Visual Acuity Is Reduced in Alpha 7 Nicotinic Receptor Knockout Mice

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PURPOSE. Nicotinic receptors (nAChRs) are part of a heterogeneous family of pentameric ligand-gated ion channels that are widely expressed in the visual system. The impact of α7 homomeric nAChRs on visual function was investigated using mutant mice lacking the α7 nicotinic receptor subunit.

METHODS. The spatial resolution limit was measured in α7 knockout (α7 KO) and age-matched control mice using three independent methods: an operant behavioral visual task (Prusky maze), cortical visual evoked potentials (VEPs), and the pattern electroretinogram (PERG) evoked by alternating gratings of different spatial frequencies and contrasts.

RESULTS. Visual acuity measured by means of the visual water maze task was significantly decreased in the α7 KO mice and, concurrently, there was a reduction of the cortical spatial resolution limit measured by VEPs. However, the PERG was normal in α7 KO mice, compared with control mice. The use of fluorescently tagged cholera toxin showed that projections from the retina segregate normally in α7 KO mice and, in line with this, the visual cortical responses elicited by stimulating either eye were normally balanced in both visual cortices and showed no retinotopic anomalies.

CONCLUSIONS. These findings indicate that the absence of the α7 nicotinic subunit reduces visual acuity. Because the cortical output has an abnormal spatial resolution but retinal output is preserved, it can be concluded that the low visual acuity was due to a deficit specifically present in the visual cortex.

Visual Neurophysiology

Nicotinic receptors (nAChRs) represent a heterogeneous family of ion channels that are differently expressed in the nervous system. There are 12 subunit genes that derive from a common ancestral gene: 9 α subunits (α2 to α10) and 4 β subunits (β2 to β4). Neuronal nAChRs fall into two main classes: homomeric or heteromeric α-bungarotoxin (α-Bgt)-sensitive receptors consisting of α7, α8, or α7-α8 and/or α10 subunits and α-Bgt-insensitive heteromeric receptors consisting of α2-α6 and β2-β4 subunits. Nicotinic AChRs are permeable to Na+, K+, and Ca2+ ions, and their cation permeability is influenced by their subunit composition. The α7 subunit that forms native pentameric homomeric receptors is highly expressed in the hippocampus, hypothalamus, and neocortex of rodents.1 Alpha-7 knockout (KO) mice, which are characterized by the loss of α-Bgt receptors and the lack of nicotine-evoked fast desensitizing currents in neurons,2 are viable with apparently normal brain anatomy.

In terms of behavior, the performance of α7 KO mice in the classic Morris water maze test, the Pavlovian conditioned fear test and the prepulse inhibition paradigm are similar to those of wild-type (WT) mice, which suggests that the absence of α7 nAChRs has little impact on normal, baseline behavioral responses.3 Moreover, α7 KO mice show a reduced anxiety-related response,3 whereas only mice lacking both the α7 and the β2 nAChR subunits show impaired learning and memory in a passive avoidance test, and enhanced motor activity on the rotarod.3 More recently, a deficit in a modified Morris water maze task (delayed matching to place preference) has been described in mice lacking the α7 subunit,5 thus suggesting a minor impairment in episodic/working memory such as that observed in previous hippocampus studies based on the infusion of an α7 subunit antagonist.6

Although α7 KO mice are widely used in behavioral studies, no attempt has been made to characterize their visual function. The α7 subunit is widely expressed in the visual system, particularly in the retina and retinal input recipient regions such as the dorsolateral geniculate nucleus (dLGN), the superior colliculus, and the visual cortex.1,7,8 Recent studies have shown that visual system development and function require the activity of nAChRs: in particular, the β2 subunit is required for the formation of eye-specific layers at the thalamic level, a process that depends on retinal waves of spontaneous activity.9,10 However, the impact of a complete lack of the α7 nicotinic subunit on visual system development and functional phenotype acquisition is still unknown.

