Postnatal development of dendrodendritic inhibition in the mammalian olfactory bulb

Shelby B. Dietz*, Foivos Markopoulos and Venkatesh N. Murthy*

Department of Molecular and Cellular Biology, Center for Brain Science, Harvard University, Cambridge, MA, USA

Edited by:
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*Correspondence:
Shelby B. Dietz, Department of Neurobiology and Behavior, Cornell University, Seeley Muudd Hall, Ithaca, NY 14853, USA.
e-mail: sdietz@cornell.edu;
Venkatesh N. Murthy, Department of Molecular and Cellular Biology, Harvard University, 16 Divinity Avenue, Cambridge, MA 02138, USA.
e-mail: vnmurthy@fas.harvard.edu

The mitral–granule cell (MC–GC) reciprocal synapse is an important source of auto- and lateral-inhibition in the olfactory bulb (OB), and this local inhibition is critical for odor discrimination. We may gain insight into the role of MC autoinhibition in olfaction by correlating the functional development of the autoinhibition with the postnatal development of olfactory function. We have studied the functional development of the MC–GC reciprocal synapse using whole-cell patch-clamp recordings from MCs and GCs in acute OB slices from 3- to 30-day-old rats. The magnitude of dendrodendritic inhibition (DDI) measured by depolarizing a single MC and recording recurrent inhibition in the same cell increased up to the fifteenth day of life (P15), but dropped between P15 and P30. The initial increase and later decrease in DDI was echoed by a similar increase and decrease in the frequency of miniature inhibitory post-synaptic currents, suggesting an accompanying modulation in the number of synapses available to participate in DDI. The late decrease in DDI could also result, in part, from a decrease in GC excitability as well as an increase in relative contribution of N-methyl d-aspartate (NMDA) receptors to γ-amino butyric acid (GABA) release from GC synapses. Changes in release probability of GABAergic synapses are unlikely to account for the late reduction in DDI, although they might contribute to the early increase during development. Our results demonstrate that the functional MC–GC circuit evolves over development in a complex manner that may include both construction and elimination of synapses.

Keywords: olfaction, autoinhibition, mitral

INTRODUCTION

The olfactory bulb (OB) is the first processing center for odor information in the mammalian brain (Shepherd and Greer, 1998; Mori et al., 1999; Schoppa and Urban, 2003). Mitral and tufted cells receive excitatory input from the olfactory receptor neurons in the olfactory epithelium and project their axons to the olfactory cortex and other higher brain regions. Mitral cell (MC) firing is modulated in the bulb by inhibition from glomerular interneurons, which form GABAergic synapses on the primary dendritic tufts of MCs (Parrish-Aungst et al., 2007; Shao et al., 2009), as well as from local GABAergic interneurons called granule cells (GCs) that form reciprocal synapses with MC lateral dendrites (Rall et al., 1966; Price and Powell, 1970; Yokoi et al., 1995). Local inhibition is critical for odor discrimination in insects and vertebrates (Stopfer et al., 1997; Gheusi et al., 2000; Nusser et al., 2001; Abraham et al., 2010).

In rats, some olfactory function is present at birth and is necessary to initiate nursing (Teicher and Blass, 1977; Brunet et al., 1996). Pups are capable of appetitive learning at P1 (Johanson and Hall, 1979), can respond to odors with sniffing and increased respiratory rates at P2 (Welker, 1964), and can perform simple odor discriminatory tasks as early as P3 (Armstrong et al., 2006). These early olfactory abilities manifest well before the number of GCs and the number of MC–GC reciprocal synapses approach adult levels. The number of GCs rises from 10% of adult levels at birth to nearly 100% at P30, whereas MCs are slightly overabundant at birth and are pruned away to form the adult complement of cells by P10–20 (Rosselli-Austin and Altman, 1979; Frazier and Brunjes, 1988). There have been few anatomical studies of synapse formation during development in the OB; the best available measurements were made in mice by Hinds and Hinds (1976) using electron microscopy. MC-to-GC synapse density in the external plexiform layer (EPL) starts at about 10% of the adult level at P0 and reaches the adult level at about P15; GC-to-MC synapse density lags behind slightly, reaching 90% of adult levels by P30 but continuing to increase into adulthood (Hinds and Hinds, 1976). In other areas of the brain, anatomical studies can over- or underestimate the number of functional synapses (Pun et al., 1986; Isaac et al., 1995; Liao et al., 1995; Durand et al., 1996), and physiological studies have been necessary to demonstrate functionality.

