Importance of the GluN2B carboxy-terminal domain for enhancement of social memories

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The N-methyl-D-aspartate (NMDA) receptor is known to be necessary for many forms of learning and memory, including social recognition memory. Additionally, the GluN2 subunits are known to modulate multiple forms of memory, with a high GluN2A:GluN2B ratio leading to impairments in long-term memory, while a low GluN2A:GluN2B ratio enhances some forms of long-term memory. Here, we investigate the molecular motif responsible for the differences in social recognition memory and olfactory memory in the forebrain-specific transgenic GluN2A overexpression mice and the forebrain-specific transgenic GluN2B overexpression mice by using two transgenic mouse lines that overexpress chimeric GluN2 subunits. The transgenic chimeric GluN2B subunit mice were tested for their ability to learn and remember fruit scents, male juveniles of the same strain, females of the same strain, male juveniles of another strain, and rodents of another species. The data presented here demonstrate that the GluN2B carboxy-terminal domain is necessary for enhanced social recognition memory in GluN2B transgenic overexpression mice. Furthermore, the GluN2A carboxy-terminal domain is responsible for the impaired long-term olfactory and social memory observed in the GluN2A overexpression mice.

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The N-methyl-D-aspartate (NMDA) receptor is known to play a crucial role in multiple forms of learning and memory including object recognition memory, emotional memory, social memory, extinction memory, and spatial memory (McHugh et al. 1996; Tonegawa et al. 1996; Tsien et al. 1996; Rampon et al. 2000a,b; Szapiro et al. 2003; Cui et al. 2004, 2005; Li and Tsien 2009; Wang et al. 2009; Kuang et al. 2010; Zimmerman and Maren 2010; Mei et al. 2011; Zhang et al. 2013).

NMDA receptors have been shown to be necessary for social memory in mice. Direct evidence of this was found in a post-natal deletion of GluN1 in which the knockout animals were unable to remember the stimulus partner for even a short amount of time in a habituation–dishabituation paradigm (Belforte et al. 2010). Further, Gao et al. (2009) found that both short- and long-term social recognition memory were impaired in mice treated with the NMDA antagonist ketamine. Similarly, MK-801 administration also impaired social recognition memory (van der Staay et al. 2003; Cui et al. 2004, 2005; Li and Tsien 2009; Mei et al. 2011; Zhang et al. 2013). The modulation of the GluN2B subunit composition in the NMDA receptor complex also modulated olfactory memory, and social recognition memory (White and Youngentob 2004; Jacobs and Tsien 2012, 2014). Interestingly, genetically increasing the GluN2B/GluN2A subunit ratio increases the ability for the transgenic GluN2B overexpression mice to learn and remember mice of another strain and animals of another rodent species (Jacobs and Tsien 2012). Remarkably, decreasing the GluN2B/GluN2A subunit ratio significantly impairs long-term social recognition memory of conspecifics, females, mice of another strain and rodents of another species (Jacobs and Tsien 2014).

Although it is known that an increased GluN2B/GluN2A subunit ratio results in increased object recognition memory, emotional memory, social memory, and spatial memory, little is known about the exact molecular mechanism responsible for this enhancement. It is of great interest to determine if the enhanced memory in the GluN2B transgenic animals is a result of the amino-terminal domain or the carboxy-terminal domain function. While GluN2 knockout models provide a platform for identifying the contribution for individual subunits by eliminating the unit completely (Ikeda et al. 1995; Sakimura et al. 1995; Kutsuwada et al. 1996; Hillman et al. 2011), a complete GluN2B knockout is perinatally lethal, thereby preventing analysis of the adult brain functions (Kutsuwada et al. 1996). Open channel blockers and antagonists act on the receptor complex which...

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does not allow for the separation of the amino-terminal and membrane domain functions from the carboxy-terminal domain functions (Albensi and Ilkanich 2004; Wyllie et al. 2013; Vyklicky et al. 2014). Approaches to make chimeric subunits provide an alternative means to assess how the amino terminus and carboxy terminus would contribute to memory enhancement or impairment. Recently, it was determined, using transgenic mice overexpressing GluN2 chimic subunits, that the GluN2B carboxy-terminal domain was necessary for the enhanced object recognition memory, long-term emotional memory and extinction memory observed in the transgenic GluN2B overexpression mice regardless of whether it was coupled to the GluN2B or GluN2A amino-terminal domains (Jacobs et al. 2014). On the other hand, the GluN2A carboxy-terminus would impair long-term memory when it was coupled to the GluN2B amino-terminal domain.

