neuropsychiatric conditions (8).

Neuronal nAChRs consisting of α4 and β2 subunits assemble in two stoichiometries: ((α4)2(β2)3 and (α4)3(β2)2 (9). The initial pharmacological distinction between the (α4)2(β2)3 and (α4)3(β2)2 isoforms was based on their sensitivity to acetylcholine (ACh). ACh potency (EC50) is ~1 μM at the (α4)2(β2)3 nAChR and ~100 μM at the (α4)3(β2)2 nAChR; the two isoforms are thus referred to as the high- and low-sensitivity α4β2 nAChRs (10). Both isoforms contain two high-affinity agonist binding site (ABS) located in the two α4:β2 intersubunit interfaces in the extracellular domain, whereas the (α4)3(β2)2 nAChR has a third, low-affinity ACh-binding site located in the α4:α4 subunit interface. Subsequent studies on heterologously expressed (α4)3(β2)2 and (α4)2(β2)3 nAChRs revealed a number of key differences in channel functional properties and pharmacological selectivity to exogenous nAChR ligands (11–15). Assembly of both α4β2 nACh isoforms has been reported in vivo (16). The (α4)3(β2)2 nAChR isoform is considered the major isoform expressed in the cortex (17), whereas the (α4)2(β2)3 nAChR isoform contributes to nicotine dependence and is selectively upregulated and stabilized after chronic nicotine exposure (18–21).

High-throughput screening has identified several nAChR subtype-selective positive allosteric modulators (PAMs) (12, 22–26). CMPI (3-(2-chlorophenyl)-5-(5-methyl-1-(piperidin-4-yl)-1H-pyrazol-4-yl)isoxazole) and NS9283 (3-[3-(3-pyridinyl)-1,2,4-oxadiazol-5-yl]benzonitrile) have been identified as potent nAChR PAMs that preferentially potentiate the (α4)3(β2)2 isoform (12, 23). At first glance, the pharmacology of CMPI and NS9283 (location of binding site and effect on ACh concentration-response curve) at the (α4)3(β2)2 nAChR resembles that of benzodiazepines at the GABA_A receptor (27). However, unlike benzodiazepines and GABA at the GABA_A receptor, CMPI, NS9283, and ACh share an overlapping binding site at the α4:α4 subunit extracellular interface in the (α4)3(β2)2 nAChR (12, 28), raising the possibility that CMPI and NS9283 could act as agonists at the α4:α4

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Indeed, NS9283, which binds at the α4:α4 interface and at the β2:α4 pseudo-agonist site (29), was found to enhance nAChR (α4)3(β2)2 channel activity by transitioning the channel into a preactivated state (30).

In this study, we investigated the properties of CMPI and other nAChR ligands to delineate the pharmacology of the α4:α4 binding site and to elucidate allosteric interaction between the α4:β2 and α4:α4 interface-binding sites. Current responses from the (α4)3(β2)2 nAChR elicited by a series of nAChR agonists, alone or in combination with CMPI, were analyzed using a modified Monod-Wyman-Changeux (MWC) allosteric activation model (31–33). Our results indicate that CMPI is a high-affinity, high-efficacy agonist at the α4:α4 binding site. It binds to the α4:α4 interface with a higher affinity than ACh, cytisine, or nicotine, and efficaciously potentiates receptor responses to subsaturating concentrations of these agonists. The gating efficacy of CMPI at the α4:α4 site is equivalent to that of ACh, whereas weak direct activation of the (α4)3(β2)2 nAChR in the presence of CMPI is accounted for by a single binding site mediating its action. Thus, CMPI enhances channel gating triggered by ACh at the α4:β2 sites by providing ligand occupancy at the α4:α4 site, which is otherwise vacant or only occupied at very high (hundreds of μM) ACh concentrations. The data presented here enhance our understanding of ligand-binding properties and functional contributions of the “noncanonical” α4:α4 subunit interface to

Figure 1. Effects of CMPI and NS9283 on (α4)3(β2)2 nAChR current responses elicited by subsaturating and saturating agonist concentrations. The traces show whole-cell currents elicited by 10 s applications of low or saturating concentrations of various nAChR agonists alone or in the presence of 1 μM of CMPI or NS9283. Representative traces for ACh and TC-2559 are shown in (A and B), respectively. C, for each agonist concentration, the peak currents were normalized to the peak current elicited by agonist alone. The data obtained from several oocytes were plotted as mean ± SD with values of individual oocytes shown in open circles. The probability (P) that calculated potentiation ratio differs from no potentiation (PR = 1) was analyzed using one-way ANOVA with multiple comparisons versus control group (Holm–Sidak method, SigmaPlot, and Systat Software Inc). The effects of 1 μM of CMPI or NS9283 on current elicited by 10 μM ACh, 1 μM Nicotine, 3 μM SI-A85380, 1 μM Cytisine, 1 μM TC 2559, 10 μM TC 2559, and 30 μM TC 2559 were statistically significant with a p < 0.001. The data for 10 μM ACh + 1 μM CMPI and 10 μM ACh + 1 μM CMPI contain data from oocytes that were reported previously (28). ACh, acetylcholine; CMPI, 3-(2-chlorophenyl)-5-(5-methyl-1-(piperidin-4-yl)-1H-pyrazol-4-yl)isoxazole; nAChR, nicotinic acetylcholine receptor.
nAChR channel gating and facilitate structure-based design of novel therapeutics that selectively target the (α4)3β2 nAChR.

