Brian G. Burton, Michael N. Economo, G. Jenny Lee and John A. White *J Neurophysiol* 100:3144-3157, 2008. First published Oct 1, 2008; doi:10.1152/jn.90424.2008

You might find this additional information useful...

This article cites 74 articles, 27 of which you can access free at: http://jn.physiology.org/cgi/content/full/100/6/3144#BIBL

This article has been cited by 1 other HighWire hosted article:

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

A. Boehlen, U. Heinemann and I. Erchova *J. Neurosci.*, March 31, 2010; 30 (13): 4585-4589. [Abstract] [Full Text] [PDF]

Updated information and services including high-resolution figures, can be found at: http://jn.physiology.org/cgi/content/full/100/6/3144

Additional material and information about *Journal of Neurophysiology* can be found at: http://www.the-aps.org/publications/jn

This information is current as of April 29, 2010.

Development of Theta Rhythmicity in Entorhinal Stellate Cells of the Juvenile Rat

Brian G. Burton, Michael N. Economo, G. Jenny Lee, and John A. White

Department of Biomedical Engineering, Center for Memory and Brain, Center for BioDynamics, Boston University, Boston, Massachusetts

Submitted 31 March 2008; accepted in final form 29 September 2008

Burton BG, Economo MN, Lee GJ, White JA. Development of theta rhythmicity in entorhinal stellate cells of the juvenile rat. JNeurophysiol 100: 3144–3157, 2008. First published October 1, 2008; doi:10.1152/jn.90424.2008. Mature stellate cells of the rat medial entorhinal cortex (EC), layer II, exhibit subthreshold membrane potential oscillations (MPOs) at theta frequencies (4–12 Hz) in vitro. We find that MPOs appear between postnatal days 14 (P14) and 18 (P18) but show little further change by day 28+ (P28-P32). To identify the factors responsible, we examined the electrical responses of developing stellate cells, paying attention to two currents thought necessary for mature oscillation: the h current I_h , which provides the slow rectification required for resonance; and a persistent sodium current I_{NaP} , which provides amplification of resonance. Responses to injected current revealed that P14 cells were often nonresonant with a relatively high resistance. Densities of I_h and I_{NaP} both rose by about 50% from P14 to P18. However, $I_{\rm h}$ levels fell to intermediate values by P28+. Given the nonrobust trend in I_h expression and a previously demonstrated potency of even low levels of I_h to sustain oscillation, we propose that resonance and MPOs are limited at P14 more by low levels of I_{NaP} than of I_{h} . The relative importance of I_{NaP} for the development of MPOs is supported by simulations of a conductancebased model, which also suggest that general shunt conductance may influence the precise age when MPOs appear. In addition to our physiological study, we analyzed spine densities at P14, P18, and P28+ and found a vigorous synaptogenesis across the whole period. Our data predict that functions that rely on theta rhythmicity in the hippocampal network are limited until at least P18.

INTRODUCTION

One of the most prominent electrical signals in the hippocampus and adjacent entorhinal cortex (EC) is the theta field potential (Kramis et al. 1975; Mitchell and Ranck 1980; Vanderwolf 1969). This rhythm, typically covering the 4- to 12-Hz band in vivo, is most commonly observed during voluntary movements and under certain conditions of alert wakefulness. Theta unit or population activity, phase-locked to hippocampal theta, has also been reported in a number of other brain areas, including the medial septum, the amygdala, the prefrontal cortex, and the nucleus accumbens (Bland and Oddie 2001; Seidenbecher et al. 2003; Siapas et al. 2005).

The function of this rhythm is unclear, although the suggestions are numerous, including the coordination of information flow between brain areas (Dickson et al. 2000a), sensory inhibition (Sainsbury 1998), sensorimotor integration (Bland and Oddie 2001), memory (Seidenbecher et al. 2003; Winson 1978), phase-dependent coding of information (Huxter et al. 2003; Jensen and Lisman 2000; O'Keefe and Recce 1993), and

phase-dependent synaptic plasticity (Larson et al. 1986; Orr et al. 2001). Most recently, studies of the medial EC in the rat suggest a specific role in spatial coding. So-called grid cells in the superficial layers of medial EC activate as the animal enters points in space separated at regular intervals on an abstract lattice (Hafting et al. 2005). The periodic nature of this representation has led some to argue that the theta rhythm is critically involved (Blair et al. 2007; Burgess et al. 2007; Giocomo et al. 2007). Many of these functions are not mutually exclusive but, given their diversity, it is of interest to determine when and how the theta rhythm develops in the young animal. One might then be able to clarify the role of theta by correlating its emergence with particular cognitive or motor skills. Knowledge of the timing of theta development with respect to other developmental events may also aid our understanding of the mechanisms of theta generation in the mature animal.

A natural focus for a developmental study of theta generation is the stellate cell of layer II in the medial entorhinal cortex. Stellate cells are the principal projection neurons from the EC to the hippocampal formation (Steward and Scoville 1976; Witter 1993). They may also be grid cells because they are the most numerous principal neuron in layer II, which possesses the highest density of grid cells of all EC layers (Alonso and Klink 1993; Klink and Alonso 1997; Sargolini et al. 2006). When depolarized, stellate cells from medial EC produce clusters of spikes, loosely phase-locked to an intrinsic subthreshold membrane potential oscillation (MPO) in the theta range (Alonso and Klink 1993; Alonso and Llinás 1989). If combined with the appropriate synaptic interactions these cells could thus make an important contribution to theta rhythmogenesis and act as a direct source for hippocampal theta (Alonso and García-Austt 1987; Kocsis et al. 1999; Mitchell and Ranck 1980).

Theta-frequency MPOs in entorhinal stellate cells are believed to be generated by the interaction of an h current (I_h) and a persistent sodium current (I_{NaP}) (Alonso and Llinás 1989; Dickson et al. 2000b; Dorval and White 2005; Klink and Alonso 1993; White et al. 1998; see Nolan et al. 2007 for a divergent view and the DISCUSSION for more on this topic). According to current models, both I_h and I_{NaP} contribute to resonance by providing rectification and amplification, respectively (Dickson et al. 2000b; Hutcheon and Yarom 2000). Additionally, persistent sodium channels have a large conductance but occur in low numbers and contribute substantially to membrane noise (Magistretti and Alonso 1999; White et al. 1998). MPOs in current clamp are then the product of an I_h -

Address for reprint requests and other correspondence: J. White, Dept. of Bioengineering, Univ. of Utah, 20 S. 2030 E., Salt Lake City, UT 84112 (E-mail: john.white@utah.edu).

The costs of publication of this article were defrayed in part by the payment of page charges. The article must therefore be hereby marked "advertisement" in accordance with 18 U.S.C. Section 1734 solely to indicate this fact.

and $I_{\rm NaP}$ -induced resonance, ringing in response to $I_{\rm NaP}$ -induced noise (Dorval and White 2005; White et al. 1998). $I_{\rm h}$, supplemented by a calcium-dependent K⁺ current, is also thought to be involved in spike clustering (Fransén et al. 2004; Nolan et al. 2007). It is therefore reasonable to hypothesize that the expression profiles of $I_{\rm h}$ and $I_{\rm NaP}$ play a role in the development of theta rhythmicity in EC stellate cells. Here we examine the physiological development of stellate cells in rat medial entorhinal cortex in vitro and correlate this with expression of $I_{\rm h}$ and $I_{\rm NaP}$. We also examine the morphological development of these cells, particularly with regard to spine development as an indication of capacity to support theta at the network level.

METHODS

Preparation and recording

Long–Evans rats were anesthetized with 1.5 ml 2-chloro-2-(difluoromethoxy)-1,1,1-trifluoro-ethane (isoflurane; Abbott Laboratories, Chicago, IL), diffused in a 4-L container, and were decapitated. Their brains were removed and placed dorsal side down in a chamber filled with ice-cold, oxygenated artificial cerebrospinal fluid (ACSF; see following text for details). Horizontal slices, 300 μ m thick, of the hippocampal formation and entorhinal cortex were cut using a vibrating microtome (Vibratome 1000 Plus; Vibratome, St. Louis, MO). These were transferred to a second chamber of ACSF at room temperature and bubbled continuously with a 95% O₂-5% CO₂ mix for \geq 1 h. All procedures involving animals were approved by the Charles River Campus Institutional Animal Care and Use Committee, Boston University.

For recording, slices were transferred to a third chamber of oxygenated ACSF held at 34°C. The tissue was imaged under infrared differential interference contrast optics (Carl Zeiss, Thornwood, NY) and viewed with the aid of an infrared camera (CCD 100; Dage-MTI, Michigan City, IN) and a TV monitor. Recordings were from medial entorhinal cortex, layer II. Stellate cells were identified by location in superficial layer II, where other cell types are rare or nonexistent (Klink and Alonso 1997; Wang and Lambert 2003); the presence of a round soma and multiple thick primary dendrites (Klink and Alonso 1997); and delayed repolarizing responses ("sags") to negative current steps (Alonso and Klink 1993; Klink and Alonso 1997). During the process of isolating sodium currents, stellate cells were identified by the activation of an inward current in response to a depolarizing voltage ramp or step (Magistretti and Alonso 1999). In all cases in which cells were filled for anatomical reconstructions (see following text), cells selected using these criteria were confirmed to be stellate

Patch-clamp recordings were made with 4- to 6-M Ω borosilicate electrodes prepared with the Sutter P-97 puller (Sutter Instruments, Novato, CA). Series resistance and cell capacitance were measured in voltage clamp after canceling the whole cell capacitive transients evoked by 10-mV steps. These quantities were typically 10–30 M Ω and 80–120 pF, respectively. The effects of both on voltage-clamp fidelity were compensated by 60%. Capacitance neutralization was set to 6 pF. Acceptable cells were those with a resting potential less than -70 mV in current-clamp experiments, a resistance >300 M Ω in voltage-clamp experiments measuring persistent sodium currents, and a resistance >200 M Ω in h-current experiments. Prior to breaking into the cell, patch resistance was always >2 G Ω , indicating a tight seal between pipette and membrane.