While social memory is highly important for survival, it has been much less studied using genetic approaches. It is often thought that social recognition memory may involve different mechanisms than other types of memory (van der Kooij and Sandi 2012; Olson et al. 2013). In this study, we investigated the molecular mechanisms responsible for the enhanced social recognition memory seen in the GluN2B transgenic overexpression mice, as well as the long-term memory impairments observed in the transgenic GluN2A overexpression mice, by using transgenic mouse lines which overexpress chimeric GluN2 receptors. In one of the chimeric mouse lines the GluN2A amino-terminal and membrane domain is paired with the GluN2B carboxy-terminal domain (hereafter referred to as the GluN2A2B(CTD) mice). The second chimeric mouse line overexpresses chimeric GluN2 receptors with the GluN2B amino-terminal and membrane domain with the GluN2A carboxy-terminal domain (hereafter referred to as the GluN2B2A(CTD) mice). Both lines have expression of the chimeric subunits targeted to the forebrain areas. We tested the ability of the chimeric mice to learn and remember nonsocial olfactory (fruit) scents, male juvenile conspecifics, female conspecifics, mice of a different strain, and rodents of a different species, the rat.

Results

Nonsocial olfactory recognition

Because of the importance of olfaction in social recognition, we first tested the transgenic mice in an olfactory recognition task. This testing paradigm was similar to the social recognition paradigms in that one scent is presented at each trial. In the training session for the 1-h retention experiment, all groups of mice tested displayed similar investigation times of the scent (GluN2A2B(CTD): n = 12, 47.28 ± 7.03 sec; GluN2B2A(CTD): n = 18, 57.01 ± 3.32 sec; Wt: n = 16, 63.79 ± 10.41 sec; Fig. 1A). After training, the mice were split into two groups, one cohort was given the identical scent as in the training session (familiar scent), the second cohort was given a novel scent at the 1-h retention session. At the retention session all three groups of mice presented with the familiar scent showed significant reductions in the exploration times of the scent (GluN2A2B(CTD): n = 6, 16.82 ± 4.08 sec, P ≤ 0.01; GluN2B2A(CTD): n = 9, 26.12 ± 7.92 sec, P ≤ 0.01; Wt: n = 8, 30.61 ± 4.17 sec, P < 0.05) indicating a memory of the scent from the training session. Interestingly, the GluN2A2B(CTD) mice spent significantly less time investigating the familiar scent than the wild-type mice (P = 0.04). As expected, the two transgenic mouse groups and their wild-type littermates did not decrease the amount of time they spent investigating the novel scent (GluN2A2B(CTD): n = 6, 58.32 ± 13.62 sec; GluN2B2A(CTD): n = 9, 50.28 ± 11.57 sec; Wt: n = 8, 56.36 ± 16.55 sec).

We also investigated the long-term olfactory memory in our GluN2A2B(CTD) and GluN2B2A(CTD) chimeric mice. Similar to the 1-h testing, all groups of mice spent similar amounts of time exploring the scent in the training session (GluN2A2B(CTD): n = 12, 48.92 ± 5.60 sec; GluN2B2A(CTD): n = 17, 54.39 ± 5.18 sec; Wt: n = 16, 68.36 ± 9.22 sec; Fig. 1B). After training, the mice were split into two cohorts, one cohort was presented with the familiar scent at the 24-h retention session; the other was presented with a novel scent. Interestingly, the GluN2B2A(CTD) mice were unable to form a strong memory of the scent and spent a similar amount of time investigating the familiar scent as they did in the training session (GluN2B2A(CTD): n = 8, 42.86 ± 12.50 sec, P = 0.3). However the GluN2A2B(CTD) mice and the wild-type mice did significantly reduce the amount of time they spent investigating the
familiar scent (GluN2A\textsubscript{2B(CTD)}: \(n = 6\), 7.58 ± 1.73 sec, \(P \leq 0.0001\); Wt: \(n = 7\), 36.47 ± 7.89 sec, \(P < 0.05\)). Interestingly, ANOVA analysis revealed that the GluN2A\textsubscript{2B(CTD)} mice spent significantly less time investigating the familiar scent than either the wild-type mice or the GluN2A\textsubscript{2A(CTD)} (\(P = 0.03\)). When the mice were presented with a novel scent they spent similar amounts of time investigating the new scent as they did the novel scent in the training session (GluN2A\textsubscript{2B(CTD)}: \(n = 6\), 42.21 ± 10.01 sec; GluN2A\textsubscript{2A(CTD)}: \(n = 9\), 45.96 ± 4.09 sec; Wt: \(n = 9\), 69.24 ± 10.25 sec). These data demonstrate the significantly impaired long-term olfactory memory in the GluN2A\textsubscript{2A(CTD)} mice.

