The Range of Intrinsic Frequencies Represented by Medial Entorhinal Cortex Stellate Cells Extends with Age

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In both humans and rodents, the external environment is encoded in the form of cognitive maps. Neurons in the medial entorhinal cortex (mEC) represent spatial locations in a sequence of grid-like patterns scaled along the dorsal–ventral axis. The grid spacing correlates with the intrinsic resonance frequencies of stellate cells in layer II of mEC. We investigated the development of frequency preferences in these cells from weaning to adulthood using patch-clamp and sharp microelectrode recordings. We found that the dorsal–ventral gradient of stellate cell properties and frequency preferences exists before animals are able to actively explore their environment. In the transition to adulthood, cells respond faster and become less excitable, and the range of intrinsic resonance frequencies in the population expands in the dorsal direction. This is likely to reflect both the growth of the brain and the expansion of the internal representation caused by new exploratory experience.

Introduction
The neuronal correlates of the development of spatial representation of the external environment are not yet fully understood. The medial entorhinal cortex (mEC) has become the main focus of research into this topic after the discovery in rodents that a large proportion of cells in this area represent external space as an internal sequence of multiple triangular “grids” (Fyhn et al., 2004; Hafting et al., 2005). The resolution of a grid in relation to external space can vary from a few meters to tens of centimeters (Brun et al., 2008). The majority of grid cells are found in layer II of mEC, and the size of the grid changes along the dorsoventral axis (Sargolini et al., 2006). Interestingly, the properties of the principal neurons in this area (often called stellate cells because of their distinctive shape) are defined by the composition of voltage-dependent channels embedded in their membrane and change as a function of their position along the dorsoventral axis (Giocomo et al., 2007). The frequencies of both resonance and membrane potential oscillations depend on the position of the neuron along this axis (Giocomo et al., 2007) as does the expression pattern of HCN channels (Garden et al., 2008; Giocomo and Hasselmo, 2008a). This suggests that the spacing of the grid may be determined by the intrinsic properties of these cells (Hasselmo and Brandon, 2008).

The development of stellate cell excitability in the first postnatal weeks has been recently addressed by Burton et al. (2008). Here, we extend their study by investigating the development of the gradient in stellate cell properties from weaning (postnatal days 21–25 (P21–P25) to adulthood when rats most actively explore their environment (Whishaw and Kolb, 2004; McCutcheon and Marinelli, 2009). Throughout the study, we used both patch clamp and sharp microelectrode recordings to distinguish developmental changes from variability in the measurements introduced by different recording techniques.

Materials and Methods
Male Wistar rats were decapitated under halothane (in accordance with United Kingdom Home Office procedures) or ether anesthesia (Berlin Animal Ethics Committee; T 0068/02), and horizontal slices (400 μm) containing the hippocampus and the perirhinal and entorhinal cortex were prepared as previously reported (Erchova et al., 2004). After gluing the cortical surface to the mount, slices were cut from ventral to dorsal towards the bregma. The slice position was measured in relation to the interaural line (Paxinos and Watson, 1998). Slices were incubated in the recording solution for at least 1 h before being transferred to a submerged (patch clamp) or an interface humidified (sharp microelectrode) recording chamber at 34 ± 1°C. Excitatory and inhibitory synaptic transmission was blocked. In a subset of experiments, the identity of recorded cells was confirmed histologically as previously reported from our laboratory (Gloveli et al., 1997; Erchova et al., 2004). The topographical location of recorded cells was taken as the position of the slice plus one-half of slice thickness. Recorded cells were sorted into two groups (ventral, 2.0–4.0 mm; dorsal, 4.1–6.2 mm).