Physiological responses were recorded on computer via the Multi-Clamp 700A amplifier (Axon Instruments, Union City, CA). Adjustments of the voltage- and current-clamp settings, including holding signals, were performed with the MultiClamp software provided with the amplifier. The feedback resistors on the headstage for voltage and

current clamps were 500 M Ω and 5 G Ω , respectively. Stimulus presentation and data acquisition were performed with custom software written in Matlab (The MathWorks, Natick, MA). Physiological signals were filtered at 800 Hz with a Bessel filter and sampled at 2 kHz. The general holding potential in voltage clamp was -40 mV for h-current recordings and -50 mV for persistent sodium recordings, after adjustment for junction potentials (see following text).

Recording solutions

The extracellular solution (ACSF) for slicing, current-clamp recording, and staining was (in mM): 126 NaCl, 1.25 NaH₂PO₄, 2 MgCl₂, 26 NaHCO₃, 2 CaCl₂, and 25 D-glucose. The intracellular (patch pipette) solution for current-clamp recording and filling was (in mM): 10 KCl, 120 K gluconate, 10 HEPES, 4 Mg-ATP, 0.3 Tris-GTP, and 10 Na₂ phosphocreatine plus 20 U·ml⁻¹ creatine kinase. During the process of filling cells, biocytin was added to the pipette at 0.5%.

For the experiments isolating the h current, the extracellular solution was (in mM): 80 NaCl, 26 NAHCO₃, 5 KCl, 3 MgCl₂, 2 CaCl₂, 2 BaCl₂, 2 CoCl₂, 0.4 CdCl₂, 40 tetraethylammonium (TEA) chloride, 4 4-aminopyridine (4-AP), 0.001 tetrodotoxin (TTX), and 10 Dglucose. The intracellular solution was the same as the slicing solution

For the experiments isolating the persistent sodium current, the extracellular solution was (in mM): 34 NaCl, 26 NAHCO₃, 5 KCl, 3 CsCl, 3 MgCl₂, 2 CaCl₂, 2 BaCl₂, 2 CoCl₂, 0.4 CdCl₂, 80 TEA, 4 4-AP, and 10 D-glucose. The intracellular solution was 10 KCl, 55 K gluconate, 50 Cs methanesulfonate, 10 HEPES, 10 EGTA, 4 Mg-ATP, 0.3 Tris-GTP, and 10 Na₂ phosphocreatine plus 20 U·ml⁻¹ creatine kinase.

Additionally, 10 μ M bicuculline methiodide and 4 mM kynurenic acid were added to all extracellular solutions to block synapses. The pH was adjusted to 7.4 with HCl and NaOH. Osmolality was adjusted to 300 mOsm with sucrose. Intracellular solutions were adjusted to a pH of 7.3 with HCl and KOH. Osmolality was 290 mOsm. Junction potentials were measured according to the method of Neher (1992). These were 12.1 mV (SD = 5.77 mV, n = 4) for the standard solutions, 14.5 mV (SD = 3.50 mV, n = 6) for the h-current solutions, and 14.7 mV (SD = 2.45 mV, n = 4) for the persistent sodium current solutions. Data were adjusted for these potentials.

Anatomy

Slices (300 μ m thick) were prepared as described earlier. Stellate cells were patched with electrodes containing 0.5% biocytin (Molecular Probes, Eugene, OR), filled for 1 h (Horikawa and Armstrong 1988), and fixed in 4% paraformaldehyde overnight. Following fixation, the slice was washed in 0.1 M phosphate-buffered saline (PBS) for 45 min and prepared for either brightfield or epifluorescence microscopy.

For bright-field specimens, cells were stained with diaminobenzidine (DAB). Endogenous peroxidase activity was quenched by exposure to 1% hydrogen peroxide ($\rm H_2O_2$) and 10% methanol in PBS for 30 min. Following a further 30 min in fresh PBS, the slice was reacted with avidin-biotinylated horseradish peroxidase (ABC Elite kit; Vector Labs, Peterborough, UK) in 0.75% Triton-X100 (Sigma–Aldrich, St. Louis, MO) PBS for 2 h. The slice was soaked in a solution of 5 μ l per ml of DAB in PBS for 30 min and the stain was developed for 2–3 min under a microscope with the addition of 1 μ l per ml $\rm H_2O_2$. The slice was washed in PBS for a further 45 min and mounted on a slide with Mowiol 4-88 (Polysciences, Warrington, PA) (Longin et al. 1993). Cells were imaged using a MagnaFIRE digital camera (Optronics, Goleta, CA) attached to an Olympus BX51 microscope (Olympus, Melville, NY).

For the fluorescence microscopy, washed slices were incubated for 4 h in a mixture of 0.75% Triton-X100 and 2 μ g·ml⁻¹ streptavidin-Alexa Fluor 488 (Molecular Probes). Slices were mounted in Mowiol.

Images of filled cells were obtained using a confocal laser scanning microscope (Zeiss LSM 510, Carl Zeiss) with a $\times 40$ objective lens. Laser excitation was at 458 nm (argon) and emitted light was low-pass filtered at 475 nm. Image contrast was adjusted after scanning to suppress background staining.

Analysis

VOLTAGE POWER AND IMPEDANCE SPECTRA. Power spectra of perithreshold voltages were calculated from traces of cells firing between 0 and 0.1 Hz in current clamp. At these frequencies, long periods of robust subthreshold theta oscillation may be observed between spikes. Traces were divided into overlapping segments of 2-s duration, each shifted by 0.2 s from the previous trace. Ignoring segments within 150 and 300 ms before and after any spikes, a cell's power spectrum was calculated as the mean power spectral density of all its segments (truncated cosine window function; Papoulis 1991). Unless otherwise stated, overall power was determined as the integral of the power spectrum between -25 and 25 Hz, excluding 0 Hz.

Impedance spectra were calculated from the voltage responses of stellate cells to white noise current stimuli at 5 mV above rest. Stimuli were 12.5 s long, but only the last 10 s were analyzed to avoid contamination by the cell's natural (transient) response. Each cell received a different stimulus, white to a frequency of 25 Hz. Stimulus SD ranged from 7 to 40 pA and was adjusted to keep the response within a peak-to-peak range of about 5 mV. As for voltage power, the stimulus and the mean response of a cell to eight presentations of the stimulus were divided into overlapping 2-s segments. A cell's impedance spectrum was calculated as the Fourier transform of the response divided by the Fourier transform of the stimulus, averaged across segments.

ACTIVATION CURVES. The steady-state h conductance $g_{\rm h}(V)$, as a function of membrane potential, was determined from the tail currents $I_{\rm tail}(V,\,t)$, recorded following voltage steps to potentials V, from a holding potential of $-40~{\rm mV}$, as illustrated in Fig. 5. For each step, we fitted the model

$$I_{\text{tail}}(V, t) = A_1(V) \exp(-t/\tau_1) + A_2(V) \exp(-t/\tau_2)$$
 (1)

with adjustable parameters A and τ describing the amplitudes and time constants of the two components of the tail current. $g_h(V)$ is then given by

$$g_{\rm h}(V) = \frac{A_1(V) + A_2(V)}{V_{\rm holding} - E_{\rm h}}$$
 (2)

where $V_{\rm holding}$ is the holding potential. $E_{\rm h}$ is the effective reversal potential of the h current, calculated using the Goldman–Hodgkin–Katz equation to be -28 mV, assuming a permeability ratio of Na⁺ to K⁺ ions of 0.4 (Dickson et al. 2000b). Assuming $\tau_1 < \tau_2$, the quotient β was formed as $\beta(V) = A_1(V)/[A_1(V) + A_2(V)]$.

The steady-state persistent sodium conductance $g_{\text{NaP}}(V)$ was estimated from current responses to either voltage ramps or steps as illustrated in Fig. 6. If I(V) is the residual current recorded in the ramp or step protocols after subtraction of TTX-insensitive currents, $g_{\text{NaP}}(V)$ is given by the relation

$$g_{\text{NaP}} = \frac{I(V)}{V - E_{\text{Na}}} \tag{3}$$

where $E_{\rm Na}$ is the sodium Nernst potential, calculated from intracellular and extracellular concentrations in the persistent sodium recordings to be $+29~{\rm mV}$.

The estimates of the conductance (activation) curves, $g_{\rm h}(V)$ and $g_{\rm NaP}(V)$, for the two channels were modeled as Boltzmann functions of the form

$$g(V) = \frac{G}{1 + \exp[-(V - V_{50})/\lambda]} \tag{4}$$

with three parameters—G, V_{50} , and λ —describing, respectively, the maximum conductance of the channel, the potential of half-maximum activation, and the channel's activation range.

SPINE DENSITY. Spine densities were calculated from DAB-stained slides, using a grid superimposed over the dendritic field by camera lucida. Grid squares of 40×40 - μ m dimensions were selected at random and the spine density in each was calculated manually as the number of spines present divided by the total dendritic length. Length was measured with a caliper resolution of 8 μ m. Squares without dendrites were ignored. When counting spines, long, thin filopodia, thought to represent spine precursors (Fiala et al. 1998), were not included.

STATISTICS. For each measured parameter, the overall effect of age across the study period was assessed by ANOVA. Additionally, a linear trend analysis was applied to the data from P14 to P18 (Maxwell and Delaney 2004). Values at P28+ were compared with those at both P18 and P14 by *t*-test. In those cases for which data were available only at P14, P18, and P28+, the linear trend analysis was replaced by a *t*-test. To control for the family-wise error introduced by performing three comparisons, *P* values were interpreted with reference to an alpha level of 0.05/3 = 0.0167 (Howell 1997). All *F* statistics presented in the text refer to the results of an ANOVA. Student's *t*-statistics refer to either a linear trend analysis or a *t*-test or a test of a Pearson's correlation coefficient, as appropriate. All quoted measures of spread are SEs.