### Habitation–dishabitation social memory paradigm

To test the ability of our GluN2A\textsubscript{2B(CTD)} and GluN2B\textsubscript{2A(CTD)} chimeric mice to remember a conspecific that they have had several short encounters with within a short period of time, we tested our mice in a social habituation–dishabitation paradigm. The GluN2A\textsubscript{2B(CTD)} and GluN2B\textsubscript{2A(CTD)} chimeric mice and their wild-type littermates showed little differences in the habituation–dishabitation paradigm. The chimeric and wild-type mice tested spent similar amounts of time investigating the stimulus mouse in the initial exposure (GluN2A\textsubscript{2B(CTD)}: \(n = 12\), 40.63 ± 2.51 sec; GluN2B\textsubscript{2A(CTD)}: \(n = 11\), 37.07 ± 3.23 sec; Wt: \(n = 12\), 33.46 ± 2.81 s; Fig. 2). In the second exposure, all three groups spent significantly less time investigating the same juvenile in a second exposure 10 min after the first (GluN2A\textsubscript{2B(CTD)}: 28.67 ± 3.19 sec, \(P < 0.01\); GluN2B\textsubscript{2A(CTD)}: 26.92 ± 3.11 sec, \(P < 0.05\); Wt: 24.14 ± 2.21 sec, \(P < 0.05\)). All three groups continued to decrease the amount of time that they spent investigating the familiar mouse in the third and fourth exposures (3: GluN2A\textsubscript{2B(CTD)}: 22.28 ± 1.34 sec; GluN2B\textsubscript{2A(CTD)}: 17.69 ± 0.23 sec; Wt: 12.50 ± 1.60 sec, \(P < 0.001\); 4: GluN2A\textsubscript{2B(CTD)}: 10.68 ± 1.20 sec, \(P < 0.001\); GluN2B\textsubscript{2A(CTD)}: 12.88 ± 2.78 sec; Wt: 7.90 ± 1.88 sec).

As a control for fatiguing effects, a fifth trial was conducted with a novel mouse. A novel conspecific was placed into the test cage with the subject mouse for 5 and 10 min after the fourth trial. In this trial, all groups of animals tested spent significantly more time investigating the novel mouse than they did the familiar mouse in the fourth trial (GluN2A\textsubscript{2B(CTD)}: 37.52 ± 3.04 sec, \(P < 0.001\); GluN2B\textsubscript{2A(CTD)}: 31.41 ± 3.27 sec, \(P \leq 0.001\); Wt: 21.79 ± 3.32 sec, \(P < 0.001\)).

#### Social memory for male conspecifics

In order to test the short- and long-term social memory for a male conspecific, we utilized a common social recognition paradigm in which a stimulus mouse is presented to the subject mouse for 5 min. Then, at the described time either the familiar mouse is presented again or a novel mouse is presented. A significant reduction in the amount of time that the subject mouse spends investigating the familiar mouse indicates a memory of that animal.

We used two time points in our investigation, 1 h, to test short-term memory and 24 h, to test long-term memory. To test the short-term memory in our chimeric mice we presented a set of male subject mice with a juvenile male conspecific. In the training session for the 1-h memory task, all groups of mice tested spent similar amounts of time investigating the juvenile stimulus mouse (GluN2A\textsubscript{2B(CTD)}: \(n = 13\), 114.74 ± 9.50 sec; GluN2B\textsubscript{2A(CTD)}: \(n = 13\), 95.46 ± 9.12 sec; Wt: \(n = 18\), 95.63 ± 7.74 sec; Fig. 3A). After training the mice were split into two cohorts. One cohort was presented with the familiar conspecific from the training session, the second cohort was presented with a novel conspecific. The subject mice presented with a familiar conspecific spent significantly less time exploring the familiar animal in the retention session than in the training session (GluN2A\textsubscript{2B(CTD)}: \(n = 7\), 74.78 ± 8.91 sec, \(P < 0.01\); GluN2B\textsubscript{2A(CTD)}: \(n = 6\), 53.31 ± 10.49 sec, \(P < 0.01\); Wt: \(n = 10\), 62.19 ± 9.05 sec, \(P < 0.01\)). As expected, the subject mice spent similar amounts of time investigating the novel stimulus mouse as they did in the training session (GluN2A\textsubscript{2B(CTD)}: \(n = 6\), 115.15 ± 10.48 sec; GluN2B\textsubscript{2A(CTD)}: \(n = 7\), 92.62 ± 12.04 sec; Wt: \(n = 8\), 85.56 ± 14.62 sec).