Results

CMPI potentiation of (α4)3β2 nAChR currents induced by subsaturating and saturating agonist concentrations

Our initial characterization of CMPI-induced potentiation of the (α4)3β2 nAChR employed the neurotransmitter ACh as the agonist. Coapplication of CMPI increased current responses to EC50 of 0.18 ± 0.03 μM (28). CMPI at 1 μM produced a ~100 fold left-shift of the ACh concentration-response curve enhancing ACh potency with no apparent effect on the ACh maximal response (15, 28). ACh is known to potentiate responses to ACh, nicotine, cytisine, and TC-2559 when the agonist concentrations below 1 μM; however, at saturating concentrations greater than 1 μM, CMPI potentiation of cytisine responses gradually declined reaching no effect at 100 μM cytisine (I100 μM cytisine + 1 μM CMPI = 91 ± 5%). In contrast, CMPI significantly increased TC-2559 efficacy at (α4)3β2 nAChRs (Emax + 1 μM CMPI = 576 ± 26% of that of 100 μM TC-2559 control) with less pronounced effects on TC-2559 potency (EC50 = 0.34 ± 0.04; EC50 + 1 μM CMPI = 0.43 ± 0.04 μM; Emax + 1 μM NS9283 = 141 ± 9%), cytisine (EC50 + 1 μM NS9283 = 0.013 ± 0.002 μM; Emax + 1 μM NS9283 = 188 ± 5%; I100 μM cytisine + 1 μM NS9283 = 113 ± 9%), and TC-2559 (EC50 + 1 μM NS9283 = 0.1 ± 0.02 μM; Emax + 1 μM NS9283 = 468 ± 30%) (data not shown).

Figure 2. Effects of coapplication of CMPI on the concentration-response curves of nicotine, cytisine, and TC-2559 at the (α4)3β2 nAChR. The whole-cell current elicited by 10 s applications of increasing concentrations of nicotine (A), cytisine (B), or TC-2559 (C) in the absence or presence of 1 μM CMPI. For each drug application, the peak currents were normalized to the peak current elicited by 100 μM nicotine (A), 100 μM cytisine (B), or 10 μM TC-2559 (C) applied in the same recording run. The recording runs from same oocyte were combined and each point plotted are mean ± SD of data obtained from at least three oocytes. The data were fit to a single site model using Equation 1. CMPI, 3-(2-chlorophenyl)-5-(5-methyl-1-(piperidin-4-yl)-1H-pyrazol-4-yl)isoxazole; nAChR, nicotinic acetylcholine receptor.
CMPI pharmacology at (α4)3(β2)2 nAChRs

Dissimilar effects of CMPI on the concentration-response curves for nicotine, cytisine, and TC-2559 were not solely reflections of the differences in agonist efficacies. There was no correlation between agonist efficacy and the extent of CMPI potentiation ($E_{\text{max}}$) or CMPI concentration that produced half-maximum potentiation ($EC_{50}$) (Fig. 3). The current responses produced by a saturating concentration (100 μM) of cytisine and nicotine were 10 ± 1 and 73 ± 3% of current elicited by 1 mM ACh. CMPI potentiated the responses induced by the coapplication of cytisine or nicotine (which elicited 2.3 ± 0.4 and 2.9 ± 0.4% of current induced by 1 mM ACh, respectively) with similar potencies ($EC_{50}$s of 0.34 ± 0.04 and 0.54 ± 0.02 μM, respectively) but has lower apparent efficacy at (α4)3(β2)2 than (α4)2(β2)3 nAChRs ($I_{30}$ μM TC-2559 = 11 ± 2% and 152 ± 21% of current induced by 1 mM ACh, respectively) (Fig. 4). This is consistent with TC-2559 activating the receptor through the two α4β2 ABS in both isoforms. The α4α4 interface (Fig. 4B) is formed by residues from a (+) face of one α4 subunit and a (−) face of the adjacent α4 subunit. Amino acid residues forming the (−) face of α4 are unique and impose an additional layer of selectivity on agonist binding at the α4α4 subunit interface (11, 35). Furthermore, amino acid substitutions at the α4 subunit (−) face have been shown to enable binding of agonists with larger molecular volumes at the α4α4 subunit interface (34). Alanine substitution at α4H116 (α4H142 when amino acid numbering includes the signal peptide) within the α4 subunit (−) face allows TC-2559 to bind to the α4α4 site (34) and results in increased TC-2559 efficacy ($E_{\text{max}}$ of 182 ± 5% of current induced by 1 mM ACh versus ~10% in WT) (Fig. 4C).

We have previously shown that amino acid substitutions at positions α4K64 and α4E66, but not α4H116, significantly reduce CMPI-mediated potentiation of ACh-induced currents in the (α4)3(β2)2 nAChR (28). Similarly, CMPI did not potentiate TC-2559-induced current responses in (α4)3(β2)2 nAChR containing α4K64T or α4E66I (Fig. 5A). In addition, mutations at α4H116, which abolish potentiation by NS9283, did not affect CMPI potentiation of current responses of (α4)3(β2)2 nAChR induced by submaximal TC-2559 concentrations.

