The novel $\alpha_7\beta_2$-nicotinic acetylcholine receptor subtype is expressed in mouse and human basal forebrain: Biochemical and pharmacological characterisation

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Non-standard abbreviations: α-Bgtx, α-bungarotoxin; α-Cbtx, α-cobratoxin; AD, Alzheimer’s disease; C C4, 1,2-bis-N -cystinylethane; DHβE, dihydro-β-erythroidine; E pi, epibatidine; MECA, mecamylamine; MLA, methyllycaconitine; nAChR, nicotinic acetylcholine receptor(s); PMSF, phenylmethylsulphonylfluoride; SDS-PAGE, sodium dodecyl sulphate-polyacrylamide gel electrophoresis; SN/VTA, substantia nigra / ventral tegmental area; TBS, Tris-buffered saline;
TEVC, two-electrode voltage-clamp.
ABSTRACT:
We examined α7β2-nicotinic acetylcholine receptor (α7β2-nAChR) expression in mammalian brain and compared pharmacological profiles of homomeric α7-nAChR and of α7β2-nAChR. α-bungarotoxin affinity purification or immunoprecipitation with anti-α7 subunit antibodies (Abs) were used to isolate nAChR containing α7 subunits from mouse or human brain samples. α7β2-nAChR were detected in forebrain, but not other tested regions, from both species, based on western blot analysis of isolates using β2 subunit-specific Abs. Ab specificity was confirmed in control studies using subunit-null mutant mice or cell lines heterologously expressing specific, human nAChR subtypes and subunits. Functional expression in Xenopus oocytes of concatenated pentameric (α7)5-, (α7)4(β2)1-, and (α7)3(β2)2-nAChR was confirmed using two-electrode voltage-clamp recording of responses to nicotinic ligands. Importantly, pharmacological profiles were indistinguishable for concatenated (α7)5-nAChR or for homomeric α7-nAChR constituted from unlinked α7 subunits. Pharmacological profiles were similar for (α7)5-, (α7)4(β2)1-, and (α7)3(β2)2-nAChR except for diminished efficacy of nicotine (normalized to acetylcholine efficacy) at α7β2- vs. α7-nAChR. This study represents the first direct confirmation of α7β2-nAChR expression in human and mouse forebrain, supporting previous mouse studies that suggested relevance of α7β2-nAChR in Alzheimer’s disease etiopathogenesis. These data also indicate that α7β2-nAChR subunit isoforms with different α7:β2 subunit ratios have similar pharmacological profiles to each other, and to α7 homopentameric nAChR. This supports the hypothesis that α7β2-nAChR agonist activation predominantly or entirely reflects binding to α7/α7 subunit interface sites.
INTRODUCTION

Several nicotinic acetylcholine receptor (nAChR) subtypes are expressed widely along the entire neuraxis, and are involved in many of the physiological functions of the central and peripheral nervous systems (Albuquerque et al., 2009; Hurst et al., 2013). nAChR activity controls important aspects of synaptic function and brain development, including the proliferation and differentiation of neural progenitors, neural migration, and neuronal maturation (Griguoli and Cherubini, 2012; Picciotto et al., 2012; Yakel, 2013). Furthermore, nAChR dysfunction may play an important role in a variety of neurological diseases including neurodegenerative and psychiatric diseases (Gotti and Clementi, 2004; Lewis and Picciotto, 2013).

\[ \alpha_4\beta_2 \] and homomeric \[ \alpha_7\text{-nAChR} \] are the most widely-expressed subtypes in mammalian brain. The latter are thought to contain five identical agonist binding sites located at subunit interfaces in extracellular domains (Gotti and Clementi, 2004; Whiteaker et al., 2007). Pharmacological hallmarks of \[ \alpha_7\text{-nAChR} \] are their high sensitivity to antagonism by snake venom-derived polypeptide toxins such as \[ \alpha\text{-bungarotoxin (\(\alpha\text{-Bgtx} \))} \] and \[ \alpha\text{-cobratoxin (\(\alpha\text{-Cbtx}) \)}, \] and their sensitivity to choline (a product of ACh hydrolysis) as an agonist (Albuquerque et al., 1997; Albuquerque et al., 2009). \[ \alpha_7\text{-nAChR} \] are highly expressed in the cortex, hippocampus and subcortical limbic regions, and (at lower levels) in the thalamus and basal ganglia. \[ \alpha_7\text{-nAChR} \] that are located on or near nerve terminals are involved in control of neurotransmitter release, whereas \[ \alpha_7\text{-nAChR} \] on dendrites or soma apposed to cholinergic synaptic endings play roles in classic neurotransmission. In both cases, \[ \alpha_7\text{-nAChR} \]’s high calcium permeability may also result in altered intracellular signalling and gene transcription (Albuquerque et al., 2009; Dajas-Bailador and Wonnacott, 2004). \[ \alpha_7\text{-nAChR} \] also may be associated with extrasynaptic volume transmission (Lendvai and Vizi, 2008).

Affinity purification of nAChR using snake-venom \(\alpha\)-toxins has been performed from brain tissue of various species. Extracts from whole rat brain appear to be predominantly
composed of homomeric α7-nAChR (Drisdel and Green, 2000). However homomeric α7- and α8-nAChR (and heteromeric α7α8-nAChR) have been identified in chick CNS extracts (Gotti et al., 1994; Keyser et al., 1993). Further, studies using heterologous systems have shown that α7 subunits can form functional channels when combined with α5 (Girod et al., 1999), β2 (Khiroug et al., 2002), β3 (Palma et al., 1999) or β4 subunits (Criado et al., 2012). Fluorescently tagged nAChR α7 and β2 subunits have recently been used to characterize the formation of α7β2-nAChR, and functional differences between α7- and α7β2-nAChR have been suggested (Murray et al., 2012). Co-expression of β2 and α7 subunits caused a significant decrease in agonist-stimulated whole cell current amplitudes, but this decrease occurs without affecting the concentration-response characteristics of a range of common agonists and antagonists (Murray et al., 2012). Other studies have shown that α7 and β2 subunits are co-expressed in rat basal forebrain cholinergic neurons and appear to form heteromeric α7β2-nAChR with subtly different biophysical and pharmacological properties from those of homomeric α7-nAChR (Liu et al., 2009). In addition, interaction of these putative α7β2-nAChR with oligomeric forms of amyloid-β (Aβ1-42) may be relevant in the etiology of Alzheimer’s disease (Liu et al., 2013).

These previous studies suggest that the function and pharmacology of α7*-nAChR (where * denotes the known or possible presence of other nAChR subunits than α7 (Lukas et al., 1999)) may be more complex than previously thought, and that α7β2-nAChR expression may be restricted to forebrain areas. However, heteromeric α7*-nAChR have not yet been directly detected biochemically, nor have they been definitively identified in human brain. We used the α7-nAChR-selective ligand, α-Bgtx, to affinity purify α7*-nAChR from selected brain areas of humans or of wildtype (WT) or β2 subunit-null mutant (KO) mice. The subunit compositions of these isolated α7*-nAChR were analyzed by western blot analysis using subunit-specific anti-α7 or β2 antibodies. The results suggest how expression of α7β2-nAChR in both WT mouse and human forebrain samples, but not in brains from β2 KO mice. Moreover, concatemeric (linked subunit) constructs, the Xenopus oocyte system, and two-
electrode voltage-c lamp recording were used to confirm functional expression of α7β2-nAChR. This work defined α7 and β2 subunit stoichiometries that enable α7β2-nAChR function, and showed similar pharmacological characteristics across α7- and α7β2 nAChR subtypes. The results confirm commonalities in expression of α7β2-nAChR in man and mouse, and support hypotheses linking α7β2-nAChR, cholinergic signaling loss, and roles for Aβ1-42 in etiopathogenesis of at least a subset of human dementias.
MATERIALS and METHODS

Animals and Materials

The study involved the use of 4-6 month old male, pathogen-free, C57BL/6 wildtype (WT), α7 KO or β2 KO (Orr-Urtreger et al., 1997; Picciotto et al., 1995) mice obtained from Dr. U. Maskos (Pasteur Institute, Paris). All animal experiments were conducted in accordance with the European Community Council Directive (86/609/EEC) of 24 November 1986.