In the 24-h test training session, all groups of mice tested spent similar amounts of time investigating the juvenile conspecific (GluN2A\textsubscript{2B(CTD)}: \(n = 10\), 96.94 ± 9.36 sec; GluN2B\textsubscript{2A(CTD)}: \(n = 15\), 99.95 ± 5.96 sec; Wt: \(n = 17\), 89.59 ± 6.70 sec; Fig. 3B). Interestingly, when the GluN2B\textsubscript{2A(CTD)} mice were presented with a familiar conspecific, they spent nearly the same amount of time investigating the familiar conspecific in the retention session as they did in the training session (GluN2B\textsubscript{2A(CTD)}: \(n = 8\), 87.43 ± 7.27 sec). However, both the GluN2B\textsubscript{2A(CTD)} chimeric mice and their wild-type littermates do show significant decreases in the exploration times of the familiar juvenile (GluN2A\textsubscript{2B(CTD)}: \(n = 5\), 57.23 ± 9.36 sec, \(P < 0.05\); Wt: \(n = 8\), 46.48 ± 7.22 sec, \(P < 0.001\)). Interestingly, AVONA analysis revealed that the GluN2B\textsubscript{2A(CTD)} mice spent significantly more time with the familiar juvenile than did either the GluN2A\textsubscript{2B(CTD)} mice or the wild-type mice (\(P < 0.01\)). As expected, all groups of mice displayed similar exploration times with a novel juvenile 24 h after the training session (GluN2A\textsubscript{2B(CTD)}: \(n = 5\), 111.28 ± 7.08 sec; GluN2B\textsubscript{2A(CTD)}: \(n = 7\), 97.50 ± 12.02 sec; Wt: \(n = 9\), 86.80 ± 11.59 sec). These data indicate that the GluN2A\textsubscript{2A(CTD)} chimeric mouse are unable to form a long-term memory of the male conspecific, while the GluN2B\textsubscript{2B(CTD)} chimeric mice and their wild-type littermates can form strong memories of a juvenile male conspecific, that can last at least 24 h.

#### Social memory for female conspecifics

We also investigated the ability of our transgenic GluN2A\textsubscript{2B(CTD)} and GluN2B\textsubscript{2A(CTD)} mice to form a memory of a female conspecific. We tested the short- and long-term memory durations were tested. In the training session for the 1-h memory test, all groups of mice spent

![Figure 2](image_url)
the GluN2A2B(CTD) and wild-type mice spent significantly less time investigating the familiar male compared to the familiar juvenile, whereas the GluN2B2A(CTD) and wild-type mice spent similar amounts of time investigating the familiar juvenile as in the training session, whereas the GluN2A2B(CTD) and wild-type mice spent significantly less time investigating the familiar male conspecific than in the training session (GluN2A2B(CTD): n = 8, 64.34 ± 9.39 sec, P ≤ 0.001; Wt: n = 10, 62.97 ± 6.39 sec, P ≤ 0.001). Interestingly, AVONA analysis revealed that the GluN2B2A(CTD) mice spent significantly more time with the familiar female than did either the GluN2A2B(CTD) mice or the wild-type mice (P ≤ 0.001). As expected, all of the mice tested spent similar amounts of time investigating the second novel female mouse (GluN2A2B(CTD): n = 8, 164.71 ± 6.00 sec; GluN2B2A(CTD): n = 7, 170.00 ± 7.67 sec; Wt: n = 9, 174.38 ± 6.41 sec).

Figure 3. GluN2B2A(CTD) mice are impaired in long-term social recognition of a male conspecific. (A) All three groups of mice tested significantly decreased the exploration of the familiar juvenile male in the 1-h retention session. (B) The GluN2B2A(CTD) mice were unable to recall the familiar juvenile after 24 h and spent a similar amount of time investigating the familiar juvenile as in the training session, whereas the GluN2A2B(CTD) and wild-type mice spent significantly less time investigating the familiar male conspecific. (*) P < 0.05, (**) P ≤ 0.01, (***) P ≤ 0.001.

Social recognition across mouse strains

Animals in the wild may encounter other strains of their species and may compete with them for housing and food (Groning and Hochkirch 2008; Ord and Stamps 2009). In order to preserve their own colony and strain, mice are more likely to mate within their strain (Macbeth et al. 2009). In order to determine the ability of our transgenic mice to form a memory of a mouse of another strain, we tested the animals in a short-term (1 h) and long-term (24 h) social recognition task using white BALB/c juvenile male mice as the stimulus.

In the training session for the 1-h memory test, all groups of mice spent approximately the same amount of time investigating the white juvenile male (GluN2A2B(CTD): n = 12, 141.29 ± 9.47 sec; GluN2B2A(CTD): n = 15, 128.68 ± 7.70 sec; Wt: n = 13, 150.35 ± 7.72 sec; Fig. 3A). The GluN2A2B(CTD) mice, GluN2B2A(CTD) mice, and their wild-type littermates were able to form a short-term memory of the white BALB/c mouse, showing significant reductions in the exploration times in the 1 h retention session (GluN2A2B(CTD): n = 6, 44.30 ± 8.51 sec, P ≤ 0.001; GluN2B2A(CTD): n = 8, 77.51 ± 17.08 sec, P ≤ 0.01; Wt: n = 6, 43.86 ± 7.10 sec, P ≤ 0.001). When presented with a novel mouse after 1 h, the second cohort of mice showed no decrease in their investigation times of the novel stimulus mouse (GluN2A2B(CTD): n = 6, 125.23 ± 9.13 sec; GluN2B2A(CTD): n = 7, 124.24 ± 10.08 sec; Wt: n = 7, 133.92 ± 7.00 sec).