RESULTS

Physiological development of stellate cells

In current clamp, stellate cells of layer II entorhinal cortex (EC) may be distinguished from nearby nonstellate cells by the production of a delayed repolarization, or "sag," in response to hyperpolarizing current steps and by early spikes in response to depolarizing steps (Fig. 1, *A* and *B*) (Alonso and Klink 1993). Additionally, mature cells exhibit a subthreshold membrane potential oscillation (MPO) in response to constant depolarizing current (Fig. 1, *B* and *C*) (Alonso and Klink 1993; Alonso and Llinás 1989). Spikes tend to occur in clusters, loosely phase-locked with the oscillation and limited to one spike per cycle.

Perhaps the most striking of these features is the MPO. This is generated intrinsically and is thought to involve the interplay of two currents, a slow rectifying h-current (I_h) and a regenerative "persistent" sodium current (I_{NaP}) (Alonso and Llinás 1989; Dickson et al. 2000b; Dorval and White 2005). Although I_h is thought to be a crucial contributor to the resonance necessary to support an oscillation, I_{NaP} amplifies the resonant response and provides the channel noise sufficient to trigger and maintain "ringing" oscillations of the resonant system. In our recordings, we typically found MPOs to be in the low theta to subtheta frequency range (\sim 3 Hz), although spike frequencies within a cluster were slightly higher. This frequency is slightly lower than the in vivo theta rhythm, but is consistent with other patch-clamp recordings from ventral EC (Giocomo et al. 2007). We investigated the development of the oscillation in cells from animals aged between postnatal days 14 (P14) and 18 (P18). The properties of a second group, aged 28 to 32 days ("P28+"), represented the mature condition.

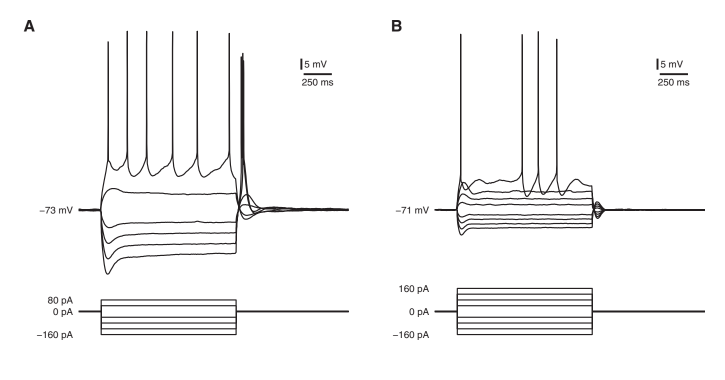




FIG. 1. Physiological characteristics of stellate cells. *A* and *B*: current-step responses of immature (postnatal day 14 [P14]; *A*) and mature (P28; *B*) stellate cells from medial entorhinal cortex, layer II. Steps start at -160 pA and increment by 40 pA until spike threshold is exceeded. Stellate cells were identified on the basis of size, location, and somatodendritic shape (Klink and Alonso 1997). However, even by P14, they were distinguishable within layer II by a repolarizing sag in response to negative current and early spikes in response to positive current. *C*: responses of a mature stellate cell to constant depolarizing current (200 pA). Same cell as in *B*. Spikes are clipped for presentation purposes. Mature stellate cells exhibit subthreshold oscillation and clustered spiking.

As illustrated in Fig. 2, MPOs were absent at P14 but gradually emerged between P14 and P18. Visual comparison of the P18 and P28+ examples will find that the oscillations at P18 are of a frequency and consistency similar to those in the adult. This observation was repeated throughout the set, although P28+ oscillations were generally of slightly lower amplitude than that at P18. To quantify these changes, we constructed power spectra of the membrane voltage at perithreshold levels of injected DC current, where the cell is spiking but at such a low rate that the "subthreshold" MPOs can be well isolated (METHODS). At P14, when MPOs were absent, voltage power fell monotonically with frequency over the theta range (Fig. 3A). On P16 (not shown), a shoulder appeared around 3 Hz and this developed into a prominent peak by P18. At older ages, overall power fell, starting at 0.288 mV^2 on P18 and ending at 0.126 mV^2 by P28+. Nevertheless, the proportion of power in the theta band was maintained at P28+, demonstrating continued theta content. Quantifying the prominence of theta as the proportion of the total power occurring between 2 and 4 Hz, there was a significant effect of age on theta prominence [ANOVA: F(5,26) = 9.25, P <0.001]. Mean theta prominence increased steadily from

 0.240 ± 0.0269 (n = 4) at P14 to 0.544 ± 0.0392 (n = 6) at P18 [trend analysis: t(26) = 5.55, P < 0.001] (Fig. 3D). At P28+, theta prominence was 0.574 ± 0.0435 (n = 4). Theta prominence at P28+ was significantly larger than the value at P14 [Student's t-test: t(26) = 5.28, P < 0.001], but not significantly different from that at P18 [Student's t-test: t(26) = 0.525, P = 0.604]. This pattern was not attributable to systematic shifts in recording potential because spike threshold (where MPOs were recorded) showed no effect of age [F(5,26) = 1.28, P = 0.303].

Although our main concern here is the development of MPOs, other changes over the study period are also evident in Fig. 2. At P14, the rebound following spike afterhyperpolarization (AHP) was slow, such that voltage rose steadily from one spike to the next and interspike interval (ISI) was fairly constant. Starting at P15, however, the AHP began to shorten in some cells and on P16 and P17 spikes began to cluster as in the mature cell. Because both AHP duration and spike clustering are thought to be influenced by $I_{\rm h}$ (Fransén et al. 2004; Klink and Alonso 1993; Nolan et al. 2007), we examined these attributes in more detail (Fig. 3, B, C, and E–G). As shown in Fig. 3B, shortening of the AHP was a general feature of our

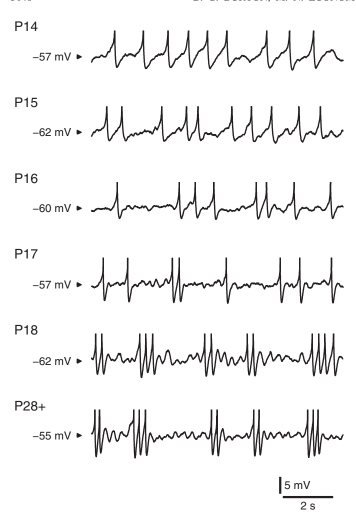


FIG. 2. The development of oscillatory behavior in stellate cells. Example traces recorded from stellate cells in response to constant current at 6 different days of development, as indicated. Traces are roughly matched for firing rate. At P14 and P15, action potentials occurred somewhat regularly and had a long afterhyperpolarization (AHP). At P16 and P17 action potentials began to cluster, the AHP shortened, and oscillations appeared between spikes. By P18 an adult physiology was obtained, resembling that at P28+ (P28–P32). Spikes were highly clustered and oscillations were strong.

data. Half-width, measured as the width of the AHP at half the minimum voltage relative to background, varied significantly over the study period as a whole [F(5,39) = 15.9, P < 0.001]and fell significantly from 174 \pm 14.7 ms (n = 4) at P14 to $89.4 \pm 3.50 \text{ ms } (n = 7) \text{ at P18 } [t(39) = 7.60, P < 0.001]. \text{ Like}$ theta prominence, however, it witnessed no further change from P18 to P28+ [t(39) = 1.06, P = 0.295] (Fig. 3E). Further features of AHP development evident in Fig. 3B are the deepening of the AHP and the development of an overshoot between days P14 and P18. Measuring overshoot as the maximum potential reached above background voltage, overshoot rose significantly from near 0 mV at P14 to 1.01 \pm 0.150 mV at P18 [t(39) = 5.48, P < 0.001] (Fig. 3F). Although overshoot fell again by P28+ (0.589 \pm 0.172 mV, n = 8), the drop did not quite reach significance after taking account of family-wise error [t(39) = 2.38, P = 0.023; see METHODS]. Furthermore, overshoot at P28+ was still significantly greater at this age than that at P14 [t(39) = 2.77, P = 0.008]. Amalgamating all ages, both AHP duration and overshoot were highly correlated

with theta prominence [Pearson's r > 0.719, t(28) > 5.47, P < 0.001].

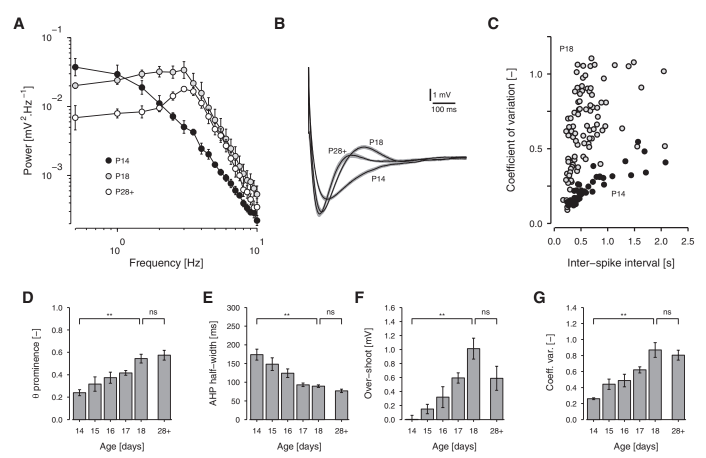
To quantify the development of clustering, we calculated the coefficient of variation (CV) in ISI from recordings of stellate cells firing over their full range of firing rates. As shown in Fig. 3C, the CV was consistently higher in P18 animals than that in P14 animals. This was true at all but the lowest ISIs, where all cells were firing regularly. Between 0.5 and 1 Hz, where clustering was most prominent in mature cells, the CV witnessed a significant change over the study period [F(5,37)]10.0, P < 0.001], increasing significantly from 0.261 \pm 0.0114 (n = 4) at P14 to 0.869 \pm 0.0926 (n = 7) at P18 [t(37) = 6.02,P < 0.001] (Fig. 3G). At P28+ CV was 0.805 ± 0.0621 (n =8) and, again, whereas the overall change from P14 to P28+ was significant [t(37) = 5.12, P < 0.001], the change from P18 to P28+ was not [t(37) = 0.718, P = 0.477]. The CV was highly correlated with both theta prominence [r = 0.790]t(27) = 6.70, P < 0.001 and AHP overshoot [r = 0.831, t(39) = 9.33, P < 0.001].