The development of the mitral–granule reciprocal synapse is also important in light of the adult neurogenesis of GCs. The OB is one of two known brain areas in which newborn neurons are found throughout adulthood (Lledo et al., 2005). New GCs are generated in the subventricular zone and migrate to the bulb, where they form functional synapses and assume adult-appearing morphology (Carlen et al., 2002; Carleton et al., 2003). Comparison of the development of the mitral–granule synapse in the first weeks after birth with its development in adult-born cells may further our understanding of how adult-born cells can integrate themselves into a pre-existing network.
We hope to gain insight into the role of MC autoinhibition in olfaction by correlating the functional development of the synapse with the postnatal development of olfactory function. Using extracellular recordings of field potentials, Wilson and Leon (1986) found that the period of suppressed firing induced in the cells near the MC layer in vivo by antidromic stimulation of MC axons en route to the olfactory cortex increases from P5 to P15–P19, as one might expect, but then decreases in adulthood. In a related study, these same authors used indirect extracellular field potential recordings to suggest that presumed GC mediated inhibition underwent complex developmental changes (Wilson and Leon, 1987). Here, we have sought to identify the cellular and synaptic bases of developmental changes in dendrodendritic inhibition (DDI) using more direct whole-cell recordings in slices.

MATERIALS AND METHODS

Three hundred micrometer horizontal slices from the OBs of rats aged P3–30, were prepared and recordings were made as previously described (Dietz and Murthy, 2005). Briefly, slices were maintained in oxygenated, buffered saline (in mM, 119 NaCl, 26.2 NaHCO3, 11 glucose, 2.5 KCl, 1 NaH2PO4, 2.5 CaCl2, 1.5 MgCl2, pH 7.3, osmolarity 290–300 mOsm) at room temperature. Animal care and experimental procedures adhered to guidelines approved by the Harvard University IACUC and conformed to NIH guidelines. Whole-cell recordings were obtained from MC somata, using 1–2 MΩ borosilicate glass pipettes containing cesium chloride (in mM, CsCl 111, TEA-Cl 13, HEPES 20, Di-tris-P-creatine 11, Na2ATP 3, Na3GTP 0.2, EGTA 0.2) and voltage-clamped at −70 mV. For extracellular stimulation, a 200-μA biphasic stimulus of 1–50 μs was delivered through a glass monopolar electrode placed in the EPL driving GC firing. For some experiments the extracellular Mg concentration was reduced to zero. GC somata were patched with 5–7 MΩ borosilicate glass pipettes containing potassium gluconate (in mM, potassium gluconate 130, HEPES 10, MgCl2 2, MgATP 2, Na2ATP 2, GTP 0.3, NaCl 4). GC resting membrane potentials were recorded in current clamp, and the somata were held below firing threshold. GC firing properties were studied by delivering consecutive current pulses, 500 ms duration each, ranging from −20 to +150 pA with a 10-pA increment. For miniature inhibitory post-synaptic current (mIPSC) recordings from MCs, 1 μM TTX, 100 μM APV, and 5 μM CNQX were added to the bath; for evoked DDI recordings, 1 μM TTX; for extracellular stimulation recordings, 100 μM APV and 5 μM CNQX; and for GC recordings 100 μM APV, 5 μM CNQX, and 100 μM picrotoxin (PTX). Currents were analyzed using MATLAB (Mathworks, Natick, MA, USA), Clampfit (Molecular Devices, CA, USA), and MiniAnalysis (Synaptosoft, Decatur, GA, USA). Rise time of a current was defined as the time it took for a current to increase from 10 to 90% of its maximum amplitude, and decay time was defined as time to decrease from 90 to 37%. For mIPSC analysis, events with rise times between 0.5 and 12 ms, a minimum area of about 30 pA·ms, and amplitude at least three times the root mean square of the noise were accepted for analysis. All mIPSCs were visually inspected before being accepted for analysis. DDI was evoked by recording a MC in whole-cell voltage clamp and depolarizing the cell with a 100-ms voltage step from −70 to 0 mV, resulting in a barrage of IPSCs, spread over several seconds, that could be recorded with the same patch pipette in the MC soma. The DDI was quantified by integrating the current over a 2-s period after the cessation of voltage step for physiological Mg, and over a 4-s period for 0 Mg. The time constant of DDI was measured by fitting a single exponential to the decay. Statistical tests included one-way analysis of variance (ANOVA), N-way ANOVA, unpaired T-test, and pairwise comparison by Tukey’s honestly significantly different (HSD) test. All errors are reported as the SEM.