(±)-[3H]-epibatidine (Epi, specific activity, 6 6Ci/mmol) and [125I]-α-Bgtx (specific activity of 200-216 Ci /mmol) were purchased from Perkin Elmer (Waltham, MA, USA). Non-radioactive α-Bgtx, Epi, and nicotine were purchased from Tocris Bioscience (Bristol, UK or Minneapolis, MN, USA), as were dihydro-β-erythroidine (DHE), and methyllycaconitine (MLA). Sazetidine-A (also known as MOP-H-OH) was kindly supplied by Dr. Alan Kozikowski (University of Illinois at Chicago, Chicago, IL, USA). 1,2-bis N cytisnylethane (CC4) also was used (Riganti et al., 2005). α-Cobratoxin (α-Cbtx) and all other reagents were sourced from Sigma-Aldrich unless otherwise specified (St. Louis, MO, USA).

Human tissues

Human cerebellum was provided by the Newcastle Brain Tissue Resource on the basis of a collaboration with Dr Jennifer Court (Newcastle upon Tyne, General Hospital, UK). Samples were all collected by the Brain Tissue Resource with informed consent and appropriate ethical approval. Case details are shown in Table 1; the approvals and method for categorizing the subjects' smoking status are outlined in the methods section of Court et al., 2005. Human basal forebrain tissue was provided by Dr. Emanuele Sher (Lilly Research Center, Windlesham, Surrey, UK), and was also collected with appropriate informed consent in accordance with all applicable laws and regulations.

Transfected cells

Human α2, α3, β2, and β4 nAChR subunit clones in the mammalian expression vector pcDNA3 were kind gifts of Dr. Sergio Fucile (University of Rome, Rome, Italy). The
human α7 nAChR subunit clone in pcDNA3 was a generous gift of Dr. Roberta Benfante (CNR Institute of Neuroscience, Milan, Italy). HEK293 and SH-SY5Y cells were transiently transfected using the Ca₃(PO₄)₂ method or the Jet-PEI reagent (Polyplus, Euroclone, Italy) transfection. For the α7 plasmid, 1.5x10⁶ cells were transfected with 6 μg of plasmid using the Jet-PEI. For each of the α2, α3, α4 and β2 or β4 subunit 20 μg of plasmids for 1.5x10⁶ cells was used, with the Ca₃(PO₄)₂ method. nAChR expression by cells was analyzed 24 h after transfection.

**Antibody Production and Characterization**

We used affinity-purified, subunit-specific, polyclonal antibodies (Abs), produced in rabbit against peptides derived from the C-terminal (COOH) or intracytoplasmic loop (CYT) of human or mouse nAChR subunit sequences, as previously described (Gotti et al., 2006; Grady et al., 2009). The Ab against the COOH peptide (SAPNFVEAVSKDFA) was used for α7 subunits in mouse and human tissues. Abs directed against the α7 mouse CYT peptide (PSGDPDLAKILEEVRYIANRFRC) or the human C YT peptide (QMQEADISGYIPNGQMQEADISGYIPNG) were used for mouse and human tissues, respectively. For the β2 subunit, we used antibodies directed against two different cytoplasmic human β2 peptides: RQRERGAGALFFREAPGADSCTY (β2(1)) and cglADHMRSEDDDDQSVREDWKYV (β2(2)).

The specificity of the affinity-purified Abs was tested by immunoprecipitation studies using α7 WT or α7 KO hippocampus and β2 WT or β2 KO mouse cortex (the results are shown in Supplementary Figure 1). The same Abs also were tested by means of western blotting (Supplementary Figure 1). In order to exclude any cross-reactivity between nAChR subunits, anti-β2 (1)- or anti- α7 human subunit Abs were also tested by means of immunoprecipitation studies and western blotting in HEK293 cells transfected to express human α2β4-, α4β2-, α4β4, or α3β4-nAChR subtypes or in SH-SY5Y cells transfected to express human α7-nAChR (see above) (the results are shown in Supplementary Figure 2).

**Purification of α-bungarotoxin-binding nAChR**
For studies using mice, ≈100 mg of basal forebrain or hippocampus tissue micro-dissected from either WT or subunit-null mice were pooled in every experiment. The tissue was homogenised in 10 ml of 50 mM Na phosphate, pH 7.4, 1 M NaCl, 2 mM EDTA, 2 mM EGTA and 2 mM phenylmethylsulfonylfluoride (PMSF; to covalently inactivate serine protease activity), and the homogenates were diluted and centrifuged for 1.5 h at 60,000g. The entire membrane homogenisation, dilution and centrifugation procedure was then repeated, and the resulting pellets were collected, rapidly rinsed with 50 mM Tris HCl, pH 7, 120 mM NaCl, 5 mM KCl, 1 mM MgCl₂, 2.5 mM CaCl₂ and 2 mM PMSF. The washed pellets were then resuspended in 2 ml of the same buffer, further supplemented with 20 µg/ml of each of the following protease inhibitors: leupeptin, bestatin, pepstatin A and aprotinin. Triton X-100 at a final concentration of 2% was added to the washed membranes, which were extracted for 2 h at 4°C. The extracts were centrifuged for 1.5 h at 60,000g, recovered, and an aliquot of the supernatants was collected for protein measurement using the BCA protein assay (Pierce Biotechnology, Inc., Rockford, IL, USA), with bovine serum albumin as the standard. Extracts (2 ml) were incubated with 200 µl of Sepharose-α-Bgtx (concentration of coupled toxin 1 mg/ml of gel) and shaken overnight at 4°C. The following day, the beads were centrifuged, the supernatant was recovered, and the resins were washed 4-6 times by resuspension followed by centrifugation. After washing, the Sepharose-α-Bgtx beads with bound nAChR (purified α-Bgtx-binding receptors) were incubated with one-two volumes of Laemmli sample buffer (125 mM Tris phosphate, 4% SDS, 20% glycerol, 0.02% bromophenol blue and 10% 2-mercaptoethanol pH 6.8) and boiled for 2 min. The supernatant was then recovered by centrifugation.

In the case of human tissue, α-Bgtx-binding sites were purified using the same procedure as that used for mouse tissue, starting from 600 mg of tissue (See Table 1 for subject details).

**Binding studies**

[^125I]-α-Bungarotoxin
The binding of $[^{125}\text{I}]-\alpha$-Bgtx to 2% Triton X-100 extracts of mouse tissues was determined by collection onto DEAE-Sepharose™ Fast Flow (GE Healthcare, Uppsala, Sweden). Triton extracts (250 µl) from each experimental group were incubated overnight with a saturating concentration (5 nM) of $[^{125}\text{I}]-\alpha$-Bgtx at 20°C in the presence of 2 mg/ml bovine serum albumin. Specific radioligand binding was defined as total binding minus the non-specific binding determined in the presence of 1 µM unlabeled $\alpha$-Bgtx. Non-specific binding averaged 30-40% of total binding. Binding to $\alpha^7$-nAChR could also be measured in an immunoprecipitation assay format. Receptor extracts were labeled with $[^{125}\text{I}]-\alpha$-Bgtx (5 nM in the presence or absence of 1 µM unlabeled $\alpha$-Bgtx to define total and non-specific binding). The labelled extract could then be bound to protein A beads via anti-$\alpha^7$ subunit Abs (described later in Methods). Similar amounts of specific binding were recorded in either assay format, and non-specific binding was between 10-15% of total binding.