METHODS

Animals

All the experiments were carried out in accordance with the European Community Council Directive concerning the treatment of animals using mice at least 2 months old. We used α7 KO mice homozygous for the ChRNA7−/− mutation (Jackson Laboratory, Bar Harbor, ME) and, in parallel, mice of the same mixed genetic background (C57BL/6J) as controls. To exclude the possibility that the mixed genetic background of the mutant mice may affect the results of the study, the α7 KO mice were backcrossed for 10 generations onto the C57BL/6 strain (Jackson Laboratory).

Binding Assay and Immunoprecipitation

The sequences of polyclonal antibodies against the α2, α3, α4, α5, α6, β2, β3, and β4 peptides were raised and characterized as previously.
the tissue extracts was performed by incubating the extracts with 2 nM 
from binding to the subtypes containing this subunit. The binding to 
scried.11,12

The eyes and visual cortex were dissected, immediately frozen in liq-
uid nitrogen, and then stored at −80°C for later use. In each experi-
ment, the eyes or visual cortex were separately homogenized in an 
excess of 50 mM Na phosphate (pH 7.4), 1 M NaCl, 2 mM EDTA, 2 
mM EGTA, and 2 mM phenylmethylsulfon fluoride (PMSF) for 2 min-
utes in a commercial homogenizer (Ultra Turrax; IKA Works, Wilming-
ton, NC), after which the homogenates were diluted and centrifuged 
for 1.5 hours at 60,000 g. The whole eyes and visual cortex were 
homogenized, diluted, and centrifuged twice, after which the 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, and 
then resuspended in the same buffer containing a mixture of 10 μg/mL 
of each of the following protease inhibitors: leupeptin, bestatin, pep-
statin A, and aprotinin. Triton X-100 at a final concentration of 2% 
was added to the washed membranes, which were extracted for 2 hours 
at 4°C. The tissue extracts were then centrifuged for 1.5 hours at 
60,000 g, recovered, and an aliquot of the resultant supernatants was 
collected for protein measurement using the BCA protein assay (Pierce 
Protein Research Products, Thermo Fisher Scientific, Rockford, IL) 
with bovine serum albumin as the standard.

The specific activity of the [125I]-α-Bgtx (PerkinElmer, Boston, MA) 
was 150 Ci/mmol. Binding to the visual cortex membranes was per-
formed by means of overnight incubation with a saturating concentra-
tion of 5 nM [125I]-α-Bgtx at room temperature. Nonspecific bind-
ing was determined in parallel by means of incubation in the presence 
of 1 μM unlabeled α-Bgtx. After incubation, the samples were filtered 
and the bound radioactivity was directly counted in a γ counter. To ensure 
that the α7 subtype did not contribute to 3H-epibatidine (3H-Epi) 
binding, the binding to the Triton X-100 extracts and the immunopre-
cipitation experiments were performed in the presence of 1 μM 
α-Bgtx, which specifically binds to the α7 subtype and prevents Epi 
from binding to the subtypes containing this subunit. The binding to 
the tissue extracts was performed by incubating the extracts with 2 nM 
3H-Epi in the presence and absence of 100 nM cold Epi using an 
ion-exchange resin (DE52; Whatman, Maidstone, UK) as previously 
described.11,12

For the immunoprecipitation experiments, the extracts obtained 
from tissues preincubated with 1 μM α-Bgtx were labeled with 2 nM 
3H-Epi and then incubated overnight with a saturating concentration 
of affinity-purified anti-subunit-specific IgG (20 μg). The immunopreci-
pitation was recovered by incubating samples with beads containing 
bound anti-rabbit goat IgG (Technogenetics, Milan, Italy). The level of 
antibody immunoprecipitation was expressed as fmol of immunopre-
cipitated receptors/mg of protein.