RESULTS

EVOKE DDI PEAKS AT P15, AND THEN DECLINES

We assayed the functional properties of the mitral–granule reciprocal circuit by depolarizing MCs in the presence of TTX. The resulting DDI recorded in the same MC is composed of feedback inhibition from multiple reciprocally connected GCs in the form of a barrage of IPSCs (Chen and Shepherd, 1997; Isaacson and Strowbridge, 1998; Friedman and Strowbridge, 2000; Margrie et al., 2001). We measured the extent of DDI in MCs from animals of various ages by imposing voltage steps from the holding potential of −70 to 0 mV for 100 ms, which evoked a flurry of IPSCs from reciprocally connected GCs (Figure 1B). DDI increased significantly from P3 to P15, in line with expectations from anatomical studies showing an age-dependent increase in MC–GC and GC–MC synapses over this time. There was a significant decrease in DDI after P15, dropping to neonatal levels by P30. The DDI at P12 and P15 were significantly larger than at all other ages, except P17 was not significantly larger than P20 (ANOVA p = 1.0e−4; pairwise comparison by Tukey’s HSD test, p < 0.05). The most obvious explanation for the early increase in DDI is the increase in the number of anatomically identified synapses, but other factors may also contribute; the late reduction in DDI even as synapses are added is more counterintuitive and intriguing. From this point we will focus our attention on possible mechanisms for this more unexpected late fall in DDI.

VARIATIONS IN RECORDING CONDITIONS AND INPUT RESISTANCE DO NOT ACCOUNT FOR THE CHANGES IN DDI

We first investigated possible reasons for the paradoxical drop of DDI between day 15 and 30 relating to the recording techniques and the intrinsic electrical properties of the cells. One potential reason for a decrease in DDI could be a change in input resistance of MCs or GCs. The intrinsic properties of GCs will be discussed below. Although input resistance of MCs dropped significantly over the entire age range (ANOVA p = 3.5e−12), most of the decrease occurred as DDI was increasing up to P15, and there was no significant change between P15 and P30, the period of DDI decrease (Figure 2A). Another potential explanation for the decrease is a change in the access resistance of our whole-cell recordings. All cells were compensated for access resistance, but compensation may not completely correct large resistances. Access resistance increased modestly, but significantly, with age (ANOVA p = 0.05), but as access resistance was slightly positively correlated with the extent of DDI (correlation coefficient = 0.10), the increase in access resistance in older cells was still within tolerance for quality recordings and did not compromise the experiments (Figure 2B). Insufficient diffusion of the chloride-based pipette internal solution to distal sites could contribute to reduced DDI.
if distal synapses were not exposed to the high driving force of the artificial intracellular solution; we think inadequate diffusion is unlikely since the extent of DDI did not increase significantly with the time of recording in any given cell, as might be expected if diffusion to distal sites was a factor over recordings lasting up to 1 h (data not shown). Finally, measurements of IPSCs might be confounded by changes in the length of MC lateral dendrites between P3 and P30, which could lead to increasing electrotonic filtering – this is discussed below.

CHANGES IN TIME COURSE OF DDI DO NOT CONTRIBUTE TO THE LATE REDUCTION IN DDI

Dendrodendritic inhibition evoked by voltage steps decays over a period of seconds (Figure 1; Isaacson and Strowbridge, 1998; Schoppa et al., 1998), probably due to a long and variable delay of GC release back onto the MC (Kapoor and Urban, 2006). In our analysis, we measured the extent of DDI by integrating the total charge in a long time period following the voltage step. To determine if there was a systematic change in the time
course of DDI with age, we fitted a single exponential to the average time course of DDI at different ages (Figure 2C). The decay time constant was similar to that reported by Isacson and Strowbridge (1998) and did not change significantly with age (ANOVA $p = 0.37$).