$[^{3}\text{H}]$-Epibatidine.

Binding of $[^{3}\text{H}]$-epibatidine to nAChR in 2% Triton X-100 brain tissue extracts obtained was also assessed. $[^{3}\text{H}]$-Epibatidine binds to multiple heteromeric nAChR subtypes with pM affinity and to $\alpha^7$-nAChR with nM affinity. In order to ensure that the $\alpha^7$ nAChR did not contribute to $[^{3}\text{H}]$-Epibatidine binding, in solubilized extracts, binding was performed in the presence of 1 µM $\alpha$-Bgtx, which specifically binds to $\alpha^7$ nAChR (and thus prevents $[^{3}\text{H}]$-Epibatidine binding to these sites).

As for $[^{125}\text{I}]-\alpha$-Bgtx binding assays, binding sites were captured using DEAE-Sepharose™ Fast flow, following overnight incubation of 250 µl aliquots of the extracts with 1 nM $[^{3}\text{H}]$-Epi at 4°C. Non-specific binding (averaging 5-10% of total binding) was determined in parallel samples containing 100 nM unlabelled Epi.

Immunoprecipitation

For immunoprecipitation studies of heteromeric receptors present in human tissues, we used Abs specific for $\alpha_2$, $\alpha_3$, $\alpha_4$, $\alpha_5$, $\beta_2$ or $\beta_4$ subunits directed against human subunit peptides as previously described (Gotti et al., 2006). For $\alpha_6$ and $\beta_3$ subunits, we used Abs
directed against peptides of mouse subunit sequences, also as previously characterized and described (G rady et al., 2009). The immunoprecipitation capacities of the anti-human subunit Abs ranged from 90% to 100% of the [3H]-Epi labeled receptors (mean of three independent experiments). For immunoprecipitation experiments, affinity purified Abs were covalently immobilized on agarose-Protein A beads at a concentration of 4 mg/ml of wet resin. Immunoprecipitation was then performed by adding 20 µl of agarose-Protein A beads with bound, affinity-purified Abs to 200 µl of 1 nM [3H]-Epi-labeled extracts. After overnight incubation, immunoprecipitates were recovered by centrifugation and washed three times with phosphate-buffered saline containing 0.1% Triton X-100.

Immunoblotting and densitometric quantification of western blot bands

nAChR subunit contents of tissue extracts or of α-Bgtx-binding complexes were analysed by western blotting. For the extracts loaded before and after the purification 10 µg of proteins were loaded whereas for the α-Bgtx-purified receptors a constant volume (40 µl), that depending on the tissue, may represent 1/10 or 1/20 of the total recovered Laemmli sample buffer-eluted receptors was loaded onto a 9% acrylamide (Biorad, Hercules, CA, USA) gel and subjected to sodium dodecyl sulphate-polyacrylamide gel electrophoresis (SDS-PAGE). After SDS-PAGE, proteins were electrophoretically transferred to nitrocellulose membranes with 0.45 mm diameter pores (Schleicher and Schuell, Dassel, Germany). The blots were blocked overnight in 5% non-fat milk in Tris-buffered saline (TBS), washed in a buffer containing 5% non-fat milk and 0.3% Tween 20 in TBS, incubated for two hours with the primary antibody (1–2.5 mg/ml), and then incubated with the appropriate peroxidase conjugated secondary Abs (Sigma-Aldrich, St Louis, MO, USA). After 10 washes, peroxidase was detected using a chemiluminescent substrate (Pierce, Rockford, IL, USA). The signal intensity of the Western blot bands was measured using an Epson 4500 gel scanner. The developed films were scanned as a Tiff image in eight-bit gray scale format at a resolution setting of 300 dpi. All of the films obtained from the separate experiments were acquired in the same way and scanned in parallel with a calibrated optical density step tablet.
from Stouffer (Stouffer Graphics Arts, Mishawaka, IN, USA).

The images were analyzed using National Institutes of Health ImageJ software (Schneider et al., 2012). The pixel values of the images were transformed to optical density values by the program using the calibration curve obtained by acquiring the calibrated tablet with the same parameters as those used for the images. The immunoreactive bands were quantified in four separate experiments for the mouse hippocampus and basal forebrain as previously described (Grady et al., 2009)

**Concatameric α7*-nAChR constructs**

Fully-pentameric nAChR concatamers were constructed from human nAChR subunit sequences. cDNAs encoding concatamers were created using the same subunit layout we have previously employed to encode high- and low-agonist-sensitivity α4β2*-nAChR isoforms and α3β4(α5[D/N])-nAChR (Eaton et al., 2014; George et al., 2012). Subunits were arranged in the order α7-α7-α7-α7-α7 (α7 homopentamer), α7-α7-β2-α7-α7 or α7-β2-α7-β2-α7. Kozac and signal peptide sequences were removed from all subunit sequences with the exception of subunits expressed in the first position of the concatamer. Subunits were linked by alanine-glycine-serine (AGS) repeats designed to provide a complete linker length (including the C-terminal tail of the preceding subunit) of 40 ± 2 amino acids. At the nucleotide level, linker sequences were designed to contain unique restriction sites that allow easy removal and replacement of individual α7 and β2 subunits. The protein sequences for the human nAChR subunits were encoded by synthetic nucleotide sequences optimized for expression systems (GeneArt, Life Technologies, Grand Island, NY, USA). Optimization included minimization of high GC content sequence segments, improved codon usage, reduction of predicted RNA secondary structure formation, and removal of sequence repeats and possible alternative start and splice sites. Sequences of all subunits, together with their associated partial linkers, were confirmed by DNA sequencing (GeneArt). Each concatamer was subcloned into the pSGEM oocyte high-expression vector (a kind gift of Dr. Michael Hollmann; Ruhr-Universitaet, Bochum, Germany). For comparison, homomeric α7-nAChR also were expressed from unlinked individual subunits (cDNA clone also synthesized
and optimized by GeneArt). The unlinked human α7 subunit cDNA also was subcloned into the pSGEM vector.

**RNA synthesis**

Plasmids containing concatameric α7-homopentameric or α7β2 nAChR constructs, or individual α7 nAChR subunits, were linearized with NheI (2 hrs at 37 °C), and the reaction mix was treated with proteinase K (30 minutes at 50 °C). cRNAs were transcribed using mMessage mMachine T7 kit (Applied Biosystems/Ambion, Austin, TX, USA). Reactions were treated with TURBO DNase (1U for 15 minutes at 37 °C) and c RNAs were purified using the Qiagen RNeasy Clean-up kit (Valencia, CA, USA). cRNA purity was confirmed on a 1% agarose gel and preparations were stored at -80 °C.

**Xenopus oocytes and RNA injection**

*Xenopus* oocytes were purchased from Ecoyte Bioscience US (Austin, TX) and incubated upon arrival at 13° C. The tips of pulled glass micropipettes were broken to achieve an outer diameter of ~40 μm (resistance of 2-6 MΩ), and pipettes were used to inject 20-60 nl containing 10 ng of cRNA/oocyte. To improve functional expression of α7*-nAChR, Ric-3 mRNA was also co-injected (Halevi et al., 2002). A ratio of 1:50 Ric-3:α7 subunit mRNA by mass was determined to be optimally effective in pilot experiments (data not shown).