The recordings were done using either patch clamp or sharp microelectrodes (see supplemental material, available at www.jneurosci.org) in current-clamp mode, bridge balance adjusted. Patch pipettes (5–6 MΩ) were filled with an intracellular solution (in mM: 140 K-glucuronate, 2...
MgCl₂, 10 phosphocreatine, 2 Na₂ATP, 0.4 NaGTP, 10 HEPES, and 0.4% bicytion, pH 7.3). Recordings were obtained using a Multiclamp 700B amplifier, filtered at 3 kHz, digitized at 10–100 kHz, and acquired using Clampex 9.0 (Molecular Devices).

Data analyses were performed using Clampfit 9.0 (Molecular Devices) and custom software written in Matlab (version 7.4; Mathworks). Input resistance, membrane time constant, rheobase, sag potential, and action potential (AP) parameters including analysis of afterpotentials (see supplemental material, available at www.jneurosci.org) were estimated from the response of the cell to a square current pulse of 500 ms. The time course for the sag potential (SAG-Tau) was analyzed from the voltage response to -400 pA using a double-exponential function (Giocomo et al., 2007) (see supplemental Fig. S1 A, available at www.jneurosci.org as supplemental material). The AP parameters were estimated from the first spike elicited by the rheobase stimulus (for details, see supplemental Fig. S1 B, available at www.jneurosci.org as supplemental material). The resonance frequency of a cell was determined from the average voltage response (five repetitions) to a sinusoidal current injection with linearly increasing frequency from 0 to 20 Hz over 30 s (Puil et al., 1986; Hutcheson and Yarom, 2000; Erchova et al., 2004). The measurements were obtained at three levels of membrane potential (hyperpolarized, approximately -75 mV; resting; depolarized, approximately -52 mV). The amplitude of oscillating command current was adjusted to avoid changes in the membrane potential by >10 mV. The input frequency corresponding to the highest impedance was defined as the resonance frequency. The amplitude (Q value) of the resonance was determined as the ratio of the peak impedance to the impedance at zero frequency. The dominant frequency of membrane potential oscillations (MPOs) (evoked by depolarization to near threshold by injecting a constant current, 10 pA smaller than the rheobase) was determined from the power spectra analysis (Erchova et al., 2004) as the frequency between 1 and 15 Hz with the highest power. MPOs were analyzed near spiking threshold on at least 2-s-long data subsets without spikes. All values are reported as mean ± SEM. For multiple comparisons, the multivariate ANOVA and regression lines and fits of distributions were constructed using the Matlab Statistical Toolbox. The test results are abbreviated as follows: *p < 0.05; **p < 0.01; ***p < 0.001 or n.s., not significant.

Results

We examined changes in excitability of mEC stellate cells from weaning to adulthood using both patch-clamp (juvenile: P20.8 ± 0.4, P17–P25, 48 cells; adult: P46.4 ± 0.7, P34–P56, 79 cells) and sharp microelectrode recordings (juvenile: P18.0 ± 0.5, P14–P22, 28 cells; adult: P93 ± 3, P73–P110, 58 cells). The patch-clamp results are presented in the main text, whereas most of the sharp microelectrode results and the overall summary table are in the supplemental material (available at www.jneurosci.org).