These results indicate that CMPI binds in the (α4)3(β2)2 nAChR to the same site and interacts with the same amino acid residues in the presence of ACh or TC-2559. The effect of

Figure 3. Effect of coapplication of CMPI with agonist on (α4)3(β2)2 nAChR. CMPI concentration-dependent potentiation of (α4)3(β2)2 nAChR current responses induced by 1 μM cytisine or nicotine (A), 1 μM TC-2559 (B), or 10 μM TC-2559 (C) in the absence and presence of increasing concentrations of CMPI. The peak currents were normalized to the peak current elicited by agonist alone applied in the same recording run. Recording runs from the same oocyte were combined and each point plotted are mean ± SD of data obtained from at least three oocytes. The data were fit to a single site model using Equation 1. CMPI, 3-(2-chlorophenyl)-5-(5-methyl-1-(piperidin-4-yl)-1H-pyrazol-4-yl)isoxazole; nAChR, nicotinic acetylcholine receptor.
amino acid substitutions at position αH116 on CMPI potentiation of responses to saturating concentrations of TC-2559 was remarkable (Fig. 5, B–D). CMPI potentiation ratios of current induced by 30 μM TC-2559 in (α4)3(β2)2 nAChR containing amino acid substitution at αH116 to leucine (α4H116L), valine (α4H116V), or alanine (α4H116A) were 4.58 ± 0.24, 3.01 ± 0.39, and 0.74 ± 0.13, respectively (Fig. 5D). Statistical analyses of the effects of these mutations on CMPI and NS9283 potentiation of responses to saturating concentrations of ACh and TC-2559 are shown in Table 1. The effect of 1 μM CMPI on current induced by 30 μM TC-2559 in (α4H116A)3(β2)2 nAChR was significantly different from that in WT (α4)3(β2)2 and not significantly different from no potentiation (Table 1). The simplest interpretation of this decline in CMPI-potentiation ratio is that reduction in the molecular volume of the aliphatic amino acid residue at position αH116 (molecular volumes of L, V, and A are 166.7, 140.0, and 88.6 Å³, respectively) increases the affinity of TC-2559 at the α4:α4 interface, and thus at high TC-2559 concentrations, it reduces the ability of CMPI to bind at α4:α4 interface. Indeed, CMPI concentration-dependent potentiation and its effect on the TC-2559 concentration-response curve of (α4)3(β2)2 nAChR containing αH116A substitution (Fig. 6) mirrored the effect of CMPI on ACh-induced current responses of WT (α4)3(β2)2 nAChR. CMPI potentiated TC-2559-induced current at (α4H116A)3(β2)2 nAChR with EC50 of 0.2 ± 0.1 μM and to Imax of 634 ± 77% of that of control. In the absence and presence of 1 μM CMPI, the EC50 of TC-2559 at (α4H116A)3(β2)2 nAChR were 22 ± 7 and 0.97 ± 0.32 μM, respectively. The Emax were 115 ± 14 and 83 ± 5% in the absence and presence of CMPI.
Analysis of $\alpha_4^3\beta_2^2$ receptor activation in the framework of the Monod–Wyman–Changeux allosteric model

To gain mechanistic insight into receptor activation in the presence of CMPI, we analyzed the currents in the framework of a cyclic two-state (resting and active) allosteric activation model (31–33). In this model, channel opening is mediated by stabilization of the active state by an agonist that, by definition, has higher affinity to the active than resting state.

Table 1
Effect of mutations on CMPI and NS9283 potentiation of $\alpha_4^3\beta_2^2$ nAChR current elicited by saturating concentrations of ACh and TC 2559

<table>
<thead>
<tr>
<th>Agonist</th>
<th>Subunits combination</th>
<th>+1 μM CMPI</th>
<th>+1 μM NS9283</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Ave ± SD</td>
<td>$P_{\text{versus control}}$</td>
</tr>
<tr>
<td>TC 2559 (30 μM)</td>
<td>[α4]3[β2]2 WT</td>
<td>6.41 ± 2.15</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>[α4]K64T[β2]2</td>
<td>1.17 ± 0.21</td>
<td>1.000</td>
</tr>
<tr>
<td></td>
<td>[α4]E66I[β2]2</td>
<td>0.99 ± 0.15</td>
<td>1.000</td>
</tr>
<tr>
<td></td>
<td>[α4]H116L[β2]2</td>
<td>4.58 ± 0.24</td>
<td>0.004</td>
</tr>
<tr>
<td></td>
<td>[α4]H116V[β2]2</td>
<td>3.01 ± 0.39</td>
<td>0.518</td>
</tr>
<tr>
<td></td>
<td>[α4]H116A[β2]2</td>
<td>0.74 ± 0.13</td>
<td>1.000</td>
</tr>
<tr>
<td></td>
<td>[α3]3[β2]2 WT</td>
<td>0.82 ± 0.08</td>
<td>1.000</td>
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<tr>
<td></td>
<td>[α4]K64T[β2]2</td>
<td>0.83 ± 0.10</td>
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</tr>
<tr>
<td></td>
<td>[α4]E66I[β2]2</td>
<td>0.82 ± 0.05</td>
<td>1.000</td>
</tr>
<tr>
<td></td>
<td>[α4]H116L[β2]2</td>
<td>0.76 ± 0.05</td>
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<tr>
<td></td>
<td>[α4]H116A[β2]2</td>
<td>1.07 ± 0.13</td>
<td>1.000</td>
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</tbody>
</table>

The data from Figure 5D reporting current responses to agonist, agonist + 1 μM CMPI, and agonist + 1 μM NS9283 of oocytes expressing WT and mutants [α4]3[β2]2 nAChRs was analyzed using one-way ANOVA with multiple comparisons versus control group (Holm–Sidak method, SigmaPlot, Systat Software Inc.). Shown in the table are the probability ($P_{\text{versus control}}$) that calculated potentiation folds in the presence of 1 μM CMPI or 1 μM NS9283 differ from no potentiation and the probability ($P_{\text{versus WT}}$) that calculated potentiation folds in the presence of 1 μM CMPI or 1 μM NS9283 for [α4]3[β2]2 nAChRs containing the indicated amino acid mutation differ from the calculated potentiation fold for WT [α4]3[β2]2 nAChRs.
presence of multiple active compounds, each drug independently and energetically additively contributes to the stabilization of the active state.