**Two-electrode voltage-clamp recording of α7- and α7β2-nAChR function**

Two-electrode voltage-clamp recordings were made at room temperature (20 °C) in oocyte saline (OR2) solution (containing 82.5 mM NaCl, 2.5 mM KCl, 5 mM HEPES, 1.8 mM CaCl2·2H2O, and 1 mM MgCl2·6H2O, pH 7.4). Seven to fourteen days after injection, *Xenopus* oocytes expressing concatenated α7*-nAChR were voltage clamped at −70 mV with an Axoclamp 900A amplifier (Molecular Devices, Sunnyvale, CA, USA). Recordings were sampled at 10 kHz (low-pass Bessel filter: 40Hz; high-pass filter: DC), and the resulting traces were saved to disk (Molecular Devices Clampex v10.2). Data from oocytes with leak currents (Ileak) > 50 nA were excluded from recordings.
Nicotinic receptor pharmacology

Fresh stock drug solutions (agonists: ACh, choline, nicotine, sazetidine and 1,2-bis N cytisineylethane (CC4); antagonists dihydro-β-erythroidine (DHβE), methyllycaconitine (MLA), mecamylamine (MECA) and α-Cbtx) were made daily and diluted as required. Agonists and antagonists were applied using a sixteen channel, gravity-fed, perfusion system with automated valve control (AutoMate Scientific, Inc.; Berkeley, CA, USA). All solutions were supplemented with atropine sulfate (1.5 µM) to ensure that muscarinic ACh receptor responses were blocked and thus not recorded. Oocytes expressing loose subunits and/or concatemeric α7- or α7β2-nAChR were perfused with nAChR agonists for 5 seconds with 60 second washout times between each subsequent application. Oocytes were preincubated with nAChR antagonists for 2 minutes prior to activation with ACh (10 mM; 5 seconds). For experiments using α-Cbtx, bath and drug solutions were supplemented with 0.1% BSA to reduce loss of this peptide ligand by adsorption to the TEVC apparatus.

Data analysis

The expression of [3H]-Epi and [125I]-α-Bgtx receptors and the subunit contents of the [3H]-Epi receptors expressed in the mouse and human samples were statistically compared using unpaired t tests. In human cerebellum samples from smokers and non-smokers, results were compared using an unpaired t test. Statistical analyses were performed using GraphPad Prism 5.0 software (GraphPad Software, Inc., La Jolla, CA, USA).

For TEVC data, EC50 and IC50 values were determined from nAChR-mediated peak currents through non-linear least-squares curve fitting (GraphPad Prism 5.0) using unconstrained, monophasic logistic equations to fit all parameters, including Hill slopes. Desensitization / inactivation of α7*-nAChR currents in the presence of 10 mM (maximally-stimulating) ACh was also analyzed by non-linear least-squares curve fitting in GraphPad Prism 5.0. These data were best fit by a two-phase exponential decay equation. One-way ANOVA was used to compare parameters between multiple groups in each case. Tukey's
multiple comparison test was used for post-hoc analysis in order to compare the means of three or more groups (GraphPad Prism 5.0).
RESULTS

α-Bgtx-binding sites in WT or β2 KO mice

In preliminary experiments, we analyzed nAChR expression in 2% Triton X-100 extracts obtained from the hippocampus or basal forebrain of WT or β2 KO mice (Table 2). By two different approaches (immunoprecipitating [125I]-α-Bgtx-labellel receptors using anti-α7 subunit Abs and by [125I]-α-Bgtx binding to Triton extracts) we determined that the density of α7*-nAChR in mouse hippocampus is more than two times higher than that in murine basal forebrain. During α-Bgtx binding site purification, we determined that 85-95% of those sites in Triton X-100 extracts were bound by α-Bgtx-Sepharose 4B affinity resins, whereas more than 95% of high affinity [3H]-Epi binding was recovered in the flow-through.

The densities of [3H]-epibatidine-binding nAChR were similar between the two regions (Table 2). Elimination of β2 subunit expression dramatically reduced expression of [3H]-epibatidine-binding nAChR in both regions, indicating that this binding is almost entirely due to β2*-nAChR. In contrast, [125I]-α-Bgtx (α7*-nAChR) expression was not significantly different between WT and β2 KO mice in either hippocampus or basal forebrain.

Additionally, western blot analysis was performed on α-Bgtx binding sites affinity purified from the hippocampus of WT or β2 KO mice and probed with anti-α7 (top) or anti-β2 (bottom) subunit Abs (Fig. 1A). Confirming results from the binding studies, western blots also showed no significant differences in presumed α7*-nAChR levels (i.e., polypeptide labeled with anti-α7 subunit Abs) in whole extracts from WT or β2 KO mouse hippocampus (lanes 1 in top two panels). Moreover, affinity purification on α-Bgtx affinity resins isolated comparable levels of α7*-nAChR from WT or β2 KO mouse hippocampus (compare lanes 1 and 3 of Fig. 1A top) but did not isolate nAChR containing β2 subunits (compare lanes 1 and 3 of Fig. 1A, bottom left), which instead were found in the flow-through fraction (lane 2; Fig. 1A bottom left). As expected, no β2*-nAChR were found in extracts isolated on Bgtx resins, or in the flow through from that separation, for tissue taken from β2 KO mice (Fig. 1A lower right panel lanes 1,3 and 2).
In further agreement with the findings of the binding studies, western blot analysis showed that basal forebrain extracts contained fewer α7*-nAChR (i.e., immunoreactive α7 protein) than hippocampal extracts, whereas amounts of immunoreactive β2 subunits was very similar across brain regions. Analysis of the western blots probed using anti-β2 subunit Abs showed clearly-detectable β2 subunit presence in α7*nAChR isolated on Bgtx resins from tissue derived from basal forebrain, but not from the hippocampus (compare lanes 3 of the bottom-left panels of Figs. 1A and 1B). To quantify the % of α7 receptors containing the β2 subunit we loaded on the same gel 10 μg of 2% Triton extract and 1/10 of the α−Bgtx purified receptor and determined by western blotting the optical density of the immunoreactivity of β2 subunit present in the extract (corrected for the total volume used for the receptor purification) and that of the α−Bgtx purified receptor. We found that the immunoreactivity of the β2 subunit determined in the purified α-Bgtx was 2.25 ±0.6 % (n=4) of the total β2 subunit immunoreactivity measured in the basal forebrain extracts of WT mice.

α-Bgtx-binding sites in human brain

The possible presence of α7β2-nAChR in human brain was analysed using post-mortem samples of basal forebrain and cerebellum. In preliminary experiments, we characterised nAChR subtypes expressed in basal forebrain and cerebellum and their levels in 2 % Triton extracts (Table 3). The average level of [125I]-α-Bgtx-labelled (α7*) nAChR was higher in basal forebrain than cerebellum.

The level of [3H]-epibatidine-binding nAChR in cerebellum depended on smoking status. As shown in Table 3 the density of non-α7*-nAChR measured by means of [3H]-Epi binding was higher in smokers than in non-smokers (p=0.02). Based on immunoprecipitation using subunit-specific Abs, in both tissues the large majority of [3H]-Epi-binding sites contained the β2 subunit as associated with the α4 subunit (α4β2-nAChR: 75% in basal forebrain and 60% and 67% in cerebellum of smokers and non-smokers, respectively). An additional 14% of [3H]-Epi-binding sites in the basal forebrain were α2β2*-nAChR whereas
this subtype accounted for only 7% of cerebellar \([^{3}H]\)-Epi-binding sites. The largest region-to-region difference was for \(\alpha 3\beta 2\)-nAChR: whereas those sites accounted for 4.8% of \([^{3}H]\)-Epi-binding sites in the basal forebrain, they represented 32% in the cerebellum.