**CHANGES IN THE NUMBER OF SYNAPSES PARTICIPATING IN DDI MAY CONTRIBUTE TO THE INCREASE AND REDUCTION IN DDI**

The changes in DDI could also be due to a change in the number or the strength of the MC–GC synapses available to participate in DDI. An increase in the number of synapses would doubtless increase the DDI evoked by a current step. Furthermore, the same number of synapses could mediate increased DDI if they gained in strength, either presynaptically or post-synaptically. Estimating the number of synapses available for DDI and assessing their strength using electrophysiological methods requires the combination of several measurements. We began by first recording mIPSCs in MCs in the presence of TTX. The number and strength of these miniature events are related to the number and strength of synapses onto a cell (Turrigiano and Nelson, 2004). The frequency of mIPSCs can be modeled to depend on the product of the release probability $p$ and the number of release sites $n$ (Kerchner and Nicoll, 2008). An increase in mIPSC frequency could result from an increase in the number of synapses onto a cell, an increase in release probability, or an increase in the number of docked vesicles per synapse (Malenka and Nicoll, 1997; Lisman et al., 2007). We first measured changes in mIPSC frequency and properties; we then investigated some possible contributors to changes in frequency, such as release probability and silent synapses, discussed in the following sections. If other contributors to changes in mIPSC frequency can be ruled out, the change in frequency can then be ascribed to a change in synapse number. Other measurements derived from mIPSCs, such as amplitude and time course, can indicate changes in synaptic strength.

The frequency of mIPSCs remained steady at about 2.5 Hz from P3 to P12, peaked at 6.1 ± 0.42 Hz at P20, then dropped back down to about 2.4 ± 0.52 at P30 (Figure 3). The frequency at P20 was significantly higher than at P3, 6, 9, and 30 (ANOVA $p = 0.0012$, pairwise comparison by Tukey’s HSD test, $p < 0.05$). The amplitude of mIPSCs increased significantly with age (ANOVA $p = 0.007$), being higher at P30 than at P9 ($p < 0.05$), with most of the increase occurring after P15. Rise time peaked at P12, increasing from 2.4 ± 0.22 ms at P3 to 3.4 ± 0.21 ms at P12, then dropping back to 2.5 ± 0.21 by P30 (ANOVA $p = 0.0012$); the P12 value was significantly higher than those at P3 and P30, while the P15 value was significantly higher than the P3 value ($p < 0.05$). Decay time did not change significantly (Figure 3D; ANOVA $p = 0.42$). We obtained a rough estimate of the number of quanta contributing to DDI at each age by dividing the mean area of the evoked DDI by the mean area of the mIPSC for each age. As a result, the increase in the number of quanta per DDI from P3 to P15, and the subsequent decrease from P15 to P30 were even more striking than the total charge per DDI (Figure 3E). The approximate numbers of quanta in the DDI barrage were 269 at P3, 1143 at P15, and 150 at P30. The changes in mIPSC frequency suggest that either the number of available synapses, the number of release sites per synapse, or the release probability may rise and fall at roughly the same time as the DDI.

We also looked for any contribution of changes in electrotonic decay to the measured mIPSC parameters (Figure 3F). Assuming that rise times will be more readily high-pass filtered than decay times (Magee and Cook, 2000), and that rise times are less likely than decay times to be dependent on GABA_A receptor subunit assemblies (Hajos and Mody, 1997), we calculated the variance in rise times of the individual mIPSCs recorded in six cells each at P3, P15, and P30 and found no change with age. Increasing heterogeneity of electrotonic locations should lead to increasing variance of rise times when mean rise times are similar across ages. We also computed the correlation between the rise times and amplitudes of individual mIPSCs recorded in six cells each at P3, P15, and P30. We found a positive correlation between rise time and amplitude, significantly increasing from P3 to P30 (mean correlation coefficient $0.03 ± 0.06$ at P3, $0.17 ± 0.05$ at P15, and $0.38 ± 0.08$ at P30, ANOVA $p = 0.007$). One would expect that filtering would make mIPSCs from distal synapses appear artificially slow and small; the fact that the slower events tend also to be larger in P15 and P30 cells, suggests that either those distal synapses have unexpectedly large amplitudes or the distal events are not being detected at all. If distal mIPSCs are indeed not electrically detectable at the soma, their absence would suggest that an increase in passive electronic filtering or a change in active dendritic processes as the lateral dendrites lengthen with age could contribute to the reduction of DDI in older animals.