We commenced by converting the raw amplitudes of current responses to units of probability of being in the active state (\(P_A\) units). The constitutive probability of being active (\(P_{A,\text{constitutive}}\)) and peak \(P_A\) of the response to 1 mM ACh in the \((\alpha 4)3(\beta 2)2\) receptor were estimated by comparing the current responses to 100 \(\mu\)M mecamylamine, 1 mM ACh, and 1 mM ACh + 3 \(\mu\)M desformylIstrabrinobromine (dFBr). The underlying assumption in this approach (36) is that the application of the blocker mecamylamine inhibits constitutively-active receptors, thereby revealing the current level corresponding to \(P_A \sim 0\), whereas the coapplication of ACh and the allosteric activator dFBr activates all receptors in the membrane and reveals the current level with \(P_A \sim 1\). The application of mecamylamine elicited outward current with the mean amplitude of 2.1 ± 0.5% of the absolute response to 1 mM ACh, whereas dFBr potentiated the peak response to 1 mM ACh to 278 ± 36% of control. From this, we estimate a \(P_{A,\text{constitutive}}\) of 0.00755 ± 0.00118 and a \(P_{A,1 \text{ mM ACh}}\) of 0.36 ± 0.05.

The concentration-response relationships for ACh, cytisine, and nicotine were fitted to Equation 2. With \(L\) constrained to 220 (calculated as \((1 - P_{A,\text{constitutive}})/P_{A,\text{constitutive}}\)), the fitting yielded a \(K_{R,\alpha 4\beta 2}\) (equilibrium dissociation constant of ACh at the \(\alpha 4\beta 2\) site in the resting receptor) of 1.32 ± 0.35 \(\mu\)M (best-fit parameter ±SD of the fit) and a \(c_{ACh,\alpha 4\beta 2}\) (ratio of the equilibrium dissociation constant of ACh at the \(\alpha 4\beta 2\) site in the active receptor to \(K_{R,\alpha 4\beta 2}\)) of 0.239 ± 0.014, and a \(K_{R,\alpha 4\beta 2}\) of 244 ± 25 \(\mu\)M and a \(c_{ACh,\alpha 4\alpha 4}\) of 0.117 ± 0.013 (affinity and efficacy parameters of ACh, respectively, at the \(\alpha 4\alpha 4\) site). Thus, the binding of transmitter to the two \(\alpha 4\alpha 4\) sites and the single \(\alpha 4\alpha 4\) site contributes −1.69 and −1.27 kcal/mol, respectively, toward stabilization of the active state. Note that a lower value of \(c\) is associated with higher efficacy and that the single \(\alpha 4\alpha 4\) site in the presence of ACh contributes nearly as much as the combined two \(\alpha 4\beta 2\) sites in free energy change. Fitting of the cytisine concentration-response curve to Equation 2 yielded a \(K_{R,\text{cytisine,}\alpha 4\beta 2}\) of 0.63 ± 1.21 \(\mu\)M, a \(c_{\text{cytisine,}\alpha 4\beta 2}\) of 0.875 ± 0.029, a \(K_{R,\text{cytisine,}\alpha 4\alpha 4}\) of 8.3 ± 0.5 \(\mu\)M, and a \(c_{\text{cytisine,}\alpha 4\alpha 4}\) of 0.155 ± 0.104. Fitting of the nicotine concentration-response curve gave a \(K_{R,\text{nicotine,}\alpha 4\beta 2}\) of 0.08 ± 0.05 \(\mu\)M, a \(c_{\text{nicotine,}\alpha 4\beta 2}\) of 0.421 ± 0.043, a \(K_{R,\text{nicotine,}\alpha 4\alpha 4}\) of 19 ± 2 \(\mu\)M, and a \(c_{\text{nicotine,}\alpha 4\alpha 4}\) of 0.065 ± 0.013. Thus, at the \(\alpha 4\beta 2\) sites, ACh and cytisine have similar low affinities, whereas the affinity of nicotine is nearly ten-fold higher. At the \(\alpha 4\alpha 4\) site, all three agonists have significantly lower affinity. All three agonists act more efficaciously via the \(\alpha 4\alpha 4\) than a single \(\alpha 4\beta 2\) site. The concentration-response curves are given in Figure 7, and the fitting results are summarized in Table 2.

TC-2559 only interacts with the agonist-binding sites at the \(\alpha 4\beta 2\) interface (30, 34). The concentration-response curve for TC-2559 was fitted to Equation 3, which describes a model with a single class of binding sites. The fitting yielded a \(K_{R,\text{TC-2559,}\alpha 4\beta 2}\) of 0.33 ± 0.05 \(\mu\)M and a \(c_{\text{TC-2559,}\alpha 4\beta 2}\) of 0.371 ± 0.006. Thus, TC-2559 has similar to ACh affinity and efficacy at the \(\alpha 4\beta 2\) sites, and its overall lower gating efficacy (Fig. 7) is explained by its inability to contribute to channel activation via the \(\alpha 4\alpha 4\) site.