\(\alpha\)-Bgtx affinity-purified binding sites were obtained from three human basal forebrain (Fig. 2; lanes 1-3) or three human cerebellum (Fig. 2; lanes 5-7) samples. These sites were western blotted and probed with anti-\(\alpha 7\) subunit Abs (top) or two different anti-\(\beta 2\) subunit Abs (anti-\(\beta 2(1)\) Abs (middle) and anti-\(\beta 2(2)\) (bottom)) targeting different epitopes within the \(\beta 2\) subunit. Control samples were extracts from \(\alpha 4\beta 2\)-nAChR-expressing, transfected HEK cells (lane 4) or from \(\alpha 7\)-nAChR-expressing, transfected SH-SY5Y cells (lane 8), and probed with the Abs. Levels of immunoreactivity for the \(\alpha 7\) subunit were very similar in samples loaded in lanes 1, 2, 5 and 6, higher in the sample loaded in lane 3, and lower in the sample loaded in lane 7. Similar isolates from HEK-\(\alpha 4\beta 2\) cells were negative, but SH-SY5Y-\(\alpha 7\) cells contained immunoreactive \(\alpha 7\) subunits (Fig. 2 upper panel, lanes 4 and 8, respectively). Isolation of \(\alpha\)-Bgtx binding sites also yielded anti-\(\beta 2\) subunit A b-labeled proteins from basal forebrain samples but not from the cerebellum, regardless of whether the cerebellum samples were obtained from smokers or non-smokers. Such immunoreactivity was absent in extracts from SH-SY5Y-\(\alpha 7\) cells but very evident in HEK-\(\alpha 4\beta 2\) cells (Fig. 2 middle and lower panels, lanes 8 and 4, respectively). Both the \(\alpha 7\) and \(\beta 2\) subunits present in the human tissues show a slightly higher molecular weight than the corresponding transfected subunits. This is probably due to differences in glycosylation between native and transfected receptors.

Since it has been shown in a heterologous expression system that an \(\alpha 7\beta 4\)-nAChR subtype may be formed (Criado et al., 2012), we also probed human \(\alpha\)-Bgtx-purified sites with anti-\(\beta 4\) subunit Abs with proven specificity (Supplementary Figure 2, bottom). No specific labelling was observed in either the human basal forebrain or cerebellum samples, showing absence of \(\alpha 7\beta 4\)-nAChR. Collectively, these results clearly indicate that \(\alpha 7\beta 2\)-nAChR are present in the human basal forebrain but not in the cerebellum.
Functional expression of concatemeric $\alpha^7*$-nAChR from human subunits

Heterologous expression has shown assembly of functional $\alpha7\beta2^*$-nAChR (see Introduction), but the way(s) in which $\alpha7$ and $\beta2$ subunits might combine from individual, unlinked, subunits could not be defined. Accordingly, we used a linked-subunit approach to produce $\alpha7^*$-nAChR with defined subunit ratios and assembly orders. Each of the three concatemeric constructs [($\alpha7$)$_5$-nAChR homopentamer, ($\alpha7$)$_4$($\beta2$)$_1$-nAChR, and ($\alpha7$)$_3$($\beta2$)$_2$-nAChR] showed concentration dependent ACh-evoked function (representative traces shown in Fig. 3A-D). This function, while smaller than that measured in *Xenopus* oocytes expressing homomeric $\alpha7$-nAChR from unlinked human $\alpha7$ subunits (typically > 1 µA at 7 days after mRNA injection) was easily measurable ($\approx 100 - 300$ nA peak current response, depending on the construct). The time-course of desensitization / inactivation following a peak response stimulated by 10 mM ACh (maximally-stimulating concentration) was also measured for each construct. For each construct, desensitization / inactivation was best fit by a double exponential decay model. As detailed in the legend to Fig 3 no significant differences were seen between the fast desensitization / inactivation time constants calculated for each group. This is not surprising since the apparent time constants will likely reflect the relatively slow kinetics of agonist application in the apparatus, rather than the much faster kinetics of $\alpha7^*$-nAChR desensitization (Papke, 2010). Indeed the apparent $\tau_{fast}$ values are very similar to those measured for solution exchange in our apparatus (Eaton et al, 2014). However, the $\tau_{slow}$ value calculated for the ($\alpha7$)$_3$($\beta2$)$_2$ construct was significantly slower than those associated with the other groups. Thus, despite the admitted disadvantages of measuring kinetic parameters in the *Xenopus* oocyte expression system, there is some evidence that $\alpha7\beta2^*$-nAChR desensitization may be slower than that of homomeric $\alpha7$-nAChR.

**Agonist and antagonist pharmacology of concatemeric human $\alpha7^*$-nAChR**

Pharmacological parameters of selected ligands were determined at concatenated $\alpha7^*$-nAChR. Compounds chosen included the prototypical agonist s, ACh and nicotine,
choline (which is a relatively selective agonist of α7 nAChR (Alkondon et al., 1997)), and two further agonists with established selectivity for other β2*-nAChR subtypes (sazetidine-A and CC4 (Kozikowski et al., 2009; Sala et al., 2013; Xiao et al., 2006)). Agonist pharmacological profiles for (α7)5-, (α7)4(β2)1-, and (α7)3(β2)2-nAChR subtypes were largely indistinguishable from each other, and from that for non-concatemeric (loose-subunit), homomeric α7nAChR (Fig. 4; Table 4). The only exception is that nicotine has significantly lower efficacy (normalized to that of ACh) at both α7β2*-nAChR subtypes than at concatemeric (α7)5-nAChR or unlinked α7- nAChR (which are statistically indistinguishable on this measure). There was also a trend towards lower nicotine potency across all concatemeric α7*-nAChR constructs, but this did not reach statistical significance (see Table 4). The observed slight trend towards lower choline efficacy, although not significant, is suggestive of the previous observation of 50-70% efficacy of choline vs. ACh at putative α7β2 nAChR expressed from non-linked subunits (Khiroug et al., 2002; Zwart et al., 2014). Strikingly, both sazetidine-A and CC4 were very weak agonists (< 10% efficacy normalized to that of ACh) at all α7*-nAChR subtypes tested, including both α7β2-nAChR, making it impossible to reliably calculate EC50 or Hill slope values from the resulting concentration-response data.

Concentration / response relationships were explored for archetypal α7 antagonists (MLA and the snake venom α-toxin, α-Cbtx), together with the β2-selective antagonist DHβE and the non-competitive antagonist MECA (Fig. 5). The resulting pharmacological parameters are summarized in Table 5. Similarly to the agonist pharmacology, antagonist responses were statistically-indistinguishable between the α7* subtypes (including between α7-only nAChR expressed from either unlinked subunits, or from the concatenated α7 homopentameric construct).
DISCUSSION

This study provides the first direct evidence that α7β2-nAChR are expressed in the mammalian CNS. This is demonstrated by isolation of Bgtx-binding or α7 subunit-containing complexes also shown to contain β2 subunits from human or mouse forebrain samples. In addition, we have demonstrated for the first time that multiple human α7β2-nAChR isoforms of defined subunit composition have pharmacological profiles similar to each other and to homopentameric α7-nAChR.