Behavioral Assessment of Visual Acuity Using the Visual Water Task
A behavioral measure of visual acuity was obtained using the method 
described by Prusky et al.15 Starting at postnatal day 60 (P60), the α7 
KO (n = 5) and WT mice (n = 5) were trained and tested in the visual 
water task for 10 trials every day to assess their visual acuity. The visual 
water task first trains the animals to choose a low (0.05 cycle/deg) 
spatial frequency vertical grating from gray and then tests the limit of 
their discrimination ability at increasing spatial frequencies. The high-
est spatial frequency at which 70% discrimination accuracy is achieved 
is taken as the visual acuity. (See Fig. 2A later in text, which shows a 
schematic representation of the visual water task.)

VEP and ERG Recordings
Visual evoked potentials were recorded as described by Porciatti et 
al.16 Briefly, the mice were anesthetized by means of an intraperitoneal 
injection of 20% urethane and mounted in a stereotaxic apparatus 
allowing a full view of the visual stimulus. After carefully removing a 
portion of the skull overlying the binocular visual cortex while leaving 
the dura intact, a glass-pulled recording electrode filled with NaCl (3 
M) was inserted into the cortex perpendicularly to the stereotaxic 
plane. The electrical signals were amplified (10,000-fold), band-pass 
filtered (0.3–100 Hz), digitized, and averaged (at least 75 events in 
blocks of five events each). The transient VEPs in response to an abrupt 
contrast reversal (1 Hz) were evaluated in the time domain by measur-
ing the peak-to-trough amplitude and peak latency of the major com-
ponent. The visual stimuli consisted of horizontal gratings of different 
spatial frequencies and contrasts generated by a visual stimulator in-
terface (VSG2.2 card; Cambridge Research System, Cheshire, UK) and 
presented on the screen of a monitor (Sony model CPD-G520) placed 
20 cm in front of the animal. To obtain an estimate of contralateral 
ocular bias using VEPs, the electrode was positioned 2.8–3 mm later-
ally to lambda in accordance with the retinotopy and ocularity analysis 
described by Porciatti et al.14; in agreement with these previous data, the 
response to a windowed stimulus with different azimuths gave a 
maximal amplitude at nearly 10° lateral to the vertical midline.

The flash ERG (F-ERG) was evoked by means of 10-ms flashes of 
light generated by a commercial optical stimulator (Ganzfeld stimula-
tor; Lace, Pisa, Italy). The electrophysiological signals were recorded 
through gold-plate electrodes inserted under the eyelids and in con-
 tact with the cornea, which had been previously anesthetized with 
ossibuprocarbonate (Novesine; Novartis Pharma, Basel, Switzerland). The 
corneal electrode at each eye was referred to a needle electrode (one 
on each side) subcutaneously inserted into the ipsilateral region be-
hind the ear. The different electrodes were connected to a two-channel 
amplifier.

For the recordings made under scotopic conditions, the animals 
were dark adapted (300 minutes), anesthetized with an intraperitoneal 
injection of 2.2,2-tribromoethanol (Avetrin 2 mL/100 g body wt; 1.25% 
[w/v]) 2,2,2-tribromoethanol; and 2.5% [w/v] 2-methyl-2-butanol; Sig-
ma-Aldrich, St. Louis, MO), and loosely mounted in a stereotaxic 
apparatus under dim red light. Their body temperature was kept at 
37.5°C and their heart rate was monitored. Immediately before 
the recording session, the pupil was dilated by topically applying one eye 
drop containing atropine (0.5% sol; Allergan, Pomezia, Italy). The 
scotopic ERGs were recorded in response to single flashes of different 
light intensities ranging from 10−4 to 20 cd/m² × s−1 in the 
presence of constant background illumination of 15 cd/m². The b-wave 
amplitude of each eye was averaged and plotted as a function of 
increasing luminance (transfer curve); for each animal group, the mean 
b-wave amplitude was plotted as a function of different luminances and 
rearing conditions.