**CHANGES IN PAIRED-PULSE RATIO MAY CONTRIBUTE TO THE EARLY INCREASE IN DDI**

Another candidate for the reduction in the DDI between P15 and P30 is a decrease in release probability. If the release probability at either the MC-to-GC excitatory synapse or the GC-to-MC excitatory synapse was reduced, a voltage step in a MC would evoke less DDI in the older animal. A direct measurement of release probability is quite challenging. However, the depletion model of synaptic depression predicts that an increase in initial probability of release ($P$) at a synapse results in a decrease in paired-pulse ratio (PPR; Zucker and Regehr, 2002). At other rodent synapses, the PPR can increase (Bolshakov and Siegelbaum, 1995), decrease (Mori-Kawakami et al., 2003), or remain unchanged (Hsiat et al., 1998) during development. Physical substrates for a change in release probability can include a change in any parameter starting with action potential (AP) shape, calcium dynamics, vesicle pool properties, and exocytic machinery (Atwood and Karunanithi, 2002; Branco and Staras, 2009). To look for a developmental change in initial release probability, we tested the PPR of the granule-mitral synapse at ages P3, 15, and 30. We have shown previously that the combined mitral–granule synapse, as measured with DDI, displays paired-pulse depression at P9–17; we have also shown that the two sides of the synapse can be isolated using extracellular stimulation in the EPL (Dietz and Murthy, 2005). The GC-to-MC side of the synapse is a promising place to look for changes in release probability for two reasons: first, because the GC-to-MC synapse displayed characteristic paired-pulse depression that probably dominates the circuit behavior at short interstimulus intervals, and second, because a reduction in
FIGURE 3 | Changes in the properties of mIPSCs with age. (A) Examples of mIPSCs recorded in mitral cells at P3, P15, and P30 in 1 μM TTX, 100 μM APV, and 5 μM CNQX. (A1) Sample recordings at each age. (A2) Individual mIPSCs recorded in a different cell are shown aligned by their onset (blue), and the average mIPSC is shown in black. (B–D) mIPSC frequency, amplitude, rise time and decay time are shown as functions of age. \( N = 11 \) cells for P3, P6, P9, and P30; \( N = 12 \) for P12, P15, and P20. The frequency of mIPSCs remains steady for P3–P12, peaks at P20, and then drops back down at P30. The frequency at P20 is significantly higher than those at P3, 6, 9, and 30 (ANOVA \( p = 0.0012 \), pairwise comparison by Tukey’s HSD test, \( p < 0.05 \)). The amplitude of mIPSCs increases significantly with age, being significantly higher at P30 than at P9 (Tukey’s \( p < 0.05 \)). Rise time peaks at P12; the P12 value is significantly higher than those at P3 and P30, while the P15 value is significantly higher than the P3 value (ANOVA \( p = 0.0012 \), Tukey’s \( p < 0.05 \)). Decay time does not change significantly (ANOVA \( p = 0.42 \)). (E) The number of quanta in a single DDI trial was obtained by dividing the integrated charge in the DDI by the integrated charge in the average mIPSC at each age. Note that the increased amplitude of the P30 mIPSCs outweighs the moderate speeding of its decay time to give it a larger area than the P3 and P15 mIPSCs. For clarity, data for only three representative ages are shown. Error bars are not included as the values are derived from other measurements. (F) Electrotonic filtering is not a source of variability in mIPSC kinetics. (F1) Mean variance of rise time vs. age. The variance of the rise times of all mIPSCs recorded in a particular cell is calculated, and the average variance for each age is plotted (\( N = 6 \) for each age). The variance does not change with age (ANOVA \( p = 0.12 \)). (F2) Mean correlation coefficient of rise time and amplitude as a function of age. The correlation coefficient of rise time and amplitude for all mIPSCs recorded in a particular cell is calculated, and the average value for each age is plotted (\( N = 6 \) for each age). The positive correlation increases significantly with age (ANOVA \( p = 0.007 \)).
release probability at the GC-to-MC synapse would also explain the reduced frequency of mIPSCs recorded in MCs. Here, we delivered two stimuli 100 ms apart and measured the PPR as the ratio of the peak of the second response to the first (Figure 4). At P3, the synapse displayed paired-pulse facilitation (an average PPR of 1.27 ± 0.27), while at P15 and P30, it displayed depression (PPRs of 0.52 ± 0.10 and 0.35 ± 0.08, respectively; the P3 value was significantly different from the P15 and P30 values, ANOVA $p = 0.0039$; pairwise Tukey’s HSD, $p < 0.05$ and 0.01, respectively). At the granule-to-mitral synapse, the decrease in PPR suggests an increase in P, which could partially account for the increase in evoked DDI between P3 and P15, but does not account for the decrease in evoked DDI or mIPSC frequency between P15 and P30.