CMPI, which binds only to the \(\alpha 4\alpha 4\) interface, is a very weak direct activator of the \(\alpha 4\beta 2\) receptor (28). We therefore
estimated the affinity and gating parameters for CMPI by measuring its effect on the background of activity elicited by a low concentration of TC-2559. Because TC-2559 only interacts with the agonist-binding sites at the α4:β2 interface (30, 34), its activating effect was reflected in a reduced value of $L$ in Equation 3, calculated as $(1 - P_{A,TC-2559})/P_{A,TC-2559}$. Using a dataset obtained in the presence of 1 μM TC-2559 and 0.01 to 3 μM CMPI, we estimate a $K_{R,CMPI,α4:α4}$ of 0.17 ± 0.03 μM and a $c_{CMPI,α4:α4}$ of 0.090 ± 0.004. The binding of CMPI to the α4:α4 interface contributes −1.42 kcal/mol of free energy change toward stabilization of the active state. CMPI is thus as efficacious as ACh at the α4:α4 site, and its overall low efficacy is accounted for by its single binding site.

Additional estimates of the properties of CMPI were obtained by measuring receptor activation by CMPI on the background of activity elicited by 10 μM TC-2559, 10 μM ACh, 1 μM cytisine, or 1 μM nicotine. Fitting the CMPI concentration-response data obtained in the presence of 10 μM TC-2559 to Equation 2 yielded a $K_{R,CMPI,α4:α4}$ of 0.09 ± 0.02 μM and a $c_{CMPI,α4:α4}$ of 0.129 ± 0.009. The data for the combinations of CMPI with ACh, cytisine, or nicotine were analyzed using Equation 4, which describes a model in which CMPI competes with ACh, cytisine, or nicotine, respectively, at the α4:α4 site. Fitting the 10 μM ACh + CMPI data to Equation 4 gave a $K_{R,CMPI,α4:α4}$ of 0.29 ± 0.06 μM and a $c_{CMPI,α4:α4}$ of 0.145 ± 0.008, and the combination of 1 μM...
CMPI pharmacology at (α4)3(β2)2 nAChRs

Table 2
Summary of mechanistic analyses

<table>
<thead>
<tr>
<th>Receptor</th>
<th>Ligand</th>
<th>Site</th>
<th>K_R (μM)</th>
<th>c</th>
<th>K_R,CMPI (μM)</th>
<th>c_CMPI</th>
</tr>
</thead>
<tbody>
<tr>
<td>(α4)3(β2)2 WT</td>
<td>ACh</td>
<td>α4β2</td>
<td>1.32 ± 0.35</td>
<td>0.239 ± 0.014</td>
<td>0.29 ± 0.06</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Cytisine</td>
<td>α4α4</td>
<td>244 ± 25</td>
<td>0.117 ± 0.013</td>
<td>0.145 ± 0.008</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Nicotine</td>
<td>α4β2</td>
<td>0.63 ± 1.24</td>
<td>0.875 ± 0.029</td>
<td>0.43 ± 0.17</td>
<td>0.074 ± 0.009</td>
</tr>
<tr>
<td></td>
<td>TC-2559</td>
<td>α4β2</td>
<td>8.3 ± 0.5</td>
<td>0.155 ± 0.104</td>
<td>0.40 ± 0.10</td>
<td>0.065 ± 0.007</td>
</tr>
<tr>
<td>(α4H116A)3(β2)2</td>
<td>ACh</td>
<td>α4β2</td>
<td>0.08 ± 0.05</td>
<td>0.421 ± 0.043</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>TC-2559</td>
<td>α4β2</td>
<td>19 ± 2</td>
<td>0.065 ± 0.013</td>
<td>0.40 ± 0.10</td>
<td>0.065 ± 0.007</td>
</tr>
<tr>
<td></td>
<td>α4α4</td>
<td>0.33 ± 0.05</td>
<td>0.371 ± 0.006</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
</tbody>
</table>

The table gives the fitted equilibrium dissociation constants for ACh, cytisine, nicotine, and TC-2559 in the resting receptor (K_R) and the ratios of the equilibrium dissociation constants for ACh, cytisine, or nicotine. The different approaches yielded estimates that were within a factor of 2 for K_R,CMPI and c_CMPI. One potential explanation to this relatively large range of estimates is allosteric coupling between the α4β2 and α4α4 sites. This possibility could be tested by comparing receptor activation, individually and in combination, by agents selectively interacting with the α4β2 and α4α4 sites. At present, however, we lack the tools to independently measure activation produced by selective occupation of the α4α4 site.

Discussion

Prior pharmacological studies have identified multiple ligand-recognition sites within the extracellular and transmembrane domains of the (α4)3(β2)2 nAChR including the “canonical” α4β2 and the “noncanonical” α4α4 ABS. The functional response (i.e., channel gating) to ligand occupancy at these recognition sites depends on which site(s) is occupied, the number of site(s) occupied, the allosteric coupling of the occupied site with channel gating, and the intrinsic activity of the ligand at the site(s) it occupies. The small molecule agonists including ACh, nicotine, and cytisine bind with high affinity to the α4β2:2 AChR and with a much lower affinity to the α4α4:2 ACh binding (11, 34). In contrast, larger agonists with a maximum length of >7.5 Å and an accessible surface area of >300 Å (e.g., TC-2559) only bind to the α4β2:2 ACh R (30, 34). Compounds that bind with high affinity at the α4α4:2 ACh-binding site (e.g., CMPI and NS9283) have been identified. CMPI was introduced as the PAM of the (α4)3(β2)2 nAChR as it enhanced channel responses when coapplied with ACh (12, 28). Here, we have characterized the effect of coapplication of CMPI with a series of agonists with different intrinsic activities at the (α4)3(β2)2 nAChR and different binding properties at the α4α4:2 ACh-binding site. CMPI potentiated (α4)3(β2)2 nAChR responses to subsaturating concentrations of all tested agonists, independent of their intrinsic activity or α4α4 binding properties. In contrast, CMPI potentiation of (α4)3(β2)2 nAChR responses induced by a saturating agonist was dependent on the ability of the agonist to bind at the α4α4 ACh-binding site. CMPI enhanced the response to saturating TC-2559 in the WT receptor where TC-2559 only binds to the α4β2 ACh-binding site, but not in the receptor containing the α4H116A mutation that enables the binding of TC-2559 to the α4α4 ACh-binding site. Analysis of the findings in the framework of the MWC model indicates that CMPI binds at the α4α4 interface with higher affinity than ACh, cytisine, or nicotine whereas its gating efficacy at the α4α4 site is equivalent to that of ACh, cytisine, or nicotine. Therefore, the weak direct activating effect of CMPI is accounted for by a single-binding site mediating its action.