Developmental changes in electrical response

Given that MPOs in stellate cells are thought to be the product of a highly resonant membrane interacting with internal noise sources (Dorval and White 2005; White et al. 1998) one might expect the development of MPOs to be associated with increasing resonance.

To measure resonance, we injected white noise currents into a subset of cells at the key ages-P14, P18, and P28+ (Fig. 4A)—and constructed impedance spectra from the responses (METHODS). Membrane potential was depolarized to -67 mV ($\sim 5 \text{ mV}$ above rest), to bring the potential near the value normally associated with spontaneous MPOs but below the threshold to trigger action potentials. Spectra were fitted with the empirical model of Erchova et al. (2004), which treats the membrane as a low-pass capacitative circuit, representing the passive membrane, in parallel with a high-pass inductive circuit, representing a rectifying conductance. From the fits, descriptive parameters were calculated, including the frequency of maximum impedance (f_0) and the Q factor, which is the ratio of the maximum impedance amplitude to its value at the lowest frequency measured (0.5 Hz). We also extracted the cutoff frequency (f_{cut}), which is the frequency at which the passive membrane attenuates impedance power by half and is a conventional measure of membrane speed or bandwidth. (In the time domain, one would use the time constant: $\tau_{\rm m}$ = $1/2\pi f_{\rm cut}$.)

As illustrated in Fig. 4*B*, the magnitudes of impedance power spectra fell substantially from P14 to P18 and beyond. There was also a clear change in the mean shapes of spectra, from being largely low-pass at P14 to becoming resonant, with a peak at nonzero frequencies at P18 and P28+. In fact, three of eight P14 cells were completely nonresonant, having a maximum impedance power at 0.5 Hz. Overall f_0 changed significantly from 1.44 \pm 0.395 Hz (n=8) at P14 to 3.36 \pm 0.340 Hz (n=7) at P18 and 3.35 \pm 0.308 Hz (n=10) at P28+ [F(2,22)=9.93, P<0.001]. The change from P14 to P28+ was significant [t(22)=3.70, P=0.001] but the change from P18 to P28+ was not [t(19)=0.015, P=0.99] (Fig. 4*C*). Parallel changes occurred in the cutoff frequency, which rose significantly from 2.65 \pm 0.304 Hz at P14 to 4.76 0.363 Hz at



P18 and 4.13 \pm 0.458 Hz at P28+ [F(2,22) = 6.72, P = 0.005] (Fig. 4*D*). Again P14 and P28+ were significantly different [t(22) = 2.69, P = 0.013] but P18 and P28+ were not [t(22) = 1.10, P = 0.285]. The Q factor, which measures how strongly the membrane band-pass filters broadband input, followed a similar pattern. This rose from 1.16 \pm 0.062 (n = 8) at P14 to 1.35 \pm 0.072 (n = 7) at P18 and 1.34 \pm 0.049 (n = 10) at P28+, but the overall effect of age fell just short of significance [F(2,22) = 3.23, P = 0.059] (Fig. 4*E*).

These results indicate two features of stellate cell development. First, the amplitude of the resonance peak increases from P14 to P18, consistent with the hypothesis that the development of MPOs follows the development of resonance. Second, this increase in resonance is achieved despite a decrease in overall cellular input resistance. The falling magnitude of the impedance at low frequencies and the increasing bandwidth indicate that resistance is lower at P28+ than that at P14. Yet, artificially decreasing resistance

in stellate cells (P19–P31) has been shown to lower resonance and both MPO amplitude and theta content (Fernandez and White 2008). Thus there must be a strong current contributing to resonance for impedance to become more resonant from P14 to P18.

Based on past work on the subject of resonance and MPOs in mature stellate cells (e.g., Dickson et al. 2000a,b; Dorval and White 2005; Haas and White 2002; Haas et al. 2007; Klink and Alonso 1993; White et al. 1998), there are two obvious candidate membrane currents: I_h , the slow rectifying current thought to set the resonant period, and I_{NaP} , the persistent Na⁺ current that amplifies and adds noise important for MPOs. We first studied I_h in current-clamp data by examining whether the degree of sag changes from P14 to P28+. As shown in Fig. 1A, in response to negative current steps, P14 cells already showed a prominent sag, considered the signature of I_h (Klink and Alonso 1993). For step amplitudes of between -10 and -40 pA, producing re-

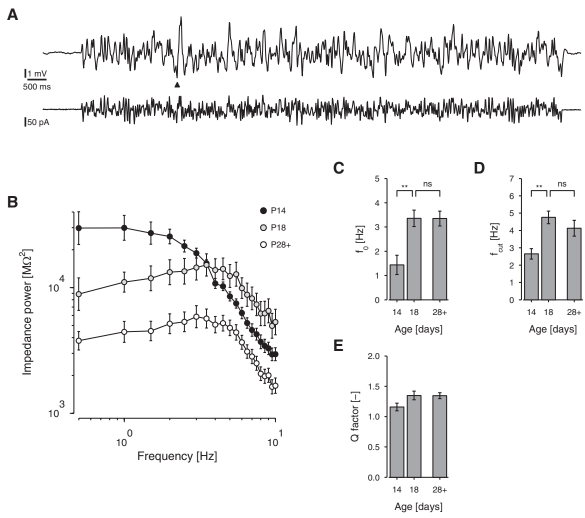


FIG. 4. Response to injected current. A: mean response (above) of an example cell from a P29 rat in response to the presentation of a white noise current stimulus (below) at -67 mV. Mean response is an average of 8. To avoid the influence of the cell's transient response on impedance measurements, only the last 10 s, starting from the arrowhead, were used to estimate membrane impedance (METHODS). B: mean impedance power spectra (square of impedance amplitude) obtained from white noise responses for cells from rats at P14, P18, and P28+, as indicated. P14 cells were mainly low-pass. C-E: impedance parameters (P14, P18, and P28+, n = 8, 7, 10). C: frequency of maximum impedance, f0. D: cutoff frequency, fcut. E: Q-factor (maximum amplitude divided by amplitude at 0.5 Hz).

sponses ending between -80 and -75 mV, steady-state resistance fell continuously from P14 to P28+ [F(5,73) = 13.6, P < 0.001] but the percentage reduction in the voltage response from peak to steady state was quite stable, with a mean of 26% [P14 to P28+: F(5,73) = 2.10, P = 0.075; P14 vs. P28+: t(73) = 0.278, P = 0.782]. That is, developmental changes in resonance near threshold were not matched by effective changes in I_h -generated sag below rest.

Expression of ionic currents

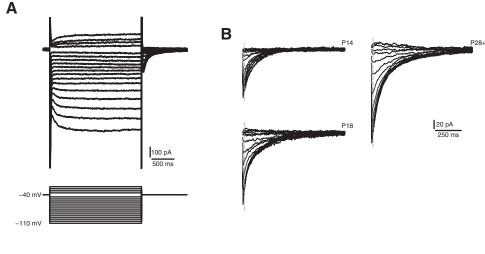
Although the preceding results may suggest that $I_{\rm h}$ is not a limiting factor in the emergence of resonance and MPOs from P14 to P18, the responses of stellate cells to injected current under normal conditions are influenced by several ionic currents, which may obscure the particular contribution of $I_{\rm h}$. To further investigate the mechanisms of MPO development, we studied $I_{\rm h}$ and $I_{\rm NaP}$ in isolation under voltage clamp (see following text). The likely effects of these currents and their relative importance in limiting MPO

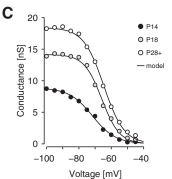
development are explored through simulation in the supplemental material.¹

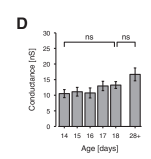
Expression of I_h

Expression of I_h was investigated using the step protocol of Dickson et al. (2000b), illustrated in Fig. 5, A and B. In the presence of blockers of K^+ , Ca^{2^+} , and Na^+ channels, voltage steps of 2-s duration were applied from a holding of -40 mV (Fig. 5A). The tail currents (Fig. 5B) predominantly reflect the slow decay of I_h from its steady-state step value to its value at the holding potential (Dickson et al. 2000b). The amplitude of each tail current was measured by fitting a double-exponential curve to the tail (Eq. I) and finding the value of the fit shortly after the tail's peak. The corresponding conductance (g_h) was calculated by subtracting the current amplitude obtained at the most depolarized step, where I_h is inactive, and adjusting the result for the effective reversal potential of the I_h channel (Eq.

¹ The online version of this article contains supplemental data.







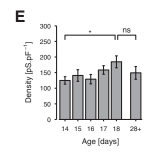


FIG. 5. Development of the expression of I_h . A and B: voltage-step protocol for isolating the h current, I_h . A: current responses of an example stellate cell (P32) to 2-s voltage steps, applied from a holding of -40 mV, in the presence of Na+, K+, and Ca2+ channel blockers. B: the magnified tail currents of 3 representative cells at P14, P18, and P28+ (same cell as in A). The vertical dashed line indicates the point at which amplitudes are measured. Tail current amplitude reflects the decay of the h conductance from its steadystate step value to its value at the holding potential. C: activation curves of the h-conductance, g_h , for the cells in B. The tail current magnitude for the highest voltage step is subtracted from all others and the results are adjusted for electromotive force (emf) to indicate conductance as a function of voltage (discs). The data are then fitted with a Boltzmann curve (model). D and E: histograms of conductance amplitude (D) and density (E) of I_h from P14 to P18 and at P28+ (n = 15, 18, 8, 15, 18, 5). D: the absolute I_h conductance increased throughout the period studied. E: I_h conductance density increased somewhat from P14 to P18 but was intermediate at P28+.