**CHANGES IN THE BALANCE OF AMPA AND NMDA CURRENTS MAY CONTRIBUTE TO THE LATE REDUCTION IN DDI**

Another mechanism that could account for the decrease in DDI with age would be an increase in the proportion of NMDA-only silent synapses at the excitatory MC-to-GC side of the reciprocal synapse. Silent synapses would appear anatomically, but would not be activated by glutamate release unless the magnesium block of the NMDA receptors on the post-synaptic GC were relieved. The magnesium block would normally be relieved by depolarization of the post-synaptic cell in vivo, but in vitro the block can be relieved simply by recording in magnesium-free extracellular solution. If DDI in 0 Mg did not show the decrease after P15, we would conclude that some of the synapses appearing late in development were indeed NMDA-only silent synapses.

In 0 Mg, peak currents of up to 10 nA could be evoked with a 100-ms step to 0 mV (Figure 5). The mean current evoked in physiological Mg (1.5 mM) was 10% of the mean current recorded in 0 Mg at P15. NMDA-mediated currents are known to be unusually important for fast synaptic transmission at this synapse when their Mg block is relieved by concomitant AMPA receptor activation; as a result of this strong influence of NMDA receptors we found DDI to be greatly enhanced in the absence of Mg at all ages, as has previously been shown in older animals (Isaacson and Strowbridge, 1998; Schoppa et al., 1998). We measured DDI at five age groups in 0 Mg. Evoked DDI peaked at P20, with DDI changing significantly with age and P15 being significantly larger than P3–6 (ANOVA $p = 0.044$, pairwise Tukey’s HSD, $p < 0.05$). The DDI data for physiological Mg and 0 Mg are plotted in Figure 5B; they are also shown normalized to their respective values at P3 in Figure 5C, with the data obtained with physiological Mg from Figure 1 binned to compare with the 0 Mg data. The curves were similar in shape, but the effect in the absence of Mg was much less pronounced. It is also interesting to note that the peak DDI arrives at P20 in 0 Mg, later than the P15 in physiological Mg. If these large currents evoked in 0 Mg were saturating, the largest currents at P15 and P20 might be blunted and the effect of age on DDI masked. However, a shorter, 5 ms voltage step in these same cells evoked smaller currents that displayed the same age-dependent curve (Figure 5B). Since the phenomenon of reduced DDI between P15–20 and P30 was greatly dampened in 0 Mg, an increase in the relative contribution of NMDA receptors may indeed contribute to the reduction in DDI between P15 and P30.

**GC EXCITABILITY CONTRIBUTES ONLY MODERATELY TO THE LATE REDUCTION IN DDI**

Another possible contributor to the changes in DDI would be a change in GC excitability; if the GCs become less excitable with age, a voltage step in the MC releasing the same quantity of glutamate at a given synapse may be less likely to evoke reciprocal GABA release in the GCs. In a slice preparation, increasing the excitability of GCs via blockade of mGluR activity reduces MC autoinhibition (Heinbockel et al., 2007), and decreasing the excitability of GCs via muscarinic receptors facilitates the long-term inhibition of MCs (Pressler et al., 2007), and decreasing the excitability of GCs via the blockade of mGluR activity reduces MC autoinhibition (Heinbockel et al., 2007). Although no direct developmental studies of GC excitability have been previously attempted, newborn GCs in adults are moderately more excitable than older cells (Saghatelyan et al., 2005), but adult-born GCs become electrically indistinguishable from older cells once they become fully mature (Carleton et al., 2003). We tested the excitability of GCs at ages P5, 15, and 30 by making whole-cell current-clamp recordings.