The pattern electroretinograms (PERGs) were recorded as de-
scribed in Porciatti et al.15,16 using a recording electrode made of a thin 
silver wire (diameter, 0.25 mm) configured as a semicircular loop, with 
a radius of 2 mm, that was gently placed on the corneal surface by 
means of a micromanipulator under microscopic control. To maximize 
the PERG amplitude, patterned visual stimuli were used as described 
earlier at a mean luminance of 30 cd/m², with different spatial frequen-
cies and contrasts.

Fluorescent Tracing of Retinogeniculate Projections
Adult mice were anesthetized with 2,2,2-tribromoethanol (Avetrin 2 
ml/100 g body wt) and 2 μL of 1% Alexa Fluor 488–conjugated 
(invitrogen) cholera toxin B (CTB) subunit was injected into one eye 
using pulled glass capillaries inserted just behind the corneoscleral 
marg in. After 48 hours, the mice were anesthetized and perfused with 
4% paraformaldehyde in PBS. Their cryoprotected brain was cut into 
40-μm coronal sections using a freezing microtome. The images of 
the sections were acquired using an optical microscope (Nikon) and ana-
lyzed using ImageJ software (developed by Wayne Rasband, National 
Institutes of Health, Bethesda, MD; available at http://rsb.info.nih.gov/
RESULTS

The α7 KO mice were anatomically and morphologically indistinguishable from the mice of the same mixed genetic background (C57BL/6J mice) used as controls.

Although the findings of a considerable number of studies of α7 KO mice have been published, little is known about the developmental changes in nAChR subunits occurring in the brains of α7-deficient mice. We investigated the levels of different nAChR subunits in the retina and cortex of α7 KO and WT mice, and analyzed the presence of α7-containing receptors by means of binding with the specific α7 ligand, [3H]Bgtx. As expected, specific binding of [3H]Bgtx was observed only in the visual cortex of the WT but not in α7 KO mice (17.8 ± 1.3 vs. 0.3 ± 0.1 fmol of specifically bound [3H]Bgtx/mg of protein). The expression pattern of heteromeric receptors in the retina and visual cortex of the WT and α7 KO mice was very similar (Fig. 1), with no statistical differences in the levels of receptors containing the α2, α3, α4, α5, α6, β2, β3, and β4 subunits. Our findings in mouse retina are in line with those previously determined in rat[17] and exclude the possibility that the expression of non-α7 nicotinic subunits is altered in α7 KO mice.

Because previous studies of α7 KO mice have found minor alterations in the learning of spatial tasks, we used the visual water task developed by Prusky et al.[13] to generate a behavior-based measure of visual acuity (Fig. 2). The test involves learning an associating task in a visual water maze (see schematic drawing in Fig. 2A). The behavioral visual acuity of the WT controls was in line with previous findings,[13] and there was no difference in learning ability between the control and α7 KO mice: 90% of correct choices was achieved in 10 sessions by both groups. However, visual acuity was significantly reduced in the α7 KO mice (0.38 ± 0.01 cycle/deg, n = 5, vs. 0.51 ± 0.02 cycle/deg, n = 5; P < 0.05).

VEP recordings were used to relate behaviorally assessed visual acuity to cortical electrophysiological responses to patterned visual stimuli. The VEPs were recorded in at least three spaced positions in the binocular portion of the primary visual cortex.[14] At low spatial frequencies (0.06 - 0.1 cycle/deg, contrast = 90%), the VEPs consisted of a major negative component with a mean latency of 108 ± 5 ms in WT and 104 ± 4 ms in α7 KO mice (P > 0.05; typical examples are shown in Fig. 3A). Spatial frequencies of <0.06 cycle/deg were not used because it was technically impossible to include a sufficient number of cycles (at least three) on our computer screen. In the control mice, VEP amplitudes decreased at spatial frequencies of >0.06 cycle/deg until the signal became indistinguishable from the noise level (calculated for 0% contrast; see Fig. 3A).