2). Conductance as a function of voltage was then fitted with a Boltzmann model whose parameters $G_{\rm h}$, $V_{50\,\rm h}$, and $\lambda_{\rm h}$ describe, respectively, the maximum conductance of $I_{\rm h}$ (i.e., the conductance when $I_{\rm h}$ is fully activated), the half-activation potential of $I_{\rm h}$, and the range of activation (Fig. 5C) (Eq. 4).

Figure 5, D and E shows respectively the maximum conductance magnitude (G_h) and the corresponding conductance density (or conductance per unit capacitance) of I_h across the period of our study, where capacitance was read from the MultiClamp amplifier (METHODS). Although the former indicates the absolute expression levels of I_h , the latter is more interesting from a functional point of view because it indicates the effectiveness of I_h to drive the membrane. Both measures showed a general increase with age (see also Fig. 5, B and C). However, the overall effect for conductance density was weak [F(5,73) = 1.74, P = 0.136] because capacitance rose from P18 to P28+ (not shown). Between P14 and P18, conductance density increased from $125 \pm 12.0 \text{ pS} \cdot \text{pF}^{-1}$ (n = 15) at P14 to $184 \pm 19.2 \text{ pS} \cdot \text{pF}^{-1}$ (n = 18) at P18, about 47%. The trend was significant [t(73) = 2.73, P = 0.008] but density at P28+ was intermediate at $149 \pm 20.3 \text{ pS} \cdot \text{pF}^{-1}$ (n = 5) and was not significantly different from either the P14 value [t(73) = 0.717,P = 0.476] or the P18 value [t(73) = 1.08, P = 0.286]. It is therefore not easy to correspond I_h expression with MPO theta

In addition to conductance, we analyzed the other parameters of the Boltzmann model of the conductance curve as well as the parameters of the double-exponential model of tail current, including the time constants of the two components τ_1 and τ_2 , and the proportion (β) of current amplitude contributed

by the faster (τ_1) component. None of these showed a statistical trend across the period of the study. The grand means for these parameters were: half-activation potential, -66.9 ± 0.456 mV; activation range, 6.25 ± 0.119 nS; fast time constant, 56.6 ± 3.57 ms; slow time constant, 284 ± 13.44 ms; and current quotient (β), 0.515 ± 0.0200 (n = 79). β , τ_1 , and τ_2 were averaged across responses to step voltages more negative than -70 mV, where tail currents are robust and we found β to be constant.

In summary, although there is some evidence for developing expression of the h current in the third week after birth, the overall pattern is not consistent with $I_{\rm h}$ being an important limiting factor in the appearance of MPOs. These results support our earlier finding that the strength of the voltage sag in response to negative current steps was stable from P14 onward and indicate that the poor resonance of P14 cells may be due to factors other than levels of $I_{\rm h}$ expression. Consistent with this interpretation, our simulations showed that relative changes in the levels of $I_{\rm h}$ of the order measured here have relatively little effect on the existence of MPOs compared with other factors (such as $I_{\rm NaP}$). Its role appears to be to help set the overall magnitude of the oscillation and, to a modest extent, the oscillation frequency (supplemental material).

Expression of I_{NaP}

Expression of $I_{\rm NaP}$ was investigated using either the ramp or step protocols of Magistretti and Alonso (1999). Both protocols rely on the sensitivity of $I_{\rm NaP}$ to tetrodotoxin (TTX) and its slow rate of inactivation compared with that

of the conventional transient sodium current $(I_{\rm NaT})$. In fact, the $I_{\rm NaP}$ of stellate cells inactivates in a biexponential manner and, following measurements by Magistretti and Alonso (1999), we chose stimuli that isolate the slower, more "persistent" component.

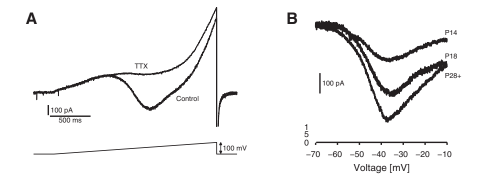
The ramp protocol is illustrated in Fig. 6A. In the presence of blockers of K^+ , Ca^{2+} , and h channels, voltage ramps were applied, rising at a speed of 50 mV·s⁻¹ from a holding potential of -95 mV to a maximum of +5 mV. At this ramp speed, $I_{\rm NaT}$ and the faster inactivating component of $I_{\rm NaP}$ are exhausted but the slower inactivating $I_{\rm NaP}$ components are relatively unaffected (Magistretti and Alonso 1999). The step protocol was similar except the stimuli were steps of various amplitudes, each applied from a holding of -80 mV for 800 ms. In both cases, the procedure was repeated in TTX and the difference between the two sets of responses was taken to reflect the amplitude of $I_{\rm NaP}$ (Fig. 6B).

The difference curve obtained was adjusted for the sodium Nernst potential $(E_{\rm Na})$, according to Eq.~3 to obtain an estimate of the persistent sodium conductance $g_{\rm NaP}(V)$, as a function of voltage V. Again, this curve was fitted with a Boltzmann model (Eq.~4) as shown in Fig. 6C, and parameters were extracted, including maximum conductance magnitude, $G_{\rm NaP}$; the half-activation potential, $V_{\rm 50~NaP}$; and the range of activation, $\lambda_{\rm NaP}$.

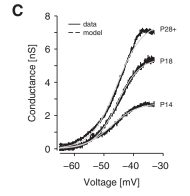
We compared the responses of cells from animals at P14 to P18 and P28+. Neither V_{50} nor λ showed any statistical effect of age, either between P14 and P18 [V_{50} : t(79) = 0.941, P = 0.350; λ : t(79) = 0.237, P = 0.813] or between P18 and P28+ [V_{50} : t(79) = 1.01, P = 0.317; λ : t(79) = 0.269, P = 0.789]. The grand means for these parameters across all animals were -45.9 ± 0.304 and 3.20 ± 0.103 mV, respectively (n = 85). We therefore focus on the maximum conductance amplitude

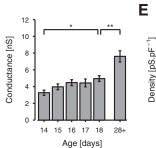
 (G_{NaP}) and corresponding conductance density. Like G_{h} , G_{NaP} increased from P14 to P18 but additionally rose substantially from P18 to P28+ (Fig. 6, B–D). Consequently, conductance density was maintained at P28+, despite the large membrane area of older cells (Fig. 6E), and the overall effect of age was significant [F(5,79)=3.75, P=0.005]. From P14 to P18, conductance density increased by 50% from 36.6 \pm 3.76 to 54.9 \pm 3.20 pS·pF⁻¹. The effect of age across this period was significant [t(79)=3.24, P=0.002] as was the difference between densities at P14 and P28+ [t(79)=2.90, P=0.005]. However, there was not a significant difference between densities at P18 and P28+ [t(79)=0.613, P=0.542]. This pattern mirrors trends in theta prominence, which reached mature levels by about P18 (Fig. 3).

Thus like I_h , the persistent sodium current I_{NaP} was present at P14 and more prominent at P18. Unlike I_h , however, the expression of I_{NaP} was well correlated with physiological development whereby densities increased from P14 to P18 and remained high at P28+. These results are consistent with a hypothesis that P14 cells fail to show MPOs because they lack sufficient excitability normally provided in the adult by I_{NaP} . Supplemental simulations indicate that changes in I_{NaP} density of the order measured here have dramatic effects on the occurrence of MPOs, mainly through their role in amplification but also due to the changes in noise levels they produce (supplemental material). In simulations, the amplification produced by a 50% increase in $I_{\rm NaP}$ conductance density (mimicking the change from P14 to P18) was so large that the only way to account for the observed trends in MPO strength in simulations was to include similarly large increases in shunt conductance, consistent with decreases in input resistance and increases in membrane band width we measure from P14 to P18.



D





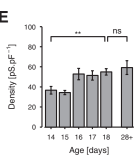


FIG. 6. Development of the expression of I_{NaP} . A: voltage-ramp protocol for isolating the persistent sodium current, I_{NaP} . Current responses of an example stellate cell (P18) to a 50 mV·s⁻¹ voltage ramp, applied from a holding of -95 mV, before (control) and after (tetrodotoxin [TTX]) the addition of 1 μM TTX to the bath. B: isolated persistent sodium currents as a function of voltage for representative cells at P14, P18 (same cell as in A), and P28+. Currents are calculated by subtracting TTX ramp responses from control ramp responses and mapping time onto voltage according to ramp speed. C: activation curve of the persistent sodium conductance, g_{NaP} , for the cells in B. The currentvoltage functions in B are adjusted for emf to indicate g_{NaP} as a function of voltage (black) and the results fitted with a Boltzmann curve (gray). D and E: histograms of conductance amplitude (D) and density (E) of I_{NaP} at P14 to P18 and P28+ (n = 11, 12, 21, 11,18, 12). D: the absolute I_{NaP} conductance increased steadily from P14 to P18 and beyond. $E: I_{NaP}$ conductance density also increased steadily from P14 to P18, but there was little further change in the mature animal.