*FIGURE 4 | Changes in paired-pulse ratio may contribute to the early increase in DDI. (A) Drawing of the recording configuration, in which GC release is evoked by extracellular stimulation in the EPL and IPSCs recorded in the MC. (B) Sample traces of responses to pairs of extracellular stimuli in the EPL with an interpulse interval of 100 ms. Traces are averages of 10, 8, and 11 trials, respectively. (C) Average paired-pulse ratio at different ages. At P3, the synapse displays paired-pulse facilitation, while at P15 and P30, it displays depression ($N = 6$ cells for each age). The P3 value is significantly different from the P15 and P30 values, ANOVA $p = 0.0039$; pairwise Tukey’s $p < 0.05$ and 0.01, respectively).*
We have shown that the strength of DDI increases from P3 to P15, and then decreases between P15 and P30. We have tested several possible mechanisms.

The increase in DDI from P3 to P15 presumably arises largely from the increase in the number of synapses, with a contribution from an increase in release probability from GCs. The decrease in DDI from P15 to P30 even as the number of anatomically identified synapses increases, may arise from several factors. A moderate reduction in GC excitability likely contributes to the fall, but may be insufficient to account for the entire, dramatic effect. The reduction in mIPSC frequency from P20 to P30 may hold the answer. A drop in mIPSC frequency can be a result of reduced synapse number, reduced number of release sites per synapse, reduced release probability, or an increase in the proportion of synapses that are functionally silent (for example, with no functional GABA_A receptors, or with a synaptic weight near zero). We have shown that release probability in the GC-to-MC direction does not change between P15 and P30. This leaves two major mechanisms: a reduction in the number of release sites available to participate in autoinhibition, and an increase in the number of functionally silent synapses.

**POSSIBLE CHANGES IN THE NUMBER OF AVAILABLE RELEASE SITES**

How might synapses anatomically identified by microscopy methods actually be ineffective or non-functional? GC–MC synapses could be made inaccessible to the voltage-step–evoked depolarization even as the total number of synapses increases. Based on our recordings of mIPSCs, the lack of an increase in variance with age, and the absence of small and slow mIPSCs seem to suggest that distal synapses are indeed becoming electrically isolated from the soma by passive electrotonic filtering. Although our detection criteria for mIPSCs were permissive for very slow events, events of very small amplitude would have eventually disappeared below the noise. It is possible that if mIPSCs are not recordable at the soma, mIPSC-sized events from those synapses are less likely to contribute to changes in membrane potential at the soma (although DDI would be less affected by passive filtering than would individual mIPSCs, as charge transfer is significantly less sensitive to changes in electrotonic distance than are rise time and amplitude; Byrne and Shepherd, 2009). Instead, these synapses may participate in local dendrodendritic computations that are entirely separate from the somatic compartment. In addition to the effect of passive filtering, active processing in the dendrites could change with age. For example a developmental change in dynamic gating
Granule cell excitability may contribute to the late reduction in DDI. (A) Drawing of the recording configuration. Consecutive 500 ms duration current injections from −20 to +150 pA were delivered into GCs, with 10 pA increments and 5 s inter-injection intervals. (B) Sample traces of the minimum current injection to evoke action potentials (APs) in GCs with age, showing the injected current and the latency to the first spike. (C) The minimum current to evoke APs does not change significantly with age (ANOVA $p = 0.50$). (D) The latency to the first AP increases with age. The current above threshold is plotted vs. time between the onset of the current step and the first spike for ages P5, P15, and P30 (N-way ANOVA, contribution of age $p = 3.4e^{-7}$). Curves are fitted to a single exponential. Asterisks indicate time points for which the P30 values are significantly different from P5 and P15 (Tukey’s $p < 0.05$). (E) Input resistance does not change significantly with age (ANOVA $p = 0.48$). (F) Resting potentials become significantly more negative with age (ANOVA $p = 0.05$, pairwise Tukey’s $p < 0.05$ indicated by asterisk). $N = 6$ for P5; $N = 25$ for P15; $N = 12$ for P30 for (D–F), $N = 3$ for P5; $N = 16$ for P15; $N = 10$ for P30 for (C).

(Lowe, 2002; Xiong and Chen, 2002) could electrically isolate even relatively proximal synapses in older animals. Proximal GC–MC synapses could be pruned as distal synapses are added, effectively reducing the number of synapses available for DDI evoked in the soma even as the total number of synapses increases. A final possibility is that many of the synapses identified microscopically in the adult animal, despite their appearance, lack critical synaptic machinery, or have synaptic weights near zero. In future studies, immunohistochemistry or electron microscopy can be used to map the development and distribution of synapses containing specific synaptic proteins in order to address these interesting possibilities.