In juvenile rats, we found that cellular excitability changed as a function of cell position. The input resistance and time constant increased from dorsal to ventral, whereas the rheo-
base decreased (Fig. 1 B–D; supplemental Fig. S2 A, B, available at www.jneurosci.org as supplemental material). No difference in resting membrane potential or voltage threshold for AP initiation was observed. The ventral cells were more excitable and needed less current to elicit a spike, but at rheobase the dorsal cells produced a spike earlier (91 ± 24 vs 144 ± 21 ms) (Fig. 1 A). In adults, the rheobase increased (p < 0.001), reflecting a decrease in cellular excitability. We attributed these changes to the reduced input resistance (p < 0.001). It was, however, also accompanied by a decrease in the AP voltage threshold (from -50 ± 1 to -53 ± 1 mV). In addition, all cells responded faster to current injections as indicated by their decreased membrane time constants (p < 0.001). We evaluated the contribution of HCN channels by estimating the fast time constant for their activation from the sag potential (Giocomo et al., 2007) (supplemental Fig. S1 B, available at www.jneurosci.org as supplemental material). The SAG-Tau was reduced for dorsal cells (p < 0.001) and decreased in adults (p < 0.01) (Fig. 1 E; supplemental Fig. S2 C, available at www.jneurosci.org as supplemental material). In addition, we measured a number of AP parameters including afterhyperpolarization (AHP) (supplemental Fig. S1 B, available at www. jneurosci.org as supplemental material). In juvenile rats, none of the AHP components [fast AHP (fAHP), depolarizing afterpotential, medium AHP (mAHP) amplitude and duration] showed any significant correlation with cell position. However, the half-width of AP increased from dorsal to ventral cells (slope, -0.06; r = -0.53; p < 0.001). In adults, the fAHP decreased from 7–8 to 4–5 mV, the gradient in AP half-width duration became steeper (slope, -0.1; r = -0.56; p < 0.001), and the duration of mAHP also became correlated with the cell position (Fig. 1 F) (for details, see supplemental Methods and Fig. S2 D, available at www. jneurosci.org as supplemental material).

We characterized the frequency preferences of cells by measuring the membrane resonance and MPOs. The resonance frequency decreased along the dorsal–ventral axis in both juvenile and adult rats (p < 0.001), but the range of frequencies in adults was larger than in juvenile rats (Fig. 2; supplemental Fig. S3, available at www.jneurosci.org as supplemental material). Interestingly, during the transition to adulthood, the most distant ventral cells remained virtually unchanged, whereas the dorsal cells changed their frequency almost twofold (p < 0.001). We found that, in adults but not juvenile rats, the mAHP duration was correlated with the resonance frequency at both resting membrane potential and just below threshold (rrest = 0.56, p < 0.001; rf dep = 0.42, p < 0.01). In general, the resonance frequency decreased with depolarization (p < 0.001) (Fig. 2 C) and input resistance increased (p < 0.001). The MPO frequency also decreased from dorsal to ventral cells (p < 0.001) (Fig. 3) similar to the resonance frequencies but with slightly lower values. The MPOs obtained in patch-clamp recordings had, however, lower frequencies (Fig. 3 A, B) compared with sharp microelectrode recordings (Fig. 3 C, D) and did not show any age dependency.

Discussion

Our results suggest that the dorsal–ventral gradient of stellate cell properties develops before the age of active exploration and is likely to be genetically determined. During the transition to adulthood, cells become less excitable and the frequency preferences of individual cells change differentially in such a way that the range of frequencies represented by the population expands in the dorsal direction.

Although it is known that the age can be a critical source of variability in the data, the age-related differences can be confounded because of the use of different recording techniques in different age groups. To clarify this issue, we performed both patch-clamp and sharp microelectrode recordings. Although our main findings were consistent across both techniques, the membrane time constants of cells measured using sharp microelectrodes were smaller because of lower input resistance. Consequently, the range of resonance frequencies recorded by sharp microelectrodes was larger (the cutoff frequency of the cell low-pass filter depends on 1/2πτm). In both age groups, the resonance frequency decreased on depolarization similar as reported by Nolan et al. (2007), although the effect was weak when measured by sharp microelectrodes, similar to the study by Erchova et al. (2004). In all cases, the resonance frequency measured near threshold was larger than the frequency of MPOs, as we previously reported (Erchova et al., 2004; Engel et al., 2008). However,
the smaller range of measured frequencies in patch-clamp data, combined with prominent voltage-dependent decrease in resonance frequency, resulted in remarkable differences between MPOs data recorded by patch clamp and sharp microelectrode.