We estimated the binding and gating properties of CMPI in the presence of TC-2559 (applied at two concentrations), ACh, cytisine, or nicotine. The different approaches yielded estimates that were within a factor of ~5 for K_R,CMPI and c_CMPI. One potential explanation to this relatively large range of estimates is allosteric coupling between the α4β2 and α4α4 sites. This possibility could be tested by comparing receptor activation, individually and in combination, by agents selectively interacting with the α4β2 and α4α4 sites. At present, however, we lack the tools to independently measure activation produced by selective occupation of the α4α4 site.

Another possible explanation is that one or more of the drugs act through other sites or mechanisms that are not incorporated into the MWC model. Several orthosteric agonists including nicotine show reduced peak response at high agonist concentrations, possibly a result of open-channel blocking mechanism (37). This inhibitory effect, which is not accounted for by our model, may be expected to predominantly affect the fitted K_R and c at the low-affinity α4α4 site. The inhibitory effect is, however, minimized when CMPI-elicited currents are recorded in the presence of low concentration (1 μM) of nicotine. Finally and, what we consider, most plausible is that the differences in estimated K_R,CMPI and c_CMPI reflect simple experimental imprecision and variability in receptor behavior. This is supported by the finding that a change in the concentration of background TC-2559 from 1 to 10 μM generates an almost two-fold change in estimated K_R,CMPI.

The estimated activation parameters presented here are dependent on the accurate measurement of the constitutive P_A of the α4β2 receptor and its peak P_A in the presence of ACh.
CMPI pharmacology at (α4)3(β2)2 nAChRs

We used mecamylamine to block constitutively active receptors and reveal the current level corresponding to a PA of 0. Mecamylamine is a nonselective, allosteric antagonist of the nAChR with IC50 in the submicromolar range (38). It acts by blocking open receptors (39). We measured the effect of 100 μM mecamylamine on holding current to estimate the current level at PA of 0. Underestimation of the effect of mecamylamine on holding current would lead to under-estimated PA,constitutive and an overestimated L (Equation 2). This would introduce an error in the estimated values of c for ACh, cytisine, TC-2559, and nicotine. The extent of error can be calculated from the relationship Ltrue × ctrue = Lestimated × cestimated. An underestimated PA,constitutive is not expected to lead to a meaningful error in the activation parameters for CMPI or the peak PA,ACh values.

The peak PA,ACh (0.36) was estimated by normalizing the peak response to 1 mM ACh to that in the presence of 1 mM ACh + 3 μM dFBr. dFBr is a brominated alkaloid, originally isolated from the marine bryozoan Flustra foliacea (40) that selectively and allosterically potentiates the α4β2 nAChR (41). In our hands, 3 μM dFBr almost tripled the peak response to 1 mM ACh. We have assumed that the response to ACh + dFBr has a peak PA indistinguishable from 1. An underestimated potentiating effect of dFBr would lead to over-estimated peak PA for ACh. This, in turn, would lead to proportional errors in estimated c for each of the tested ligands. However, the estimated relative contributions made by α4β2 and α4α4 sites would remain unaffected. The previous studies have reported a peak PA,ACh of 0.5 to >0.8 in the (α4)3(β2)2 nAChR (30, 42). Indurthi et al. (30) used an approach similar to ours, observing doubling of the peak response to 1 mM ACh in the presence of the allosteric modulator NS206, whereas Li and Steinbach (42) employed nonstationary noise analysis on human embryonic kidney cells stably expressing the α4β2 receptor.

A previous study reported that pharmacological elimination of the α4α4 site in the (α4)3(β2)2 nAChR reduced the subsequent response to saturating ACh to ~40% of the control response (34). The fraction of the high-affinity component in the ACh concentration-response relationship remains at ~15% even when the α4 subunit is expressed in excess (11), indicating that occupation of the α4β2 sites by ACh generates a functional response that is 15 to 40% of the response to saturating ACh in the (α4)3(β2)2 receptor. This is in good agreement with the data presented here. Using the KΔG,ACh and cACh values in Table 2, we calculate, using Equation 2, that occupation of the two α4β2 sites in the (α4)3(β2)2 receptor with ACh generates a peak PA of 0.072, whereas occupation of the two α4β2 sites and the single α4α4 site with ACh generates a peak PA of 0.36. We emphasize that in either case, ACh should be considered a partial agonist of the (α4)3(β2)2 nAChR given its relatively low maximal PA.