Statistics

Student’s t-test was used to compare the data obtained from the α7 KO and WT mice. A value of P < 0.05 was considered statistically significant.

FIGURE 1. Immunoprecipitation analysis of the subunit content of the [3H]-Epi receptors expressed in retina and visual cortex. Triton X-100 (2%) extracts were obtained from membranes prepared from eyes and visual cortex preincubated with 2 μM αBgtx, and then labeled with 2 nM [3H]-Epi. Immunoprecipitation was carried out using saturating concentrations (20 μg) of antisubunit antibodies. In each experiment, the amount immunoprecipitated by each antibody was subtracted from the value obtained in control samples containing an identical concentration of normal rabbit IgG. The results for retina (A) and visual cortex (B) are expressed as fmol of labeled [3H]-Epi receptor/mg of protein, and represent the mean values ± SEM of two (eyes) and three experiments performed in duplicate.

FIGURE 2. (A) Schematic representation of the visual water task. The mice are released in the long arm of the water maze and trained to reach a submerged ‘hidden platform’ associated with the compartment displaying the low-frequency grating, which is positioned in either arm of the maze in a pseudorandom sequence. Once a mouse reaches a 90% success rate in choosing the arm of the maze showing the grating, the grating’s spatial frequency is progressively increased until its discrimination rate drops to 70%. This value is used as a measure of actual visual acuity. (B) Examples of acuity assessment. (C) Mean visual acuity was significantly decreased in the adult α7 KO mice (0.38 ± 0.01 vs. 0.51 ± 0.02 cycle/deg; n = 5; *P < 0.05).
spatial resolution limit in each mouse was calculated by means of linear extrapolation of data points to the noise level, as shown in Figure 3B (VEP amplitudes were plotted as a function of spatial frequencies, semilog coordinates). The mean cortical spatial resolution limit in the control mice was 0.62 ± 0.01 cycle/deg (n = 9; Fig. 3C), which was similar to the value previously reported in the same mouse strain.14 The spatial resolution limit was significantly reduced in the α7 KO mice (0.42 ± 0.02 cycle/deg, n = 5; P < 0.05 vs. controls; Fig. 3C), in line with the data obtained using Prusky’s behavioral method13 (see Figs. 2B, 2C).

To further explore cortical visual function, the contrast threshold was measured by recording the VEP responses to gratings at different contrasts (Fig. 4). VEP amplitude decreased with decreasing contrasts (from 90% to 0%). The amplitudes of the VEPs elicited by gratings of two different spatial frequencies (0.06 and 0.2 cycle/deg) were recorded as a function of the log contrast and the contrast threshold was calculated by means of the linear extrapolation of data to the noise level (Figs. 4A, 4B). The contrast thresholds at both spatial frequencies were not significantly different between the control mice (0.06 cycle/deg: contrast threshold = 5.2 ± 1%; 0.2 cycle/deg: contrast threshold = 15 ± 2%; n = 9; Fig. 4C) and the mutant mice (0.06 cycle/deg: contrast threshold = 4.4 ± 0.5%; 0.2 cycle/deg: contrast threshold = 17 ± 1%; n = 5; Fig. 4C). The lower cortical spatial resolution limit in α7 KO mice was therefore not dependent on the change in contrast gain.

To investigate the functional characteristics of the retina, we first used optical stimulator (Ganzfeld)-generated flashes in dark- and light-adaptation to record electroretinogram (F-ERG) responses in animals aged 2 to 3 months that had been reared under normal lighting conditions (12-hour light/12-hour dark cycle, maximum light intensity < 100 lux). The results clearly showed that there was no significant difference in the scotopic (at all light intensities) or light-adapted F-ERG (photopic) between the WT and α7 KO mice (Fig. 5).