Our findings indicate that α7β2-nAChR are found in post-mortem, human basal forebrain but not in the cerebellum. Note that total amounts of α7*-nAChR are < 2-fold different in the two brain regions. Specificity of the anti-α7 or anti-β2 Abs used in western blot analysis of these nAChR is demonstrated by control studies using cell lines transfected with specific nAChR subunits, and by studies using WT and subunit-null mice. We also found α7β2-nAChR expression in mouse basal forebrain but not hippocampus. Our results agree with earlier findings of α7β2-nAChR expression in mouse basal forebrain (Liu et al., 2009), but not with the same investigators’ study in mouse hippocampus (Liu et al., 2012). There could be several explanations for these seemingly-discrepant observations. nAChR α7 and β2 subunit mRNAs are co-expressed in both basal forebrain and hippocampal cholinergic neurons (Azam et al., 2003). However, fewer than 3% of β2*-nAChR in WT mouse basal-forebrain extracts (this study) were associated with the α7 subunit. This indicates that the large majority of α-Bgtx-binding sites are homomeric α7-nAChR. Accordingly, we feel that the most-likely explanation for the lack of immunochemically-detectable α7β2-nAChR in mouse hippocampus is that it is even less prevalent than in basal forebrain. The previous electrophysiology experiments (Liu et al., 2012) used brain slices from very young mice, whereas our work used tissue from 4-6 month old mice. Therefore, it is also possible that mouse hippocampal α7β2-nAChR expression levels fall from early life into adulthood. Multiple examples of developmental modulation of nAChR subunit expression (including of α7) have previously been seen (Balestra et al., 2000; Conroy and...
The use of a linked-subunit approach allowed us, for the first time, to directly assess the effects of defined β2 nAChR subunit incorporation on α7*-nAChR function. Of critical importance, no significant differences in EC/IC₅₀ values or efficacy relative to ACh were seen between concatenated or unlinked-subunit homomeric α7-nAChR. This indicates that, as has previously been shown for α3β4*-nAChR (George et al., 2012; Stokes and Papke, 2012), α4β2-nAChR (Carbone et al., 2009; Eaton et al., 2014; Mazzaferro et al., 2011; Zhou et al., 2003), and α6β2*-nAChR (Kuryatov and Lindstrom, 2011) subtypes, introduction of appropriately-sized linkers can be performed without altering nAChR functional pharmacology. Several of these previous studies also showed that concatemeric constructs were assembled correctly. To further confirm correct that concatemers were being assembled correctly and not fragmenting and rearranging into unanticipated functional forms, we also coinjected unlinked β2 subunits containing a gain-of-function mutation (L9'S) in the second transmembrane domain. This additional control has previously been used by us and others (Carbone et al., 2009; Eaton et al., 2014). If concatemer fragments were contributing to the functional nAChR population, the β2-gain-of-function subunit would assemble into resulting α7*-nAChR as previously shown (Khiroug et al., 2002; Murray et al., 2012; Zwart et al., 2014). Therefore, if fragments containing α7 were present, this would result in appearance of a novel α7β2-gain-of-function population with distinctive (more agonist-sensitive) properties. However, no such effect was seen.

It is noted, however, that overall function was reduced when α7-nAChR homopentamers were expressed from a concatemeric construct as opposed to from unlinked subunits. This relative diminution in function of concatenated nAChR constructs has been noted in the previous publications just cited and appears to be a regular feature of using concatemeric nAChR constructs. Importantly, however, both (α7)₄(β2)₁-and (α7)₃(β2)₂-nAChR concatemer constructs expressed more function than did the (α7)₅-nAChR concatemer. This is the opposite of the situation where loose β2 nAChR subunits are co-
expressed with α7 subunits (Murray et al., 2012), and replicates an earlier finding in which co-expression of unlinked α5, α3, and β4 nAChR subunits reduced function compared to expression of loose α3 and β4 subunits alone, but incorporation of the α5 subunit into a concatemeric construct actually increased observed function of an α3β4*-nAChR pentameric concatemer (George et al., 2012). As in the previous publication, we suspect that uncontrolled assembly of an unlinked additional subunit (in this case β2) may be deleterious, but directed assembly may result in greater functional expression of the new nAChR subtype. Certainly, the current study provides direct evidence that β2 subunit incorporation into α7*-nAChR is compatible with agonist-induced function.

The pharmacological profiles of α7β2-nAChR were very similar to those of homopentameric α7-nAChR. Even agonists (sazetidine-A, CC4) and an antagonist (DHβE) previously shown to have significant β2*-nAChR selectivity had indistinguishable pharmacology across homomeric α7-nAChR and the two different α7β2-nAChR isoforms. Each of these findings match those very recently published using Xenopus oocytes expressing α7 and β2 subunits at a 1:10 ratio (Zwart et al., 2014). The only statistically-significant difference in the present study was a diminution of nicotine’s efficacy relative to that of ACh in the two α7β2-nAChR isoforms (also seen by (Zwart et al., 2014)). This nicotine partial agonism further confirms that β2 was incorporated into α7β2-nAChR concatemers as planned and may represent a pharmacological marker for the presence of α7β2-nAChR. The same may be true of the slower desensitization kinetics measured for the (α7)₃(β2)₂ (Figure 3), although it is important to note the limitations of measuring receptor kinetics in a Xenopus oocyte system (Papke, 2010). We note that the similar α7-nAChR vs. α7β2-nAChR potency of DHβE observed by us and (Zwart et al., 2014) does not match the observations made in two previous studies (Liu et al., 2009; Murray et al., 2012). The reason for this discrepancy between the pairs of studies is not clear, but two possible explanations occur. First, the differences previously measured are relatively subtle, so may be hard to reproduce. Related to this point, we note that the Hill-slopes of the α7β2-nAChR DH βE
CRCs (Figure 5A) are shallower than those measured for other competitive antagonists (≤ 1, as opposed to significantly > 1 for MLA and α-Cbtx). This would tend to obscure fine differences in IC₅₀ values. Second, other α7 and β₂ subunit associations are possible, in addition to those used in the α7β₂ nAChR concatemers deployed in this study. It is possible that an α7β₂*-nAChR population expressed from unlinked subunits may assemble differently, giving rise to the slightly-different DHβE sensitivity previously measured. This would match the previous experience in which α3β₄α5-nAChR pharmacology perfectly matched between concatenated and unlinked-subunit α3β₄-only nAChR, but that of loose-subunit constructs (George et al., 2012). Further work may be needed to understand the (admittedly subtle) pharmacological differences between alternative α7β₂-nAChR subunit stoichiometries and association orders.

Overall, however, the functional pharmacology of α7-nAChR and α7β₂-nAChR subtypes is remarkably similar. This observation indirectly supports the concept that activation of α7β₂-nAChR may be predominantly or exclusively mediated only through agonist binding sites at α7/α7 (not α7/β₂) interfaces (Murray et al., 2012). If this is true, it seems unlikely that any competitive agonist could exhibit a significantly-different potency between α7-nAChR and α7β₂-nAChR. However, antagonists capable of disrupting the allosteric transitions required for nAChR activation (Celie et al., 2005), and of selectively binding to α7/β₂ interfaces, could be valuable in this regard as could other non-competitive ligands. In the concatemeric (α7)₄(β2)₁-nAChR construct (subunit order α7-α7-β₂-α7-α7), only three α7/α7 subunit interfaces will be retained (between the first two subunits, the last two subunits, and between the first and last subunits which will assemble together to complete the pentameric nAChR structure). In the (α7)₃(β2)₂-nAChR construct, only the α7/α7 interface formed between the first and last subunits will be retained. At first glance, it may seem remarkable that an α7*-nAChR containing such a diminished complement of putative agonist binding sites could be effectively activated. However,
elegant recent work indicates that nAChR, including α7-nAChR, can be activated effectively by as few as one agonist binding site (Andersen et al., 2013; Rayes et al., 2009; Williams et al., 2011).