We also investigated the ability of the chimeric overexpression mice to remember a mouse of a different strain and fur color for 24 h. In the initial training session, all three groups spent similar amounts of time exploring the BALB/c stimulus mouse (GluN2A2B(CTD): n = 12, 121.81 ± 11.13 sec; GluN2B2A(CTD): n = 16, 135.43 ± 8.39 sec; Wt: n = 15, 138.54 ± 9.32 sec; Fig. 5B). Interestingly, at the retention session, neither the GluN2A2B(CTD) mice nor the wild-type mice were able to form a strong memory of the BALB/c mouse and showed no reduction in the investigation times of the stimulus mouse (GluN2A2B(CTD): n = 7, 139.21 ± 12.67 sec; Wt: n = 8, 129.63 ± 9.34 sec). Remarkably, the GluN2B2A(CTD) mice were able to form a memory of the mouse of a different strain, having significantly less exploration times in the retention session than in the training session (GluN2B2A(CTD): n = 6, 48.28 ± 14.11 sec, P ≤ 0.01). Interestingly, AVONA analysis revealed that the GluN2A2B(CTD) mice spent significantly less
time with the familiar cross-strain mouse than did either the GluN2B2A(CTD) mice or the wild-type mice (P ≤ 0.001). As expected, when another cohort was presented with a novel BALB/c mouse, the exploration times were not reduced from the initial encounter (GluN2A2B(CTD); n = 6, 122.64 ± 7.54 sec; GluN2B2A(CTD); n = 9, 141.79 ± 10.70 sec; Wt: n = 7, 132.51 ± 11.94 sec). These data demonstrate that the GluN2A2B(CTD) mice have significantly improved long-term memories for a mouse of another strain.

Figure 4. GluN2B2A(CTD) mice are impaired in the long-term social recognition of a female conspecific. (A) In the 1-h retention session, the three groups of mice tested were able to remember the previously presented female and spent less time investigating the female in the retention session. (B) Although the GluN2A2B(CTD) and the wild-type mice spent significantly less time investigating the familiar female in the 24-h retention session, the GluN2B2A(CTD) mice were unable to form a long-term memory of the female conspecific and spent as much time exploring the female in the retention session as in the training session. (*) P < 0.05, (**) P < 0.01, (****) P ≤ 0.001.

Social memory for a different rodent species
Many species are able to recognize individuals of other species (Staples et al. 2008a,b). This allows animals to remember frequent visitors to their habitat, and potential predators (Apfelbach et al. 2005; Staples et al. 2008a,b; Okamoto and Grether 2013). For this reason, we also investigated the ability of the GluN2A2B(CTD) and GluN2B2A(CTD) mice and their wild-type littermates to form a memory of another rodent species, the rat. To protect our transgenic mice overexpressing chimeric receptors with the GluN2B amino-terminal domain and the GluN2B carboxy-terminal intracellular domain is essential for enhanced social recognition memory. Transgenic mice overexpressing chimeric subunits with the GluN2B amino-terminal domain and the GluN2A carboxy-terminal domain demonstrated impaired long-term social recognition and olfactory recognition memory, similar to what was previously described in the transgenic GluN2A overexpression mice (Jacobs and Tsien 2014).

Conclusions
Here we have shown that the GluN2B carboxy-terminal intracellular domain is essential for enhanced social recognition memory. Transgenic mice overexpressing chimeric receptors with the GluN2A amino-terminal domain and the GluN2B carboxy-terminal domain showed enhanced cross-strain and cross-species recognition, similar to the GluN2B overexpression mice previously tested (Jacobs and Tsien 2012). Additionally, transgenic mice overexpressing the chimeric subunit with the GluN2B amino-terminal domain and the GluN2A carboxy-terminal domain demonstrated impaired long-term social recognition and olfactory recognition memory, similar to what was previously described in the transgenic GluN2A overexpression mice (Jacobs and Tsien 2014).
The NMDA receptor is highly expressed in the forebrain areas (Hestrin 1992b; Monyer et al. 1994; Sheng et al. 1994) and has been shown to be necessary in the formation of memory (Tsien et al. 1996; Gao et al. 2010; Gupta-Agarwal et al. 2014) and the consolidation of long-term memory (Shimizu et al. 2000; Santini et al. 2001; Roesler et al. 2005; Wu et al. 2007; Liu et al. 2014). While social memory is a different form of memory involving different mechanisms than olfactory memory, several of the same brain areas are involved in both types of memory. In general, there are two major olfactory subsystems, the main olfactory system and the vomeronasal system. The main olfactory system detects mainly volatile odorants and MHC molecules (Spehr et al. 2006) via the main olfactory epithelium which projects to the accessory olfactory bulb as well as the main olfactory bulb and anterior olfactory nucleus. The vomeronasal system is important for the detection of nonvolatile pheromones and major histocompatibility complex (MHC) molecules (Leinders-Zufall et al. 2004) by activation of the vomeronasal epithelium which projects to the accessory olfactory bulb (Matsuo et al. 2015). The two subsystems provide complementary information when the animal is confronted by another animal. Deficits in long-term olfactory memory, such as those observed in the GluN2B<sub>2A(CTD)</sub> mice are likely to contribute to deficits in long-term social memory. Additionally, the transgene used in the GluN2B<sub>2A(CTD)</sub> mice and GluN2A<sub>2B(CTD)</sub> mice is targeted to the forebrain areas (Tang et al. 1999; Cui et al. 2013; Jacobs et al. 2014) which include the main olfactory bulb and accessory olfactory bulb as well as several downstream targets of these areas including the amygdala, hippocampus, and olfactory cortex (Sanchez-Andrade and Kendrick 2009) which will also contribute to the observed impairments in social recognition. Similarly, increased long-term olfactory memory, is also likely to contribute to enhanced social recognition observed in the GluN2A<sub>2B(CTD)</sub> mice. This enhancement is due to the increased GluN2B subunit carboxy-terminal domain and intracellular signaling cascades present in these transgenic mice. Similar increases in long-term social memory have been observed in transgenic mice overexpressing the GluN2B subunit in the forebrain regions (Jacobs and Tsien 2012).