In our study, near-threshold MPOs showed no consistent significant voltage-dependent variations in frequency. However, the original study by Alonso and Llina’s (1989) reported an increase in MPO frequencies on depolarization. This was recently confirmed by Giocomo and Hasselmo (2008b), who also specified that this effect was mainly seen in dorsal, but not ventral cells. Previously, we observed a similar increase in a subset of cells when both subthreshold and perithreshold MPOs were considered (Erchova et al., 2004).

It is well accepted that the establishment of receptive fields and refinement of synaptic circuitry depends on sensory experience (Moody and Bosma, 2005). However, changes in synaptic strength might be accompanied by changes in intrinsic excitability as a means to maintain stability in neuronal circuits (for review, see Desai, 2003; Nelson and Turrigiano, 2008; Yu and Goda, 2009). To this end, the maturation of synapses would require faster cellular responses and an overall reduction in cell excitability, as we observed. The situation is more complex when intrinsic excitability is differentially regulated across the cellular population (for review, see Sjöström et al., 2008), adjusting its sensitivity to the temporal aspects of the input pattern. It is usually difficult to dissociate the effects of sensory experience and genetically prescribed patterns of postnatal maturation. Burton et al. (2008) found that dendritic spine density of mEC layer II stellate cells increased almost threefold from the second to the fourth postnatal week without changes of stellate cell excitability beyond P18. However, as our data show, the intrinsic excitability continues to change differentially within the population.

In stellate cells, frequency preferences are defined mainly by HCN and potassium channels (White et al., 1995; Fransén et al., 2004; Schreiber et al., 2004). It is therefore interesting that Burton et al. (2008) found a continuous increase in the expression of HCN channels up to the end of their study at P28, although they did not find a clear physiological correlate of this process. This pattern agrees with data by Richter et al. (1997) and with our own findings showing age-related acceleration of the sag potential. It
is not yet clear whether this is caused by a systematic variation in activation kinetics [as suggested by Giocomo and Hasselmo (2008a)] or by a pure difference in the number of HCN channels. In the rat hippocampus, for example, the HCN current both increases and accelerates because of the substitution of the slow HCN4 by the fast HCN1 isoform (Surges et al., 2006). It is interesting, then, that in HCN1-deficient mice the observed gradient in stellate cells properties flattens (Giocomo and Hasselmo, 2009), suggesting that regulation of this subunit might be essential.

At present, little is known about how potassium currents shape resonance in stellate cells; we found that both AP half-width and mAHP duration were positively correlated with resonance frequency in adults, although only AP width correlated with resonance frequency in juvenile cells. This suggests that some of the currents shaping mAHP and spike-activated currents might also contribute to the increase in resonance frequency.

Many cortical areas increase considerably in size during postnatal development, but it is not clear how this impacts the remodeling of neuronal circuits. One intriguing possibility suggested by our study is that the range of frequencies represented by stellate cells might extend without necessarily affecting the frequency preferences of already established cells. The anatomy of EC favors extension in the dorsal direction (Steffenach et al., 2005), because ventral mEC contains several lateral bands, with only one band...
associated with the representation of space, whereas the dorsal mEC is homogeneous in its spatial function (Dolorfo and Amaral, 1998; Kerr et al., 2007).

Although it is not yet precisely known how the population of grid cells encodes a location, the oscillatory interference model (Burgess et al., 2007) would predict that receptive fields of dorsal cells in young animals would have slightly larger grids compared with adults, given that synaptic connections are formed and receptive fields are established. However, in the modular code suggested by Fiete et al. (2008) in which an animal’s position is encoded by the population with a large number of unique spatial frequencies, the total number of represented locations grows exponentially with the number of spatial frequencies available. Accordingly, an extension of the range of frequencies might reflect both an expansion of internal space representation in accordance with new exploratory experiences and refinement of existing representations.

To conclude, our data suggest that intrinsic mechanisms allow for the rapid formation of a cognitive spatial map that can readily rescale during development, preserving already acquired representations, increasing the spatial resolution of the map, and allowing for the encoding of novel locations.

References