That α7β2-nAChR are relatively scarce in basal forebrain does not imply that their role is necessarily insignificant. For example, α6β2*-nAChR expression on SN/VTA dopamine projections comprises < 10% of all β2*-nAChR in dopaminergic regions (Gotti et al., 2005; Whiteaker et al., 2000), but this subtype is extremely important in controlling local neuronal behaviour and signal processing (Exley et al., 2008; Exley and Cragg, 2008). Cholinergic neurons constitute only a fraction (10-15%) of basal forebrain neurons (Semba 2000) and the proportion of α7β2-nAChR in these neurons may therefore be relatively large. The basal forebrain cholinergic system provides many cholinergic innervations to limbic and cortical brain structures, and expresses nAChR that participate in the cholinergic transmission and cognitive processes associated with learning and memory (Hernandez et al., 2010; Voytko et al., 1994). One of the most marked pathological changes in AD brain is the degeneration of this cholinergic projection and the consequent reduction in the number of nAChR (Dumas and Newhouse, 2011; Pinto et al., 2011). A number of studies have found that the beta-amyloid (Aβ) peptide (a hallmark of AD) plays a critical role in neuronal degeneration and subsequent memory deficits (Capsoni et al., 2000; Dolga et al., 2009; Fraser et al., 1997; Holtzman et al., 1992; Price et al., 1985; Wenk, 1993). Further, a recent electrophysiological study has demonstrated that Aβ binds with higher affinity to α7β2-nAChR than to α7-nAChR, and that this can produce hippocampal neuronal hyperexcitation (through α7-nAChR upregulation) and subsequent neurodegeneration (Liu et al., 2013).

Post-mortem tissue is an under-used substrate for genetic and/or preclinical studies, and provides a translational element that is difficult to recapitulate in animal models alone (McCullumsmith et al., 2014). This study’s definitive evidence that α7β2*-nAChR are found in human, as well as mouse, basal forebrain provides valuable support for the concept that
this subtype may be relevant to the study and etiology of Alzheimer's disease. The similarities in human- and mouse-brain basal forebrain α7β2*-nAChR expression are also supportive of the use of mouse models in this context.
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AUTHORSHIP CONTRIBUTIONS:

Participants in research design: Gotti C., Zoli M., George A.A., Lukas R.J., Whiteaker P.

Conducted experiments: Moretti M., Pistillo F., and George AA.

Contributed new reagents or analytic tools: Maskos U., Pistillo F., and Whiteaker P.

Performed data analysis: Whiteaker P., Moretti M., Pistillo F., George AA., and Whiteaker P.

Wrote or contributed to writing of the manuscript: Gotti C., Zoli M., Lukas R.J., Whiteaker P.
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FOOTNOTES:

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†M.M., M.Z., and A.A.G. contributed equally to this work.

§P.W. and C.G. contributed equally to this work and are joint corresponding authors.
FIGURE LEGENDS:

Figure 1. Western blot analysis of nAChR subunit content in α-Bgtx-purified receptors prepared from 2% Triton X-100 extracts of WT and β2 KO mouse hippocampi (A) and basal forebrain samples (B).

A) α-Bgtx-purified receptors were prepared from mouse hippocampi by incubating extracts with Sepharose 4B covalently bound with α-Bgtx. The bound receptors were recovered from the beads using Laemmli sample buffer. Western blot analysis of 10 µg of 2% Triton X-100 extracts of the hippocampus before (lane 1) and after α-Bgtx purification (lane 2; supernatant), and 1/20 of the corresponding α-Bgtx purified receptors (lane 3; recovered from beads). The blots were probed with an anti-α7 Ab (top) or β2(1) Ab (bottom).

B) α-Bgtx purified receptors were prepared as described in the legend of Figure 1. Western blot analysis of 10 µg of 2% Triton X-100 extracts of the basal forebrain before (lane 1) and after α-Bgtx purification (lane 2; supernatant), and 1/10 of the corresponding α-Bgtx purified receptors (lane 3; recovered from beads). The blots were probed with an anti-α7 Ab (top) or β2(1) Ab (bottom).

Figure 2. Western blot analysis of α-Bungarotoxin-purified nAChR prepared from human basal forebrain and cerebellum.

α-Bgtx-binding nAChR were purified from the same volume of 2% Triton X-100 extracts of basal forebrain and cerebellum by incubating them with Sepharose 4B covalently bound with α-Bgtx. The bound receptors were eluted using sample buffer, and an identical volume of purified receptors was loaded on the gel. The Western blots were probed with anti-α7 Ab (top) or anti-β2 Ab (bottom).

Figure 3. Representative traces and maximum function (I_{max}) comparison for α7*-nAChR pentameric concatemer constructs.
Oocytes were injected with mRNA encoding unlinked α7-nAChR subunit monomers (Panel A) concatenated α7 homopentamers (Panel B), α7β2 nAChR with the β2 subunit in position 3 (Panel C), or α7β2 nAChR with the β2 subunit in positions 2 and 4 (Panel D). Representative two-electrode voltage-clamp recordings are shown in each case, for ACh concentration-response determinations (see Methods for details). Black bars above each trace represent 5 s applications of ACh at a range of concentrations. The time course of receptor desensitization / inactivation during stimulation with a maximally-effective dose of ACh (10 mM) was also investigated for each nAChR construct, using additional groups of oocytes. In each case, the time course was best fit by a double-exponential decay. The fast time constants (τfast) for desensitization / inactivation were statistically indistinguishable by one-way ANOVA across all four groups (unlinked α7, 436 ± 85 ms; α7-only concatemer, 214 ± 80 ms; α7β2(p3), 312 ± 55 ms; α7β2(p2,4), 247 ± 35 ms; F[3,11] = 2.06, p = 0.16; n = 3 in each group). In contrast, the slow time constant (τslow) for desensitization / inactivation of the α7β2(p2,4) construct was significantly longer than of the other groups; no other differences were detected by Tukey's post hoc comparison (p < 0.05). Values were: unlinked α7, 5109 ± 800 ms; α7-only concatemer, 3130 ± 585 ms; α7β2(p3), 5073 ± 638 ms; α7β2(p2,4), 6318 ± 365 ms; F[3,11] = 5.29, p = 0.02; n = 3 in each group.

Panel E, summary of maximal function (I\text{max}) measured in each concatameric nAChR group by stimulation with the full agonist ACh (10 mM). Bars represent mean ± SEM (n = 3). I\text{max} values were: α7-only, 83.9 ± 18.6 nA; α7β2(p3), 285 ± 11 nA; α7β2(p2,4), 216 ± 45 nA. Analysis using one-way ANOVA with Tukey's post hoc comparison showed that incorporation of β2 subunits resulted in a statistically-significant increase in I\text{max} (F[2,6] = 12.7, p = 0.007; denoted by *). The I\text{max} values obtained from the two α7β2-nAChR constructs were statistically indistinguishable from each other.