The GluN2A and GluN2B subunits are the main subunits available in excitatory synapses in the forebrain regions for receptor complex formation (Monyer et al. 1992; Sheng et al. 1994) and are ideal for coincidence detection due to their strong Mg<sup>2+</sup> dependency (Dingledine et al. 1999; Cull-Candy and Leszkiewicz 2004; Erreger et al. 2005). Recombinant GluN2B-containing complexes in vitro display longer excitatory postsynaptic potentials than the GluN2A-containing complexes (Hestrin 1992a,b; Monyer et al. 1992). Incorporation of GluN2B into the receptor complex increases the time-window for synaptic coincidence over GluN2A diheteromeric complexes. Genetic overexpression of the GluN2B subunit in the forebrain areas indeed leads to longer channel-opening duration, as well as enhanced learning and memory (Tang et al. 1999, 2001; Philpot et al. 2001; Wang et al. 2009; Cui et al. 2011). This reinforces the idea that the NMDA receptor mediates learning and memory at the synaptic level by controlling the time-window of coincidence detection. This implies that the channel-opening duration of the NMDA receptor should be the most critical feature for optimal learning and memory.

Interestingly, biochemical studies have shown that the intracellular carboxy-terminal domains of the GluN2A and GluN2B subunits do not show sequence homology and may preferentially interact with different downstream molecules (Kennedy et al. 2005). Conversely, studies using genetically truncated GluN2 intracellular carboxy-terminal domains demonstrate that the carboxy-terminal connections are essential for receptor function (Sprenge et al. 1998). Interestingly, the phenotype of each carboxy-terminal truncated mouse very closely resembled that of a GluN2B subunit knockout mouse line. Most relevant to the present study are the GluN2B<sub>2C/ΔC</sub> and the GluN2A<sub>2C/ΔC</sub> mice. In homozygous GluN2B<sub>2C/ΔC</sub> mice, death occurred shortly after birth, as in the GluN2B<sub>ΔC/ΔC</sub> knockout mice, limiting a detailed analysis of physiology and behavior (Kutsuwada et al. 1996).
Physiological studies of GluN2B−/− knockout embryonic neocortical neurons found significantly reduced mean peak amplitude of excitatory postsynaptic currents, and reduced peak open probabilities (Mohrmann et al. 2002). Interestingly, the GluN2A−/− mice showed impaired contextual fear conditioning and altered synaptic function. This is similar to what has been observed in GluN2A−/− knockout mice as well (Sakimura et al. 1995; Ito et al. 1996). Furthermore, carboxy-terminal truncation studies have demonstrated that the carboxy-terminal domain may have a regulatory role in gating properties (Punnakkal et al. 2012), and synaptic transmission and plasticity by controlling the open probability of the receptor (Rossi et al. 2002). In addition to intracellular signaling cascades, the carboxy-terminal domain can cause alterations in the function of the amino-terminal domain through phosphorylation by cAMP/protein kinase A (PKA). Multiple phosphorylation sites along the carboxy-terminal domain have been found to modulate gating and Ca2+ receptor permeability (Aman et al. 2014; Murphy et al. 2014). PKC phosphorylation is also known to modulate binding of the aCAMKII binding the NMDA receptor complex (Gardoni et al. 2001), as well as trafficking of the receptor complex to the synapse (Chung et al. 2004; Tang et al. 2010). These data highlight the importance of the carboxy-terminal domain of the NMDA receptor both in intracellular signaling and regulation of channel properties.