In sum, we report here that ACh binds with high affinity (KΔG,ACh,α4β2 = 1.32 μM) to the α4β2 agonist-binding sites where it acts with relatively low efficacy (cACh,α4β2 = 0.239; ΔGgating,total = −1.69 kcal/mol or −0.84 kcal/mol per site). ACh binds with low affinity (KΔG,ACh,α4α4 = 244 μM) to the α4α4 site where it acts with relatively high efficacy (cACh,α4α4 = 0.117; ΔGgating = −1.27 kcal/mol). The nicotinic receptor PAM CMPI has high affinity to the α4α4 site (KΔG,CMPI,α4α4 = 0.28 μM) and efficacy comparable to that of ACh (cACh,α4α4 = 0.101; ΔGgating = −1.35 kcal/mol). CMPI enhances channel-gating activation triggered by ACh occupancy at the α4β2 agonist-binding sites by binding to the α4α4 subunit interface which becomes occupied by ACh only at high concentrations. Overall, these results indicate that exposure to agonists targeting the α4α4 binding site in the (α4)3(β2)2 nAChR is expected to increase the efficacy of the transmitter ACh, that may be therapeutically beneficial in conditions associated with decline in the output of nAChR in the brain.

Experimental procedures

Materials

Acetylcholine chloride was purchased from Sigma-Aldrich. Other ligands of the nAChR (CMPI, NS9283, dFBr, TC-2559, cytisine, and mecamylamine) were from Tocris Bioscience R&D. Collagenase type 2 was from Worthington Biomedical. The stock solutions were prepared for ACh (1 M in water) and other nAChR ligands (10 mM in water or DMSO) and stored in aliquots at −20 °C until used. The final working solutions were prepared in recording buffer on the day of experiments.

Expression of (α4)3(β2)2 and (α4)2(β2)3 nAChRs in Xenopus oocytes

Oocytes-positive female Xenopus laevis were purchased from NASCO, and all procedures were performed according to an animal use protocol approved by the Institutional Animals Care and Use Committee of The University of Texas Health Science Center at Tyler. Ovarian lobules were surgically harvested, treated with collagenase type 2, and Stage V and VI oocytes were visually selected and maintained at 18 °C in modified ND96-gentamicin buffer (96 mM NaCl, 2 mM KCl, 1.8 mM CaCl2, 1 mM MgCl2, 5 mM Hepes, and 50 μg/ml gentamicin, pH 7.6).

pSP64 Poly(A) plasmids with cDNA encoding for human α4 or β2 nAChR subunit were used to prepare cRNA transcripts suitable for oocyte expression. The plasmids were linearized with Asel (hα4) and PvuII (hβ2), then cRNA transcripts were prepared in vitro using mMESSAGE mMACHINE high yield capped RNA transcription kits (Ambion, Thermo Fisher Scientific), purified on NucAway Spin column (Invitrogen, Thermo Fisher Scientific), and stored at −80 °C until used. Point mutations within the plasmid encoding the α4 nAChR subunit were introduced using QuikChange II Site-Directed Mutagenesis Kit (Agilent Technologies), as described previously (28). To generate amino acid substitutions (K64T, E66L, H116V, H116L, and H116A), two custom-designed complementary oligos containing the desired mutation were used (Integrated DNA Technologies). The forward primers were as the following with the codon for mutated amino acids are underlined and nucleotide(s) changes are bolded and italicized:
two-electrode voltage-clamp recordings

Two-electrode voltage-clamp recordings of ACh- or TC-2559-induced responses of Xenopus oocytes were performed, as described in (28). 24 to 72 h after cRNA injection, Xenopus oocytes were placed in a custom-made recording chamber that is connected to an eight-channel automated perfusion system (Warner Instruments) and perfused with recording buffer (100 mM NaCl, 2 mM KCl, 1 mM CaCl2, 0.8 mM MgCl2, 1 mM EGTA, and 10 mM Hepes, pH 7.5). Unless otherwise specified in figure legends, each recording run included several drug applications (10 s of an agonist with or without CMPI or NS9283) separated by 3 to 4 min buffer wash intervals. Between recording runs, the oocytes were washed with recording buffer for at least 5 min. The oocytes were voltage-clamped at −50 mV using Oocyte Clamp OC-725B (Warner Instruments). The currents were digitized using Digidata 1550A (Axon Instruments, Molecular Devices), and the peak currents were quantified using pCLAMP 10 (Axon Instruments), then normalized and analyzed using Excel 2010 (Microsoft) and SigmaPlot 11.0 (Systat Software). For NS9283 and CMPI potentiation of agonist-induced responses, the peak currents were normalized to current elicited by agonist alone applied within the same recording run. For the effect of copl application of 1 μM NS9283 or CMPI on agonist concentration-response curve, the peak currents were normalized to current elicited by saturating concentration of agonist applied within the same recording run. Mean ± SD of N oocytes were plotted and fit to the following equation:

\[ I_X = I_0 + \frac{I_{max}}{1 + \left( \frac{EC_{50}}{X} \right)^h} \]  (1)

where \( I_X \) is the normalized agonist-induced current in the presence of NS9283 or CMPI at concentration \( x \), \( I_{max} \) is the maximum potentiation of current, \( h \) is the Hill coefficient, and \( EC_{50} \) is the of NS9283 or CMPI concentration producing 50% of maximal potentiation. \( I_0 = 100 \) for NS9283 and CMPI potentiation of agonist-induced responses and \( I_0 = 0 \) for agonist concentration-response experiments. The best-fit values for \( I_{max} \) and \( EC_{50} \) ± SD are presented.