J Neurophysiol • VOL 100 • DECEMBER 2008 • www.jn.org

Anatomical development of stellate cells

Theta field potentials observed in the EC in vivo (Alonso and García-Austt 1987; Mitchell and Ranck 1980) suggest that the intrinsic rhythmicity of stellate cells may be used either to receive and resonate with theta input from elsewhere or to synchronize cells within layer II at theta frequencies through local interactions. This model supposes that the stellate cell has the appropriate complement of synaptic sites to receive driving input. Both in hippocampal neurons (Harris et al. 1992; Kirov et al. 2004; Pokorný and Yamamoto 1981) and stellate cells in other parts of the rat brain (Ferrer and Martinez-Matos 1981) the period of our study is associated with synaptogenesis. It is therefore of interest to establish the morphological development of EC stellate cells, particularly with regard to synaptic growth.

We examined the anatomy of stellate cells at P14, P18, and P28+ stained with either DAB or streptavidin-Alexa Fluor 488. P14 stellate cells already possessed many of the features of the mature cell (Fig. 7A) (Klink and Alonso 1997; Lingenhöhl and Finch 1991). There were prominent primary dendrites, projecting superficially and basally in a wide fan and extending \leq 700 μ m parallel to the pial surface, tip to tip. The axon usually projected from a basal dendrite. In three of eight P14 cells the axon was preserved to at least layer VI and axon collaterals could be seen in one or two of layers III-VI. Collaterals were always beaded and could extend medially or laterally. In addition, six P14 cells possessed a fine net of beaded axon collaterals in layer II (Fig. 7A, arrowheads). This net mainly covered a spherical region of radius 50 to 100 μ m, centered on the soma, although one to three processes could often be seen extending $\leq 100 \, \mu \text{m}$ beyond the dendritic field in either the medial or lateral direction, often both.

P14 cells were, however, less complicated than older cells, having fewer dendrites and many fewer spines (Fig. 7*B*). We quantified the change in spine density in cells stained with

DAB. Mean spine density, measured across randomly selected 40×40 - μ m boxes, increased from 71.2 ± 5.53 mm⁻¹ (n = 3) at P14 to 180.0 ± 13.0 mm⁻¹ (n = 3) at P18 and 295.0 ± 33.6 mm⁻¹ (n = 5) at P28+ (Fig. 7C). This trend was significant [F(2,8) = 40.8, P < 0.001, based on a \log_{10} transform], indicating a vigorous synaptogenesis during the period of our study. Thus low network interconnectivity before P28+ may serve as an additional factor that limits expression of the theta rhythm in the juvenile in vivo.

DISCUSSION

Physiological development of entorhinal stellate cells

We have found that the intrinsic rhythmic properties of stellate cells in layer II of the medial entorhinal cortex (EC) first develop in the third week after birth. Between postnatal days 14 (P14) and 18 (P18), subthreshold membrane potential oscillations (MPOs) at theta frequency appear, reaching strengths indistinguishable from those observed in the adult (P28+). At the same time, spikes associate with the peaks of the oscillations; the spike afterhyperpolarization (AHP) deepens, shortens, and develops an overshoot; and spikes occur in clusters on consecutive MPO cycles. These data contribute to our understanding of the events that characterize development in these cells. They also serve a practical purpose. In vitro physiologists, who often seek to use young tissue for its resilience to hypoxia, should be aware that the mature condition may be studied only in EC stellate cells taken from animals at least 3 wk old.

Mechanisms of spontaneous MPO generation

The simultaneous appearance of MPOs, AHP overshoot, and clustering between P14 and P18 is probably not a coincidence because they all reflect a propensity for resonance. MPOs, for

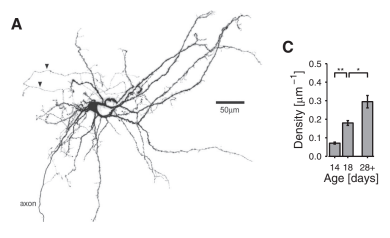
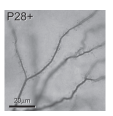


FIG. 7. Anatomical development of stellate cells. A: stellate morphology. P14 stellate cell filled with Alexa Fluor 488 and displaying typical stellate morphology, including strong primary dendrites extending superficially to the pial surface, dendritic spines, and fine axon collaterals (arrowheads) in the vicinity of the soma. An axon projects from a basal primary dendrite, eventually to send collaterals in layers III and VI (cropped). B and C: spine growth. B: photomicrographs of example DAB-stained cells at P14, P18, and P30, showing an increase in spine density. C: histogram of mean spine density in cells at P14 (n=3), P18 (n=3), and P28+ (n=5).



В





example, are generally the product of amplified resonance, whereby a basic resonance supplied by a slow rectifying current is amplified by the positive feedback of a regenerative current (Hutcheon and Yarom 2000). An overshooting (underdamped) response to a perturbation is a property of any resonant system and in EC layer II stellate cells the robust rebound and overshoot of the AHP have been shown to be essential for spike clustering (Fernandez and White 2008; Nolan et al. 2007). We may therefore characterize development between P14 and P18 as being one of increasing resonance.

Direct evidence for this resonance was obtained from our impedance measurements, which showed that P14 cells were less resonant than both P18 and P28+ cells at potentials between rest and spike threshold, some P14 cells showing no resonance at all. The increased resonance with age appears not to be the result of simple changes in passive resistance. In the mature stellate cell, artificially decreasing resistance results in a less resonant response (Fernandez and White 2008), whereas pharmacological block of background conductance amplifies oscillation (Dickson et al. 2000b). We found resistance to fall throughout development, suggesting that resonance was achieved despite membrane shunting. This observation would explain how theta prominence remained constant from P18 to P28+, despite a loss in overall power.

To investigate the cause of resonance, and thus the development of MPOs, we examined the developmental expression of two currents thought to generate MPOs in the adult, the h-current I_h and the persistent sodium current I_{NaP} (Alonso and Llinás 1989; Dickson et al. 2000b; although see following text). According to current models, I_h and I_{NaP} fulfill respectively the roles of slow rectifier and regenerative current described earlier (Dickson et al. 2000b). Additionally, I_{NaP} is a noisy current and is responsible for maintaining oscillation by continually agitating the system (Dorval and White 2005; White et al. 1998). Even at P14, both I_h and I_{NaP} were already established in EC stellate cells. This finding is consistent with the role that both these currents play in the hippocampus, such as in immature rhythms (Bender et al. 2005; Sipila et al. 2006; Strata et al. 1997). Nevertheless, both currents underwent changes in expression levels that could have affected MPO

Of the two currents, the profile of $I_{\rm NaP}$ was the more straightforward. Its conductance density increased by 50% from P14 to P18 and remained elevated at P28+. In a simulation including both $I_{\rm h}$ and $I_{\rm NaP}$ plus a shunt (leak) conductance, changes in $I_{\rm NaP}$ density of this order were found to have substantial effects on theta power both through extra noise and particularly amplification. So large was the effect that we found it necessary to substantially increase shunt conductance with age to prevent unbridled oscillation. These results indicate that $I_{\rm NaP}$ could be instrumental in the appearance of MPOs during week 3.

The interpretation of I_h development is more difficult. Although the conductance density of I_h increased significantly from P14 to P18, it was intermediate at P28+ and not significantly different from either the P14 or P18 value. It is thus not clear whether I_h expression exerts much influence over physiology beyond P14. If I_h expression were genuinely intermediate at P28+, this might explain why AHP overshoot, which is influenced by I_h (Nolan et al. 2007), fell from P18 to P28+ (albeit nonsignificantly). Nevertheless, AHP overshoot is likely

to be determined by a number of factors, including shunt conductance (see Fernandez and White 2008), which increased over the same period. Parsimoniously, we suggest it is best to assume that I_h is not a limiting factor in the development of MPOs during week 3, even if it may temporarily contribute to increasing resonance. This interpretation would be consistent with our measured responses to hyperpolarizing current steps, which showed that effective I_h -induced sag strength was constant throughout the study period. In the hippocampus, too, I_h expression undergoes only modest change in week 3 and even peaks before P14 (Bender et al. 2001, 2005; Vasilyev and Barish 2002). Furthermore, in adult EC stellate cells, large reductions in I_h levels of \geq 65% are necessary for MPOs to be abolished (Dickson et al. 2000b; Klink and Alonso 1993). The range of values we report is thus smaller than would normally be expected to affect the MPO. This conclusion is supported by our simulations in which the relative changes in I_h density we measured in vitro had only modest effects on the existence of MPOs.

According to the preceding line of argument, we may present a model for MPO development in which developing I_{NaP} expression is pivotal, whereas I_{h} is already at levels adequate for mature oscillation at P14. This does not mean, however, that other currents are unimportant. Nolan et al. (2007) recently reported that knock out of the HCN1 channel protein, which carries the majority of I_h in stellate cells, does not affect theta-range voltage fluctuations. This result is difficult to interpret because the authors did not measure the concentration of power at theta relative to other frequencies and because the data shown by Nolan et al. (2007) suggest to us that MPOs are less prominent in mice than in rats. Nevertheless, since the work of Dickson et al. (2000b) implicating I_h in MPO generation, it has been shown that the drug they applied to block I_h , ZD7288, also blocks T-type Ca²⁺ currents (Felix et al. 2003; Sánchez-Alonso et al. 2008). Because stellate cells possess these currents (Bruehl and Wadman 1999), it is possible that the abolition of MPOs that Dickson et al. (2000b) reported represents the block of I_h , T-type Ca²⁺ channels, and Ca²⁺-dependent K⁺ channels (also present; Eder et al. 1991). Interestingly, conventional blockers of $I_{\rm b}$ and Ca²⁺ channels do not eliminate MPOs when applied on their own (Klink and Alonso 1993); neither do blockers of K⁺ channels (Dickson et al. 2000b; Yoshida and Alonso 2007). It may thus be profitable in the future to consider the balance of several currents in the generation of MPOs, both in the adult and during development.