Subsequently, we used PERG recordings to rule out the possibility that the reduced visual acuity of the α7 KO mice may have been due to altered retinal function. Pattern stimulation at 1-Hz reversal generated a transient retinal response that consisted of a smaller positive peak at approximately 90 ms, and a major negative component with peak latency at 150 ms (Fig. 6A). By measuring the peak-to-trough amplitude in response to increasing spatial frequencies we obtained an estimate of retinal spatial resolution limit (Fig. 6A) using a method previously described for VEPs. The retinal spatial resolution limit measured in the control mice (0.64 ± 0.03 cycle/deg, n = 5) was similar to the cortical spatial resolution limit determined by VEPs, and not statistically different from the retinal spatial resolution limit measured in the α7 KO mice (0.64 ± 0.02 cycle/deg, n = 5). This shows that the reduced visual acuity of α7 KO mice is not caused by an abnormal spatial resolution limit at the retinal level.

Data obtained from β2 KO mice show that their reduced visual acuity is due to an alteration in the segregation of retinofugal projections in the dorsolateral geniculate nucleus.9 To rule out this possibility, we traced the retinogeniculate projections in control and α7 KO mice by means of the monococular intravitreal injection of fluorescent cholera toxin. The contralateral stained retinogeniculate terminals were distributed at the level of the “outer shell” of the dorsolateral geniculate nucleus (dLGN) located caudodorsally (Fig. 7A), and a labeling gap was visible in the dorsomedial part of the dLGN at the level of the “inner core,” a region that receives afferents
only from the ipsilateral retina. The ipsilateral labeled fibers not crossing at the level of the optic chiasm showed a complementary labeling pattern in the “inner core” of the ipsilateral dLGN (Fig. 7A). As shown in Figure 7A, the qualitative labeling of the contralateral and ipsilateral terminals in the dLGN was comparable in the control and \( \alpha^7 \) KO mice. The labeled retinal terminals were quantitatively evaluated by means of methods previously used by Rossi et al., which showed that there was no statistically significant difference in the percentage of the dLGN occupied by the ipsilateral fibers between the control mice (13.2 ± 0.1%, \( n = 3 \)) and the \( \alpha^7 \) KO mice (12.8 ± 0.1%, \( n = 3 \)) (Fig. 7B). The segregation of the retinofugal fibers is therefore normal in the LGN regions of \( \alpha^7 \) KO mice.

Finally, we evaluated the ocularity of \( \alpha^7 \) KO mice by recording the VEP responses elicited by the contralateral and ipsilateral eyes in the two cortices. The responses driven by the contralateral eye in the binocular portion of the visual cortex (OC1b, 2.8/5 mm from lambda) are larger than the responses elicited by ipsilateral eye stimulation, and this bias in favor of the contralateral eye reflects the predominance of crossed fibers at the chiasm and the distribution of LGN fibers. We measured the ratio between contralateral and ipsilateral response amplitudes in both OC1b regions, and considered it a cortical index of the balance between the inputs of the contralateral and ipsilateral eyes (ocularity). Because there was no significant difference between the \( \alpha^7 \) KO mice (3.1 ± 0.03, \( n = 7 \)) and WT mice (2.8 ± 0.03, \( n = 9 \); Fig. 7C), the reduced visual acuity of the former is not associated with any alteration in the balance between the inputs of the two eyes (ocularity) in the primary visual cortex. In addition, we evaluated cortical retinotopy by measuring VEP amplitude profiles when the electrode was moved to different distances from lambda along the mediolateral axis (see Fig. 3D); cortical retinotopy using VEP closely corresponds to that previously reported by evaluating receptive field centers of single cortical units. It is also worth noting that VEP retinotopy was normal in the \( \alpha^7 \) KO mice.

**DISCUSSION**

It has been reported that nAChRs (especially \( \alpha^7 \) nAChRs) are involved in differentiation, neuron migration, and synapse formation during brain development, and the temporal pattern of nAChR expression suggests that they play a particular role because of their high concentration during the stage of synapse formation. The results of in vitro experiments indicate that nAChRs (particularly \( \alpha^7 \)) may control the development of neuronal architecture, stabilize synapse formation, and orient and control neurite outgrowth. Availability of mutant mice lacking individual nAChR subunits provides a unique opportunity to study their function in the nervous system.