**Figure 4. Agonist concentration response profiles for α7 and α7β2 nAChR.**

Oocytes were injected with mRNA encoding unlinked α7 subunits (o), concatenated α7 homopentamers (●) or concatenated α7β2 pentameric concatemers (□ indicates α7β2.
nAChR with the β2 subunit in position 3; ■ indicates α7β2 nAChR with the β2 subunit in positions 2 and 4). Oocytes were perfused with nAChR agonists (A) acetylcholine (ACh; 10^{-5.5} to 10^{-2}; n=6), (B) choline (10^{-4.25} to 10^{-2}; n=3), (C) nicotine (10^{-5.5} to 10^{-3}; n=3), (D) sazetidine-A (10^{-7.5} to 10^{-4}; n=3) or (E) 1,2-bis-N-cytisinylethane (CC4; 10^{-6.5} to 10^{-3}; n=3). All responses within each group were normalized to an initial control stimulation with 10 mM ACh. Data points represent mean ± SEM. Drug potency and efficacy parameters were calculated by non-linear least-squares curve fitting to the Hill equation (see Methods). The resulting pharmacological parameters and statistical analyses are summarized in Table 4.

Figure 5. Antagonist concentration response profiles for α7 and α7β2 nAChR.

Oocytes were injected with mRNA encoding unlinked α7 subunits (○), concatenated α7 homopentamers (●) or concatenated α7β2 pentameric concatemers (□) indicates α7β2 nAChR with the β2 subunit in position 3; ■ indicates α7β2 nAChR with the β2 subunit in positions 2 and 4). Before antagonists were applied to each oocyte, a control 10 mM ACh-evoked response was measured. Oocytes were pre-perfused with nAChR antagonists (A) dihydro-β-erythroidine hydrobromide (DHβE; 10^{-6.25} to 10^{-3}; n=3), (B) methyllycaconitine (10^{-10.5} to 10^{-7}; n=3), (C) mecamylamine (10^{-7.25} to 10^{-4}; n=3) or (D) α-cobratoxin (α-Cbtx; 10^{-10} to 10^{-7}; n=3). The magnitudes of subsequent 10 mM ACh stimulations were compared to that of the initial control. Data points represent mean ± SEM. Drug potency and efficacy parameters were calculated by non-linear least-squares curve fitting to the Hill equation (see Methods). The resulting pharmacological parameters and statistical analyses are summarized in Table 5.
<table>
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<tr>
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<th>Number of cases</th>
<th>Age in Years</th>
<th>Postmortem delay in hours</th>
<th>Male/female</th>
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<td>Basal forebrain</td>
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<td>9.4</td>
<td>2 &gt;8</td>
<td>3/1</td>
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<tr>
<td></td>
<td></td>
<td></td>
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<tr>
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<td>4 73.0±</td>
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<td>&gt;8</td>
<td>2/2</td>
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<tr>
<td>Cerebellum non-smokers</td>
<td>4 68.7±</td>
<td>6.6</td>
<td>&gt;8</td>
<td>2/2</td>
</tr>
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**TABLE 1: Details of cases sampled for receptor analysis.** Values are means ± SEM. There were no significant differences between groups for age.
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<th>[³H]-Epibatidine</th>
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<td>β2 KO Basal forebrain</td>
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<td>1 ±2.6</td>
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**TABLE 2:** Levels of [³H]-Epibatidine and [¹²⁵I]-α-Bungarotoxin binding to 2% Triton X-100 extracts (expressed as fmol/mg of protein) in two different brain areas of WT and β2 KO mice. Values are the Mean ± SEM from three separate experiments. * = Significantly different from β2+/+ by t-test (p < 0.001).
<table>
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<th>[^{125}\text{I}]-(\alpha)-Bungarotoxin</th>
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<td>Cerebellum smokers</td>
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</tbody>
</table>

**TABLE 3:** Levels of \[^3\text{H}\]-Epibatidine and \[^{125}\text{I}\]-\(\alpha\)-Bungarotoxin binding to 2% Triton X 100 extracts (expressed as fmol/mg of protein) in the two different human brain regions. Values are mean ± SEM of the 4 samples in each group. * Denotes significant differences in cerebellar membrane \[^3\text{H}\]-epibatidine binding between smokers and non-smokers: unpaired t test: \(p = 0.02\). No significant difference was seen between \[^{125}\text{I}\]-\(\alpha\)-Bgtx binding levels in cerebellar samples taken from smokers vs. non-smokers, by the same measure.
TABLE 4: α7*-nAChR agonist pharmacological parameters. Agonist logEC50, Hill slope (nH) and efficacy values (relative to a maximally-effective (10 mM) concentration of ACh) were derived by non-linear least-squares curve fitting of the data shown in Figure 4 to the Hill model. α7-only nAChR expressed in Xenopus oocytes from unlinked subunits were used as a control group, to which the functional properties of α7-nAChR concatemeric constructs were compared (N-to-C-terminal subunit order are shown). Values are mean ± SEM of the number of indicated replicates (n=). nd = not determinable (reliable curve fitting is not possible for very low-efficacy compounds). Pharmacological parameters measured for each agonist were generally indistinguishable between all four groups of oocytes, with one exception: the relative efficacy of nicotine was lower for both α7β2 subtypes tested compared to the α7 unlinked control group (although the α7-only concatemer group was not different to the control); One way ANOVA F[3,8] = 34.2, p < 0.001, followed by Dunnett’s post-hoc test.
<table>
<thead>
<tr>
<th>Subtype</th>
<th>Dihydro-β-Erythroidine</th>
<th>Methyllycaconitine M</th>
<th>ecamylamine</th>
<th>α-Cobratoxin</th>
</tr>
</thead>
<tbody>
<tr>
<td>α7 (unlinked)</td>
<td>n= log(IC50 / M)</td>
<td>nH n= log(IC50 / M)</td>
<td>nH</td>
<td>n= log(IC50 / M)</td>
</tr>
<tr>
<td></td>
<td>3 -5.2 ± 0.05 -1.0 ± 0.1</td>
<td>3 -8.7 ± 0.1 -1.6 ± 0.2</td>
<td>3 -5.6 ± 0.2</td>
<td>3 -8.6 ± 0.1 2.4 ± 0.5</td>
</tr>
<tr>
<td>α7-α7-α7-α7-α7</td>
<td>3 -5.3 ± 0.07 -0.8 ± 0.1</td>
<td>3 -9.0 ± 0.1 -1.6 ± 0.2</td>
<td>3 -6.0 ± 0.2</td>
<td>3 -8.6 ± 0.1 1.3 ± 0.8</td>
</tr>
<tr>
<td>α7-α7-β2-α7-α7</td>
<td>3 -5.4 ± 0.10 -0.6 ± 0.1</td>
<td>3 -8.9 ± 0.1 -1.7 ± 0.1</td>
<td>3 -6.0 ± 0.1</td>
<td>3 -8.6 ± 0.1 1.8 ± 0.5</td>
</tr>
<tr>
<td>α7-β2-α7-β2-α7</td>
<td>3 -5.4 ± 0.10 -0.7 ± 0.1</td>
<td>3 -8.8 ± 0.1 -1.8 ± 0.1</td>
<td>3 -6.0 ± 0.2</td>
<td>3 -8.5 ± 0.2 1.5 ± 0.6</td>
</tr>
</tbody>
</table>

**TABLE 5: α7*-nAChR antagonist pharmacological parameters.** Antagonist log IC50 and Hill slope (nH) values were derived by non-linear least-squares curve fitting of the data shown in Figure 5 to the Hill model. Values are mean ± SEM of the number of indicated replicates. Pharmacological parameters obtained for each antagonist were statistically indistinguishable between all four groups of oocytes according to analysis with one way ANOVA.
Figure 1

(A) Hippocampus

(B) Basal Forebrain

α7

β2 WT | β2 KO
---|---
1 | 2 | 3

57 kDa

β2

β2 WT | β2 KO
---|---
1 | 2 | 3

53 kDa