The GluN2 carboxy-terminal domains are known to interact with proteins from the postsynaptic density (PSD) family of proteins affecting synaptic localization, clustering, and signal transduction (Sheng 1996; Kennedy 1997; Kennedy and Bakay 1997; Kornau et al. 1997; Steigerwald et al. 2000). Although several truncated carboxy-terminal studies have focused on the mechanisms by which the GluN2 subunits mediate NMDA receptor function, its role in learning and memory is still undefined. In addition, modulating the GluN2 receptor subunit composition has been found to modulate social recognition memory. Interestingly, a 129P2 inbred strain of mice, in which reduced Csk expression causes an elevated GluN2B subunit expression, was found to have increased social recognition memory as well as increased memory for social food preference (Sinai et al. 2012). Directly increasing the GluN2B subunit expression by genetic overexpression also increased the social recognition memory in transgenic GluN2B overexpression mice (Jacobs and Tsien 2012). Conversely, reducing the GluN2B:GluN2A subunit ratio, by genetically overexpressing the GluN2A subunit; decreased the social recognition memory in the transgenic GluN2A overexpression mice (Jacobs and Tsien 2014). However, despite accumulating evidence suggesting the importance of the GluN2 subunits for the modulation of learning and memory, little is known as to the actual contributions of the molecular motifs of the GluN2 subunits. This is due to the innate complexity of separating the amino-terminal and membrane functions from the carboxy-terminal intracellular signaling mechanisms in a single receptor complex.

It is difficult to determine how the carboxy-terminal affects the function of the receptor using current methods (i.e., subunit knockouts, carboxy-terminal truncations or pharmacological channel blockers). An alternative way to investigate the role of the carboxy-terminal domain on NMDA receptor function is to replace the carboxy-terminal cytoplasmic domain of one subunit with the carboxy-terminal domain of another subunit, such that the receptor remains functionally intact. This allowed for the amino-terminal and membrane domains to be coupled with the carboxy-terminal signaling cascade of another subunit. Recent studies using this technique have shown that, although some behaviors may be unique functions of either the GluN2A carboxy-terminal domain (i.e., locomotion and impulsivity) or the GluN2B carboxy-terminal domain (i.e., perceptual learning, anxiety, and motor learning), other behaviors may be attributed to both subunit carboxy-terminal domains (Ryan et al. 2013).

In summary, by overexpressing two different GluN2A or GluN2B chimeric receptors, we have investigated how distinct GluN2A or GluN2B molecular domains regulate several forms of social recognition memory. The behavioral data for the chimeric GluN2Aαβ(CTD) mice suggest that their long-term memory parallels that of our previously produced GluN2B transgenic mice (Jacobs and Tsien 2012), while the GluN2Bαβ(CTD) transgenic mice performed similarly to the GluN2A transgenic mice (Jacobs and Tsien 2014). This indicates that the carboxy-terminal domain
is the crucial domain for improved social memory function in the 
GluN2B transgenic mice. This provides strong neurogenetic evi-
dence that the intracellular signaling processes controlled by 
the carboxy-terminal domain of the GluN2B subunit is necessary 
and sufficient for achieving superior social memory functions.

Materials and Methods

Animals

For all experiments, adult (3–7 months old) male transgenic 
GluN2A2B(CTD) and GluN2B2A(CTD) overexpression mice, and their 
wild-type littermates were generated as previously described and 
backcrossed to the C57BL/6j wild-type line (Jacobs et al. 2014). 
Genotypes of the mice were determined by PCR analysis of a tail 
biopsies and using an SV-40 probe. All animals were maintained 
in a standard animal vivarium, group-housed (3–5 animals per 
cage) in standard cages, on a 12:12 h light–dark cycle under con-
trolled temperature and humidity. Mice were housed in separate 
rooms from rats. All animals were allowed free access to food 
and water, except during experimental procedures. Behavioral 
tests were conducted in a specially designed, noise-reduced, 
deepest animal behavior room. Protocols were approved by the Institutional 
Animal Care and Use Committee (IACUC) of Georgia Re-
gents University, and are in strict adherence with the National 
Institutes of Health (NIH) Guide for Care and Use of Laboratory 
Animals.

The subject mouse was said to be exploring the stimulus 
when the subject mouse had its head directed toward the stimulus 
within 1 in of it, or was touching, smelling, or licking the face or 
anogenital region of the stimulus animal, or following closely be-

hind (≤1 in) it (Thor and Holloway 1982). Interacting pairs were 
closely observed during social encounters for the safety of all the 
animals. For male stimuli, juvenile male animals, ∼1 mo of age, 
were used to reduce aggression of the subject mice. For testing, 
the subject mice were separated into individual testing cages, 
identical to the housing cages, 30 min prior to the experimental 
protocol for acclimation.