Mice lacking the \( \alpha^7 \) nAChR subunit are viable and apparently normal in terms of their brain anatomy and performance of basic behavioral tasks, although they show some cognitive impairment in terms of episodic/working memory and susceptibility to disease. The reduced visual acuity of \( \alpha^7 \) KO mice is not associated with any alteration in the balance between the inputs of the two eyes (ocularity) in the primary visual cortex.
sensitive to nicotinic blockers in the P1–P10 time window.29

depends on retinal waves of spontaneous activity that are
refined formation of eye-specific layers at the thalamic level

One recent study has found an increase in the number of
subunit has only a slight impact on behavioral performance
suggests a possible compensatory effect in the nervous system.

Our binding and immunoprecipitation studies of 3-month-old
mice lacking the
subunit is expressed in
the mouse retina with α-bungarotoxin labeling localized at
terminals of amacrine cells on bipolar cells30 as well as in the
colliculus and in the dLGN.1 Our PERG and F-ERG experiments
excluded the possibility that the reduced visual acuity of adult
α7 KO mice is due to abnormal retinal function. This is in line
with previous findings showing that cortical but not retinal
acuity is reduced in β2 KO mice,9 and suggests that the lack of
one specific nicotinic receptor subunit may affect visual system
development but can be compensated by retinal function in
adults. Moreover, unlike that of β2 KO mice, the reduced visual acuity of α7 KO mice is not due to an alteration in the
segregation of retinal fibers, altered cortical retinotopy, or
ocular dominance in the binocular region of the visual cortex.
Previous studies have shown that both α4 and α7 mRNA levels
in rodents accumulate from P12 before eye opening to about
P55, with α7 subunit expression increasing in all cortical layers
other than layer VI soon after eye opening. This indicates that
visual cortex stimulation by a visual input is an essential step in
normal α7 nAChR expression, and suggests that these recep-
tors may play an experience-dependent role in visual cortex
maturation.9 Unlike other genetic manipulation,31 synaptic
changes in α7 KO mice reduce visual acuity without affecting

![Figure 6](https://example.com/figure6.png)

**Figure 6.** (A) Examples of PERG recordings from WT and α7KO mice at the spatial frequency of 0.06 cycle/deg; examples of the
calculation of the retinal spatial resolution limit by means of linear extrapolation to noise level (dashed line) are shown on the right. (B)
Mean spatial resolution was not significantly different between the WT (0.64 ± 0.03 cycle/deg, n = 5) and the α7KO mice (0.64 ± 0.02, n =
3). In (A), the vertical scale bar = 5 μV and the horizontal scale bar =
100 ms.

tained attention.23,24 The fact that the absence of the α7
subunit has only a slight impact on behavioral performance
suggests a possible compensatory effect in the nervous system.

One recent study has found an increase in the number of α4- and α3-containing nAChRs in the cortex and hippocampus of
α7 KO mice, which may contribute to normal brain development; however, these changes were limited to the first 21 days
of postnatal development and there was no significant differ-
cence in subunit levels between adult α7 KO and WT mice.25

Our binding and immunoprecipitation studies of 3-month-old
mice confirmed that there is no significant difference in the
levels of non-α7 subunits in the visual cortex and the retina of
WT and α7 KO mice, thus excluding the possibility that the
absence of the α7 subunit induces changes in the expression
levels of the other nicotinic receptors.