For the enhancement of agonist-induced currents by 1 μM CMPI or NS9283 (data in Figs. 1C and 5D), the probability (P) that the calculated potentiation fold differ from no potentiation (potentiation fold = 1) or from WT (α4β2) nAChR (potentiation fold of 6.21 and 5.05 for CMPI or NS9283, respectively) was analyzed using one-way analysis of variance with Holm–Sidak post hoc test (SigmaPlot, Systat Software Inc) and reported in the legend for Figure 1C and Table 1.

Mechanistic analysis

Further analysis of electrophysiological data was conducted in the framework of the two-state concerted transition model, adapted from the MWC cyclic model originally used to describe enzyme function (31–33). The raw peak amplitudes of current responses were converted to units of probability of being in the active state (P). We used a multi-step approach where the peak P to 1 mM ACh was estimated through normalization to the peak response to 1 mM ACh +3 μM dFBr. Additional normalization was carried out by comparing responses to various agonists or agonist combinations to the peak response to 1 mM ACh in the same set of cells. The \( P_A \) of constitutive activity (\( P_A,\text{constitutive} \)) was estimated by comparing the effects of 100 μM mecamylamine and 1 mM ACh on the holding current.

The (α4)3(β2)2 receptor contains two binding sites for ACh at the α4β2 intersubunit interface and one site at the α4α4 interface (11). The same set of sites has also been shown to mediate receptor activation by the alkaloid cytisine (34). The concentration-response curves for ACh, cytisine, and nicotine were fitted to the state function:

\[ P_A = \frac{1}{1 + L} \left[ \frac{1 + [\text{agonist}]/K_{\text{agonist},\alpha/\beta}^2}{1 + [\text{agonist}]/(K_{\text{agonist},\alpha/\beta}^2 C_{\text{agonist},\alpha/\beta})} \right] \]  (2)

where \( L \) indicates the level of background activity in the absence of agonist and is calculated as \((1 - P_A,\text{constitutive})/P_A,\text{constitutive}\). [agonist] is the concentration of ACh, cytisine, or nicotine, \( K_{\text{agonist},\alpha/\beta} \) and \( K_{\text{agonist},\alpha/\alpha} \) are the equilibrium dissociation constant for the agonist in the resting receptor at the α4-β2 or α4-α4 sites, respectively, and \( C_{\text{agonist},\alpha/\beta} \) and \( C_{\text{agonist},\alpha/\alpha} \) are the ratios of the equilibrium dissociation constants for the agonist in the active receptor to that in the resting receptor. The numbers of α4β2 and α4α4 binding sites \((N_{\alpha/\beta}^2 \text{and } N_{\alpha/\alpha}^2) \) were constrained to 2 and 1, respectively.

TC-2559 activates the (α4)3(β2)2 receptor by binding to the two sites at the α4β2 interface. The concentration-
**CMPI pharmacology at (α4)3(β2)2 nAChRs**

response curve for TC-2559 was fitted to the following equation:

$$\frac{P_A}{1 + L} = \frac{1}{1 + \left(\frac{TC}{K_{R,TC-2559,α4β2}}\right)^{N_{α4β2}}}$$

(3)

The terms are as described above.

CMPI interacts with the agonist-binding site at the α4-α4 interface. Because it is a weak agonist, its affinity and gating properties at the α4-α4 site were estimated by coapplying CMPI with a fixed, low concentration of TC-2559. The concentration-response data were analyzed using Equation 3, with the value of L modified to reflect receptor activation by TC-2559, and the affinity and efficacy terms in the equation reflecting the values for CMPI. The number of binding sites for CMPI in Equation 3 was constrained to 1.

For the combinations of ACh, cytisine or nicotine, and CMPI we assumed that ACh, cytisine, and nicotine are the sole ligands at the two α4-β2 sites, whereas CMPI competes with ACh, cytisine, or nicotine at the α4-α4 site. The concentration-response curves for the combinations of ACh, cytisine or nicotine, plus CMPI were fitted to the following equation:

$$\frac{P_A}{1 + L} = \frac{1}{1 + \left(\frac{[\text{agonist}]/K_{R,[\text{agonist},α4β2]} + [\text{CMPI}]/K_{R,[\text{CMPI},α4β2]}\right)^{N_{α4β2}}} + \left(\frac{[\text{agonist}]/K_{R,[\text{agonist},α4β2]} + [\text{CMPI}]/K_{R,[\text{CMPI},α4β2]}\right)^{N_{α4β2}}}$$

(4)

where $K_{R,CMPI}$ is the equilibrium dissociation constant for CMPI in the resting receptor at the α4α4 site, $c_{CMPI}$ is the ratio of the equilibrium dissociation constants for CMPI in the active receptor to $K_{R,CMPI}$, and other terms are as described above.

Curve fitting was done using Origin 2020 (OriginLab Corp). The results are reported as best-fit parameter ± SD of the fit. All data are included in the analysis.

**Data availability**

All data are contained in the article.


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**Conflict of interest**—The authors declare that they have no conflicts of interest with the contents of this article.

**Abbreviations**—The abbreviations used are: ABS, agonist binding site; ACh, acetylcholine; CMPI, 3-(2-chlorophenyl)-5-(5-methyl-1-(piperidin-4-yl)-1H-pyrazol-4-yl)sloxazole; dFBr, desformyl-flusilatrobrone; MWC, Monod-Wyman-Changeux; nAChR, nicotinic acetylcholine receptor; PAMs, positive allosteric modulators.

**References**

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