Nonsocial olfactory recognition

Due to the importance of olfactory abilities in social recognition, 
we also investigated nonsocial olfactory recognition. Fruit juices 
were used as the nonsocial odor. A “scent cup” was constructed 
from a small plastic dish with five holes drilled into the top. 
Mice were allowed to habituate to the empty scent cup for 10 
min the day before testing. As described, the mouse was allowed to 
habitate to the testing cage, a standard housing home cage, 
for 30 min prior to testing. On the training day, 0.10 mL of lemon 
juice, lime juice, orange juice or pineapple juice was placed on 
a piece of filter paper inside the scent cup and the scent cup 
was sealed. The cup was placed into the testing cage. The time 
the mouse spent exploring the scent was recorded. At the de-
scribed time, the mouse was returned to the testing cage and ei-
ther the familiar scent or a novel scent was placed back into the 
cage for 5 min.

Habituation–dishabitation social memory paradigm

The first social memory paradigm that we used was the habitua-
tion–dishabitation paradigm (Dantzer et al. 1987; Winslow 
and Camacho 1995) in which the subject mouse is exposed mul-
tiple times in succession, to the stimulus juvenile C57BL/6j male 
mouse (1 mo of age). The reduced investigation times in subse-
quent exposures are a result of a short-term memory of the famil-
ilar stimulus mouse. The dishabitation phase consists of a fifth 
exposure session in which the subject mouse is exposed to a novel 
C57BL/6j juvenile male. This trial is used as an indicator of the fa-
tigue of the subject mouse.

To assess the habituation–dishabitation paradigm in our 
mice, we separated group-housed transgenic mice and their wild-
type littermates into individual testing cages 30 min prior to test-
ing. For the initial encounter, a juvenile 1-mo-old male C57BL/6j 
mouse was introduced into the cage of the adult male mouse for 1 
min. The amount of time the adult male mouse spent exploring 
the juvenile was measured, using the exploration criteria stated 
above. After 1 min, the juvenile mouse was removed from the test-
ing cage and placed into a holding cage. After a 10-min delay, 
the juvenile mouse was placed back into the testing cage with 
the subject mouse for the second trial. This was repeated for a total 
of four trials. For the fifth trial, a novel juvenile (1 mo old) C57BL/ 
6j male was introduced into the testing cage with the subject 
mouse for 1 min.

Social recognition memory

In the remaining social recognition experiments, subject mice 
were assigned a stimulus group (i.e., juvenile male conspecific, fe-
male conspecific, cross-strain mouse, or rat). Subject mice were 
separated from their home cages into individual testing cages, 
and allowed to acclimate for 30 min. A stimulus animal was placed 
into the testing cage with the subject mouse for 5 min. The subject 
mice were then split into a same animal recall group or a novel an-
imal recall group.

A significant decrease in the amount of time the subject 
spent exploring the familiar stimulus animal is used as the perfor-
ance index for measuring social memory. In a parallel retention 
session, a novel animal was placed into the testing cage with the 
subject mouse for 5 min. The subject mice were then assigned a 
stimulus group (i.e., juvenile male conspecific, female conspec-
ific, cross-strain mouse, or rat). Interactions between rats and mice were closely 
observed for the protection of the transgenic mice. The protocol 
was the same as described above.

Social memory for male conspecifics

To investigate the memory of our chimeric GluN2 mice for a male 
conspecific, 1-mo-old wild-type mice, from the same line-
age, were used as stimulus mice. The testing protocol was the 
same as described above.

Social memory for female conspecifics

We also investigated the social memory of our transgenic male 
mice for their female counterparts. To allow for exploration of fe-
males without mating behaviors, we placed female littermates (3– 
6 mo of age) into circular wire mesh enclosures. These enclosures 
were allowed for the mice to interact and sniff each other, while not al-
lowing for mounting and mating behaviors. The subject mice 
were allowed to habituate to the empty enclosure for 10 min the 
day before the testing and the day of testing, to reduce exploration 
as a result of a novel object. The testing protocol was the same as 
previously described above.

Social recognition across mouse strains

To further investigate the memory abilities of our chimeric trans-
genic mice, we tested their ability to remember a mouse of a differ-
ent strain and color. We used juvenile white BALB/c males (6–8 
wk old) as the stimulus mice. The testing protocol was the same 
as described above.

Social memory for a different rodent species

We also investigated the ability of our transgenic mice to form a 
memory of a different rodent species, the rat. To protect the sub-
ject mice from previously noted muricide by rats (Noack et al. 
2010), we constructed a protective enclosure of wire mesh around 
the rat. The enclosure allowed for access to all four sides and the 
top of the rat. The mice were habituated to the empty enclosure 
for 10 min in the testing cage the day before, and the day of the 
training session. Long-Evans hooded rats (4 wk old) were used as 
stimulus animals. Interactions between rats and mice were closely 
observed for the protection of the transgenic mice. The protocol 
was the same as stated above.

Statistical analysis

Graphed data are mean ± SEM. Paired Student’s t-tests were 
used to analyze significance of initial exploration versus recall
exploration. ANOVA analysis was used to evaluate differences among groups. Significance level was P < 0.05.

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