Their good performance in basic tasks allowed us to deter-
mine behaviorally the visual acuity of α7 KO mice. They were
able to learn the water maze task as quickly as the WT mice; however, despite the compensation provided by the other
nicotinic receptor subunits during cortical development,25
they were still visually impaired. Their reduced visual acuity
when performing a behavioral task was confirmed by measure-
ments of the cortical spatial resolution limit, which suggests
that the α7 subunit plays a particular role in visual system
development. It has been demonstrated that cholinergic neu-
rotransmission through the nicotinic receptors on retinal gan-
glion cells is required for retinal wave generation,26–28 and
the refined formation of eye-specific layers at the thalamic level
depends on retinal waves of spontaneous activity that are
sensitive to nicotinic blockers in the P1–P10 time window.29

Mice lacking the β2 subunit have altered retinal waves27 and,
consequently, abnormal retinofugal projections in the dorso-
lateral geniculate nucleus (dLGN) and superior colliculus that
do not segregate into eye-specific areas; they are also affected
by reduced visual acuity and altered cortical retinotopy.9

The α7 homomeric receptor is abundantly expressed in the
neocortex, hippocampus, and subcortical limbic regions.1
Concerning the visual pathways, the α7 subunit is expressed in
the mouse retina with α-bungarotoxin labeling localized at
terminals of amacrine cells on bipolar cells30 as well as in the
colliculus and in the dLGN.1 Our PERG and F-ERG experiments
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tors may play an experience-dependent role in visual cortex
maturation.9 Unlike other genetic manipulation,31 synaptic
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![Figure 7](https://example.com/figure7.png)

**Figure 7.** (A) Representative coronal sections of the contralateral and ipsilateral dLGNs (with respect to the injected eye) of WT and
α7KO mice. Labeling by means of an intraocular injection of fluores-
cently tagged cholera toxin B subunit showed that the α7KO mice
have normal retinogeniculate projections. (B) There was no difference
in the percentage dLGN area occupied by ipsilateral fibers between the
WT (13.2 ± 0.1%, n = 3) and α7KO mice (12.8 ± 0.1%, n = 3). (C)
There was also no significant difference in the ratio of contralateral and
ipsilateral VEP amplitude responses (KO, 3.1 ± 0.03, n = 7 vs. WT,
2.8 ± 0.03, n = 9).
contrast sensitivity, thus suggesting that the decreased response to high spatial frequencies may be due to an altered cortical receptive field or unbalanced excitation/inhibition at cortical synapses. One possibility is that the lack of α7 receptors at cortical synapses affects the release of other presynaptic neurotransmitters or alters calcium permeability pre- and post-synaptically. It has been clearly shown that α7 nAChRs represent a highly calcium-permeable channel that enhances calcium influx by modulating the release of neurotransmitters such as glutamate and γ-aminobutyric acid (GABA); in particular, homomeric α7 nAChRs boost the release of glutamate, thus facilitating the induction of long-term synaptic potentiation (LTP). Although the specific role of nicotinic receptors in visual cortex synaptic plasticity still needs to be thoroughly investigated, there is evidence that cholinergic system activity is involved in the plastic reorganization of neuronal connectivity in the cerebral cortex. Moreover, both LTP and long-term depression, two forms of synaptic plasticity that are thought to participate in the shaping of neuronal connections as well as learning and memory, are regulated by cholinergic transmission.

It therefore cannot be excluded that the cortical deficit in α7 KO mice may lead to the abnormal functional development of cortical circuitry as a result of defective synaptic plasticity. Our findings show that the presence of the α7 subunit during embryogenesis is not crucial for prenatal brain development, including visual system formation, but indicate that the homomeric α7 receptor plays a critical role in visual cortex refinement and functional maturation. It is worth noting that patients with homozygous or compound heterozygous deletions in 15q13.3 involving several genes (including Cbrrha7 deletions) develop neurodevelopmental disorders with severe encephalopathy, hypotonia, and visual impairment; our findings show that mice with a homozygous deletion of the gene for the α7 subunit (mice that are homozygous for the Cbrrha7−/− mutation) are affected by a visual cortical dysfunction that leads to poor vision.

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