Functional consequences of developmental changes

The function of MPOs per se in stellate cells is at present unclear. In vitro, MPOs are sensitive to synaptic conductance and are eliminated in the presence of incoherent synaptic input (Fernandez and White 2008). Nevertheless, as the product of an underlying resonance, MPOs indicate a competence for stellate cells to respond to oscillatory input. In the aroused state, stellate cells receive strong theta drive from the medial septum (Dickson et al. 1994; Mitchell et al. 1982) and most likely the EC deep layers as well (Gloveli et al. 1999). Under such conditions, it is likely that a resonant membrane will allow the stellate cell to tune its activity to theta frequencies. Thus the appearance of electrical resonance and subthreshold

MPOs in the theta range between P14 and P18 suggests that, as a whole, the EC layer II is not able to process theta effectively at P14 but becomes so within the following week.

Our anatomical data support this hypothesis. The dendritic spines of stellate cells receive synaptic input from both extrinsic sources and other stellates within layer II (Kumar et al. 2007; Witter 1993). They therefore supply a means by which the oscillatory properties of stellate cells could not only tune the cell to septal drive but also allow stellates to synchronize directly through reciprocal interactions (Alonso and Llinás 1989; Llinás 1988). At P14, dendrites were relatively free of spines, suggesting a limited capacity to receive input of any kind. By P18, however, spine density had more than doubled and by P28+ it had quadrupled. This pattern is very similar to that seen in the hippocampus CA1, where spines first appear around P11-P12 and witness a two- to fourfold increase from P15 to adulthood (Fiala et al. 1998; Harris et al. 1992). The exact age at which a full adult complement of spines is achieved is not clear because we did not study cells between P18 and P28+; thus we cannot rule out the possibility that new spines continue to develop right up to P28 or beyond. Nevertheless, almost adult spine densities and homeostasis are attained by P22 in hippocampus CA1 (Kirov et al. 2004). It is thus quite possible that both intrinsic and network sources of oscillation appear simultaneously in the EC and reach near completion by around the end of week 3.

More generally, the changes we observe fit into a trend in which immature patterns of activity involved with the formation of functional circuits give way to integrative functions in the third week after birth, contemporaneously with rapid synaptogenesis (Ben-Ari 2001; Dumas 2005). For the hippocampal formation, mature functions notably include spatial memory (O'Keefe and Nadel 1978). Several studies indicate that the theta rhythm is essential for proper memory function in the hippocampal formation, at both the synaptic and the behavioral levels (Huerta and Lisman 1993; Hyman et al. 2003; Mitchell et al. 1982). Our finding that EC layer II stellate cells develop mature MPOs between P14 and P18 may thus help to explain why many spatial memory functions are poorly developed or absent until the end of week 3 (Carew and Rudy 1991; Carman and Mactutus 2001; Green and Stanton 1987; Raunch and Raskin 1984; Rudy 1993; Rudy et al. 1987). However, perhaps the most striking feature of the EC in particular—and thus the feature of most relevance to our study—is the EC representation of space. "Grid cells" of the superficial layers of the medial EC, which presumably include stellate cells, display activity that is modulated according to the position of the rat on an abstract lattice across the laboratory floor (Hafting et al. 2005). Different cells define different grids of different orientations and phases and a gradient in grid spacing exists from narrow in the most dorsal EC to wide in the more ventral EC (Hafting et al. 2005). Presently we do not know how the grid cell representation is formed, although current theories implicate the theta rhythm (Blair et al. 2007; Burgess et al. 2007; O'Keefe and Burgess 2005). In one recent study, it was reported that stellate cells along the dorsoventral extent of EC exhibit a gradient in the frequency of theta MPOs that, when applied to the Burgess model, produced grid spacings as observed in vivo (Giocomo et al. 2007). This correspondence strongly supports the notion that the theta resonance of entorhinal stellate cells is important for spatial coding. If so, then

we would predict that grid cells appear in the rat no earlier than about P18.

ACKNOWLEDGMENTS

We thank T. Blute for training on the confocal microscope and J. Magistretti for advice on recording solutions and techniques.

Present address of J. White: Department of Bioengineering, University of Utah, 20 S. 2030 E., Salt Lake City, UT 84112.

GRANTS

This work was supported by the National Institutes of Health.

REFERENCES

- Alonso A, García-Austt E. Neuronal sources of theta rhythm in the entorhinal cortex of the rat. I. Laminar distribution of theta field potentials. *Exp Brain Res* 67: 493–501, 1987.
- **Alonso A, Klink R.** Differential electroresponsiveness of stellate and pyramidal-like cells of medial entorhinal cortex layer II. *J Neurophysiol* 70: 128–143, 1993.
- **Alonso A, Llinás RR.** Subthreshold Na⁺-dependent theta-like rhythmicity in stellate cells of entorhinal cortex layer II. *Nature* 342: 175–177, 1989.
- **Ben-Ari Y.** Developing networks play a similar melody. *Trends Neurosci* 24: 353–359, 2001.
- Bender RA, Brewster A, Santoro B, Ludwig F, Biel M, Baram TZ. Differential and age-dependent expression of hyperpolarization-activated, cyclic nucleotide-gated cation channel isoforms 1–4 suggests evolving roles in the developing rat hippocampus. *Neuroscience* 106: 689–698, 2001.
- Bender RA, Galindo R, Mameli M, Gonzalez-Vega R, Valenzuela F, Baram TZ. Synchronized network activity in developing rat hippocampus involves regional hyperpolarization-activated cyclic nucleotide-gated (HCN) channel function. *Eur J Neurosci* 22: 2669–2674, 2005.
- **Blair HT, Welday AC, Zhang K.** Scale-invariant memory representations emerge from Moiré interference between grid fields that produce theta oscillations. A computational model. *J Neurosci* 27: 3211–3229, 2007
- **Bland BH, Oddie SD.** Theta band oscillation and synchrony in the hippocampal formation and associated structures: the case for its role in sensorimotor integration. *Behav Brain Res* 127: 119–136, 2001.
- **Bruehl C, Wadman WJ.** Calcium currents in acutely isolated stellate and pyramidal neurons of rat entorhinal cortex. *Brain Res* 816: 554–562, 1999.
- Burgess N, Barry C, O'Keefe J. An oscillatory interference model of grid cell firing. *Hippocampus* 17: 801–812, 2007.
- Carew MB, Rudy JW. Multiple functions of context during conditioning: a developmental analysis. *Dev Psychobiol* 24: 191–209, 1991.
- Carman HM, Mactutus CF. Ontogeny of spatial navigation in rats: a role for response requirements? *Behav Neurosci* 115: 870–879, 2001.
- Dickson CT, Magistretti J, Shalinsky MH, Bassam H, Alonso A. Oscillatory activity in entorhinal neurons and circuits. Mechanisms and function. Ann NY Acad Sci 911: 127–150, 2000a.
- Dickson CT, Magistretti J, Shalinsky MH, Fransén E, Hasselmo ME, Alonso A. Properties and role of I_h in the pacing of subthreshold oscillations in entorhinal cortex layer II neurons. *J Neurophysiol* 83: 2562–2579, 2000b.
- **Dickson CT, Trepel C, Bland BH.** Extrinsic modulation of theta field activity in entorhinal cortex of anesthetized rat. *Hippocampus* 4: 37–51, 1994.
- **Dorval AD, White JA.** Channel noise is essential for perithreshold oscillations in entorhinal stellate neurons. *J Neurosci* 25: 10025–10028, 2005.
- **Dumas TC.** Developmental regulation of cognitive abilities: modified composition of a molecular switch turns on associative learning. *Prog Neurobiol* 76: 189–211, 2005.
- Eder C, Ficker E, Gündel J, Heinemann U. Outward currents in rat entorhinal cortex stellate cells studied with conventional and perforated patch recordings. *Eur J Neurosci* 3: 1271–1280, 1991.
- Erchova I, Kreck G, Heinemann U, Herz AVM. Dynamics of rat entorhinal cortex layer II and III cells: characteristics of membrane potential resonance at rest predict oscillation properties near threshold. *J Physiol* 560: 89–110, 2004.
- Felix R, Sandoval A, Sánchez D, Cómora JC, De la Vega-Beltrán JL, Treviño CL, Darszon A. ZD7288 inhibits low-threshold Ca²⁺ channel