The reduction in DDI amplitude from P15–20 to P30 may indicate that connections between MCs are being made more
precise over this period. Odor recognition could be improved by optimization of connections between MCs that respond to odotypes associated with the same odorant molecule, or to odors that are commonly associated in the environment. Refinement of connections between MCs could help to sharpen responses to behaviorally relevant stimuli by, for example, weakening lateral inhibition between MCs that respond to related odors, enhancing their mutual ability to fire in response to that odor.

In glomerular formation, anatomical targeting is complete by birth (Meisami and Sendera, 1993; Treloar et al., 1999). Adult-like patterns of odor-evoked glomerular activation evolve only slightly from P4 to P24, but MC activation patterns become much sparser as more GCs appear in the bulb and are activated by the odor (Guthrie and Gall, 2003). Even though fewer MC–GC synapses may be activated, they may be more effective. A single proximal GC closing off propagation along an entire dendrite may do more to influence network formation than dozens of GCs located more distally, or at less critical anatomic junctions (Lowe, 2002; Xiong and Chen, 2002). Therefore, even as the total DDI evoked by voltage step decreases, effective auto- and lateral-inhibition evoked in vivo could be increasing. It is also possible that changes in MC synapses with inhibitory interneurons in the glomerular layer over development may contribute to changing DDI, although there is little evidence for a contribution of glomerular inhibition to the DDI measured in the manner reported here (Pimentel and Margrie, 2008). MCs also display some self-excitation through dendritic glutamate receptors; although the magnitude of self-excitation is small compared to the DDI evoked with symmetric chloride currents, self-excitation could also change with age and make some contribution (Aroniadou-Anderjaska et al., 1999; Isaacson, 1999; Friedman and Strowbridge, 2000).

THE CHANGING ROLE OF NMDA WITH AGE

The MC–GC reciprocal synapse is unusually dependent on NMDA receptors for fast synaptic transmission. Isaacson and Strowbridge (1998) showed that in physiological levels of magnesium, in which NMDA currents must be released from Mg-dependent voltage block by simultaneous AMPA activation, both AMPA and NMDA channels are necessary for DDI. In Mg-free solution, in which NMDA currents are relieved from voltage-dependent block, DDI is both greatly enhanced and is mediated predominantly by NMDA receptors. While both Mg-containing and Mg-free DDI peak at P15–P20, the effect in the absence of Mg is much reduced. As a result of the much greater age-dependent change in DDI in physiological Mg, the ratio of the DDI measured in physiological Mg to 0Mg conditions peaks at P15, suggesting that NMDA currents onto GCs are greater early and late in development than in the P15–P20 window. There are many possible explanations for this relative increase in the importance of NMDA. One is an increase in NMDA-only silent synapses (Kerchner and Nicoll, 2008) late in development. Although post-synaptically silent synapses can persist in adult animals (Sametsky et al., 2010), silent synapses are usually associated with early development, converting to mixed AMPA/NMDA synapses with maturity (Isaac et al., 1995; Liao et al., 1995; Durand et al., 1996; Wu et al., 1996; Li and Sheng, 2003). Another possible but as yet undemonstrated mechanism would be synapses that contain both AMPA and NMDA increasing their proportion of NMDA receptors. A reduction in the calcium permeability of AMPA receptors with age could also reduce the effectiveness, and the relative contribution, of AMPA receptors even as they remained numerous (Kumar et al., 2002; Cull-Candy et al., 2006). Finally, a change in the distribution of pre- and post-synaptic machinery, such as increasing colocalization of NMDA channels and GABA vesicles, could enhance the effect of NMDA channels on reciprocal GABA release (Halabisky et al., 2000; although see Isaacson, 2001).

CONCLUSION

In summary, our experiments have uncovered a striking developmental profile of MC autoinhibition in the OB as measured with DDI. An initial increase in DDI is followed by a reduction by 1 month of age. We have investigated several possible mechanisms for this unexpected reduction, and conclude that a reduction in the number of available synapses; a decrease in GC excitability; and an increase in the relative contribution of NMDA receptors to GABA release from GC synapses could contribute to the reduction. An interesting question for future studies is how this biphasic development of inhibition affects or constrains odor processing in the OB.

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Diets et al.

Development of dendrodendritic inhibition
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