- activity and regulates sperm function. *Biochem Biophys Res Commun* 311: 187–192. 2003.
- **Fernandez FR, White JA.** Artificial synaptic conductances reduce subthreshold oscillations and periodic firing in stellate cells of the entorhinal cortex. *J Neurosci* 28: 3790–3803, 2008.
- Ferrer I, Martinez-Matos JA. Development of non-pyramidal neurons in the rat sensory-motor cortex during the fetal and early postnatal periods. J Hirnforsch 22: 555–562, 1981.
- Fiala JC, Feinberg M, Popov V, Harris KM. Synaptogenesis via dendritic filopodia in developing hippocampal area CA1. J Neurosci 18: 8900–8911, 1998
- Fransén E, Alonso AA, Dickson CT, Magistretti J, Hasselmo ME. Ionic mechanisms in the generation of subthreshold oscillations and action potential clustering in entorhinal layer II stellate neurons. *Hippocampus* 14: 368–384, 2004.
- **Giocomo LM, Zilli EA, Fransén E, Hasselmo ME.** Temporal frequency of subthreshold oscillations scales with entorhinal grid cell field spacing. *Science* 315: 1719–1722, 2007.
- Gloveli T, Egorov AV, Schmitz D, Heinemann U, Müller W. Carbacholinduced changes in excitability and [Ca²⁺]_i signaling in projection cells of medial entorhinal cortex layers II and III. Eur J Neurosci 11: 3626–3636, 1999.
- **Green RJ, Stanton ME.** Differential ontogeny of working memory and reference memory in the rat. *Behav Neurosci* 103: 98–105, 1987.
- Haas JS, Dorval AD, White JA. Contributions of I_h to feature selectivity in layer II stellate cells of the entorhinal cortex. J Comput Neurosci 22: 161–171, 2007.
- Haas JS, White JA. Frequency selectivity of layer II stellate cells in the medial entorhinal cortex. J Neurophysiol 88: 2422–2429, 2002.
- Hafting T, Fyhn M, Molden S, Moser M, Moser EI. Microstructure of a spatial map in the entorhinal cortex. *Nature* 436: 801–806, 2005.
- **Harris KM, Jensen FE, Tsao B.** Three dimensional structure of dendritic spines and synapses in rat hippocampus (CA1) at postnatal day 15 and adult ages: implications for the maturation of synaptic physiology and long-term potentiation. *J Neurosci* 12: 2685–2705, 1992.
- **Horikawa K, Armstrong WE.** A versatile means of intracellular labeling: injection of biocytin and its detection with avidin conjugates. *J Neurosci Methods* 25: 1–11, 1988.
- **Howell DC.** Statistical Methods for Psychology (4th ed.). Belmont, CA: Duxbury Press, 1997.
- **Huerta PT, Lisman JE.** Heightened synaptic plasticity of hippocampal CA1 neurons during a cholinergically induced rhythmic state. *Nature* 364: 723–725, 1993.
- **Hutcheon B, Yarom Y.** Resonance, oscillation and the intrinsic frequency preferences of neurons. *Trends Neurosci* 23: 216–222, 2000.
- **Huxter J, Burgess N, O'Keefe J.** Independent rate and temporal coding in hippocampal pyramidal cells. *Nature* 425: 828–832, 2003.
- Hyman JM, Wyble BP, Goyal V, Rossi CA, Hasselmo ME. Stimulation in hippocampal region CA1 in behaving rats yields long-term potentiation when delivered to the peak of theta and long-term depression when delivered to the trough. J Neurosci 23: 11725–11731, 2003.
- **Jensen O, Lisman JE.** Position reconstruction from an ensemble of hippocampal place cells: contribution of theta phase coding. *J Neurophysiol* 83: 2602–2609, 2000.
- **Kirov SA, Goddard CA, Harris KM.** Age-dependence in the homeostatic upregulation of hippocampal dendritic spine number during blocked synaptic transmission. *Neuropharmacology* 47: 640–648, 2004.
- Klink R, Alonso A. Ionic mechanisms for the subthreshold oscillations and differential electroresponsiveness of medial entorhinal cortex layer II neurons. J Neurophysiol 70: 144–157, 1993.
- **Klink R, Alonso A.** Morphological characteristics of layer II projection neurons in the rat medial entorhinal cortex. *Hippocampus* 7: 571–583, 1007
- **Kocsis B, Bragin A, Buzsáki G.** Interdependence of multiple theta generators in the hippocampus. A partial coherence analysis. *J Neurosci* 19: 6200–6212, 1999.
- **Kramis RC, Vanderwolf CH, Bland BH.** Two types of hippocampal rhythmical slow activity in both the rabbit and the rat: relations to behavior and effects of atropine, diethyl, ether, urethane and pentobarbital. *Exp Neurol* 49: 58–85, 1975.
- **Kumar SS, Jin X, Buckmaster PS, Huguenard JR.** Recurrent circuits in layer II of medial entorhinal cortex in a model of temporal lobe epilepsy. *J Neurosci* 27: 1239–1246, 2007.

- Larson J, Wong D, Lynch G. Patterned stimulation at the theta frequency is optimal for the induction of hippocampal long-term potentiation. *Brain Res* 368: 347–350, 1986.
- Lingenhöhl K, Finch DM. Morphological characterization of rat entorhinal neurons in vivo soma-dendritic structure and axonal domains. Exp Brain Res 84: 57–74, 1991.
- **Llinás RR.** The intrinsic electrophysiological properties of mammalian neurons. Insights into central nervous system function. *Science* 242: 1654–1664, 1988.
- **Longin A, Souchier C, ffrench M, Byron P-A.** Comparison of antifading agents used in fluorescence microscopy. Image analysis and laser confocal microscopy study. *J Histochem Cytochem* 41: 1833–1840, 1993
- Magistretti J, Alonso A. Biophysical properties and slow voltage-dependent inactivation of a sustained sodium current in entorhinal cortex layer-II principal neurons: a whole-cell and single-channel study. *J Gen Physiol* 114: 491–509, 1999.
- Maxwell SE, Delaney HD. Designing Experiments and Analyzing Data. A Model Comparison Perspective (2nd ed.). Philadelphia, PA: Erlbaum, 2004
- **Mitchell SJ, Ranck JBJ.** Generation of theta rhythm in medial entorhinal cortex of freely moving rats. *Brain Res* 189: 49–66, 1980.
- **Mitchell SJ, Rawlins JNP, Steward O, Olton DS.** Medial septal area lesions disrupt theta rhythm and cholinergic staining in medial entorhinal cortex and produce impaired radial arm behavior in rats. *J Neurosci* 2: 292–302, 1982.
- Neher E. Correction for liquid junction potentials in patch clamp experiments. *Methods Enzymol* 207: 123–131, 1992.
- **Nolan MF, Dudman JT, Dodson PD, Santoro B.** HCN1 channels control resting and active integrative properties of stellate cells from layer II of the entorhinal cortex. *J Neurosci* 27: 12440–12451, 2007.
- O'Keefe J, Burgess N. Dual phase and rate coding in hippocampal place cells. Theoretical significance and relationship to entorhinal grid cells. *Hippocampus* 15: 853–866, 2005.
- O'Keefe J, Recce ML. Phase relationship between hippocampal place units and the EEG theta rhythm. *Hippocampus* 3: 317–330, 1993.
- O'Keefe JA, Nadel L. The Hippocampus as a Cognitive Map. Oxford, UK: Oxford Univ. Press, 1978.
- **Orr G, Rao G, Houston FP, McNaughton BL, Barnes CA.** Hippocampal synaptic plasticity is modulated by theta rhythm in the fascia dentata of adult and aged freely behaving rats. *Hippocampus* 11: 647–654, 2001.
- Papoulis A. Probability, Random Variables and Stochastic Processes (3rd ed.). New York: McGraw-Hill, 1991.
- **Pokorný J, Yamamoto T.** Postnatal ontogenesis of hippocampal CA1 area in rats. II. Development of ultrastructure in stratum lacunosum and moleculare. *Brain Res Bull* 7: 121–130, 1981.
- **Raunch SL, Raskin LA.** Cholinergic mediation of spatial memory in the preweanling rat: application of the radial arm maze paradigm. *Behav Neurosci* 98: 35–43, 1984.
- Rudy JW. Contextual conditioning and auditory cue conditioning dissociate during development. *Behav Neurosci* 107: 887–891, 1993.
- **Rudy JW, Stadler-Morris S, Albert P.** Ontogeny of spatial navigation behaviors in the rat: dissociation of "proximal" and "distal" cue-based behaviors. *Behav Neurosci* 101: 62–73, 1987.
- **Sainsbury RS.** Hippocampal theta: a sensory-inhibition theory of function. *Neurosci Biobehav Rev* 22: 237–241, 1998.
- Sánchez-Alonso JL, Halliwell JV, Colino A. ZD7288 inhibits T-type calcium current in rat hippocampal pyramidal cells. *Neurosci Lett* 439: 275–280, 2008.
- Sargolini F, Fyhn M, Hafting T, McNaughton BL, Witter MP, Moser M, Moser EI. Conjunctive representation of position, direction and velocity in entorhinal cortex. *Science* 312: 758–762, 2006.
- Seidenbecher T, Laxmi TR, Stork O, Pape HC. Amygdalar and hippocampal theta rhythm synchronization during fear memory retrieval. *Science* 301: 846–850, 2003.
- **Siapas AG, Lubenov EV, Wilson MA.** Prefrontal phase locking to hippocampal theta oscillations. *Neuron* 46: 141–151, 2005.
- **Sipila ST, Huttu K, Voipio J, Kaila K.** Intrinsic bursting of immature CA3 pyramidal neurons and consequent giant depolarizing potentials are driven by a persistent Na⁺ current and terminated by a slow Ca²⁺-activated K⁺ current. *Eur J Neurosci* 23: 2330–2338, 2006.
- Steward O, Scoville SA. Cells of origin of entorhinal cortical afferents to the hippocampus and fascia dentata of the rat. J Comp Neurol 169: 347–370, 1976.

- Strata F, Atzori M, Molnar M, Ugolini G, Tempia F, Cherubini E. A pacemaker current in dye-coupled hilar interneurons contributes to the generation of giant GABA-ergic potentials in developing hippocampus. *J Neurosci* 17: 1435–1446, 1997.
- **Vanderwolf CH.** Hippocampal electrical activity and voluntary movement in the rat. *Electroencephalogr Clin Neurophysiol* 26: 407–418, 1969.
- **Vasilyev DV, Barish ME.** Postnatal development of the hyperpolarization-activated excitatory current $I_{\rm h}$ in mouse hippocampal pyramidal neurons. *J Neurosci* 22: 8992–9004, 2002.
- Wang X, Lambert NA. Membrane properties of identified lateral and medial perforant pathway projection neurons. *Neuroscience* 117: 485–492, 2003.
- White JA, Klink R, Alonso A, Kay AR. Noise from voltage-gated ion channels may influence neuronal dynamics in the entorhinal cortex. *J Neurophysiol* 80: 262–269, 1998
- **Winson J.** Loss of hippocampal theta rhythm results in spatial memory deficit in the rat. *Science* 201: 160–163, 1978.
- **Witter M.** Organization of the entorhinal-hippocampal system: a review of current anatomical data. *Hippocampus* 3: 33–44, 1993.
- **Yoshida M, Alonso A.** Cell-type-specific modulation of intrinsic firing properties and subthreshold membrane oscillations by the M(Kv7)-current in neurons of the entorhinal cortex. *J Neurophysiol* 98: 2779–2794, 2007.