High-Frequency Oscillations in the Output Networks of the Hippocampal–Entorhinal Axis of the Freely Behaving Rat

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Population bursts of the CA3 network, which occur during eating, drinking, awake immobility, and slow-wave sleep, produce a large field excitatory postsynaptic potential throughout stratum radiatum of the CA1 field (sharp wave). The CA3 burst sets into motion a short-lived, dynamic interaction between CA1 pyramidal cells and interneurons, the product of which is a 200 Hz oscillatory field potential (ripple) and phase-related discharge of the CA1 network. Although many CA1 pyramidal neurons discharge during the time frame (50–100 msec) of each sharp wave, each wave of a ripple (∼5 msec) reflects the synchronization of more discrete subsets of CA1 neurons.

When we used multi-site recordings in freely behaving rats, we observed ripples throughout the longitudinal extent (∼4–5 mm) of the dorsal CA1 region that were coherent for multiple cycles of each ripple. High-frequency ripples were also observed throughout the hippocampal–entorhinal output pathway that were concurrent but less coherent on a cycle-by-cycle basis. Single and multiunit neuronal activity was phase-related to local ripples throughout the hippocampal–entorhinal output pathway. Entorhinal ripples occurred 5–30 msec after the CA1 ripples and were related to the occurrence of an entorhinal sharp wave. Thus, during each hippocampal sharp wave, there is a powerful synchronization among the neuronal networks that connect the hippocampus to the neocortex. We suggest that this population interaction (1) biologically constrains theoretical models of hippocampal function and dysfunction and (2) has the capacity to support an “off-line” memory consolidation process.

Key words: hippocampus; subiculum; entorhinal cortex; oscillations; sharp waves; memory; epilepsy

The hippocampus and entorhinal cortex (EC) are substrates for the formation of enduring memories and the etiology/pathology of Alzheimer’s dementia as well as temporal lobe epilepsy. The manner and mechanisms by which these structures interact have implications for both physiological and pathophysiological processes.

Synchronous population potentials and localized oscillatory patterns occur in many neural networks (Buzsáki et al., 1983, 1992; Steriade et al., 1993; Freeman and Barrie, 1994; Gray, 1994; Singer, 1994). These events reflect synchronizing mechanisms for coordinating ensembles of neurons within distributed neural networks and for bringing neuronal aggregates within/across structures together in time (Buzsáki and Chrobak, 1995).

The physiology of the hippocampal–entorhinal networks is characterized by behaviorally regulated, macroscopic potentials that temporally organize specific subsets of network neurons. θ occurs during exploratory behavior and rapid eye movement sleep and represents a period when hippocampal circuits receive rhythmic input from neurons within the superficial layers of the EC (Mitchell and Ranck, 1980; Buzsáki et al., 1983; Boeijinga and Lopes da Silva, 1988; Stewart et al., 1992). In contrast, during sharp waves, which occur during consummatory behaviors and slow-wave sleep, the output neurons of the hippocampus and EC participate in organized population bursts (Buzsáki, 1989; Chrobak and Buzsáki, 1994; Ylinen et al., 1995). These patterns seem to serve companion processes. θ synchronizes the input pathway into the hippocampus, whereas sharp waves synchronize the output pathway from the hippocampus back to neocortical structures. Each potential is associated with more localized oscillatory field potentials that reflect the temporal organization of neuronal subsets within 5–25 msec (Bragin et al., 1995; Buzsáki and Chrobak, 1995; Ylinen et al., 1995).

The sharp wave is a large amplitude (1–3 mV), aperiodic, field potential observed most prominently in stratum radiatum of the CA1 region (Buzsáki et al., 1983; Buzsáki, 1986; Suzuki and Smith, 1987). Released from inhibitory constraints associated with θ (Leung and Yim, 1986; Fox, 1989; Soltész and Deschénes, 1993), the highly interconnected CA3 network exhibits population bursts. The burst of the CA3 network produces a field excitatory postsynaptic potential (EPSP) (a sharp wave) in the target of the CA3 Schaffer collaterals, the dendrites of CA1 pyramidal cells, and interneurons. The massive depolarization of CA1 sets into motion a short-lived, dynamic interaction between these cell populations. The product of this interaction is an oscillatory field potential (ripple) within stratum pyramidale and a phase-related discharge of the CA1 network at 200 Hz (Buzsáki et al., 1992). The synaptic currents mediating the ripple are rhythmic inhibitory postsynaptic potentials (IPSPs) near the soma of CA1 neurons produced by a high-frequency discharge of CA1 basket cells and other interneurons (Ylinen et al., 1995). Although interneurons discharge at a high frequency, CA1 pyramidal cells, excited by the CA3 input yet constrained by the interneuron barrage, typically fire once or not at all. Throughout the entire CA1 network,
however, a large number of pyramidal cells reach discharge threshold on each ~5 msec wave of the ripple.

The present study examined the interaction between this synchronized output of CA3–CA1 and the output neurons of a larger integrated network that includes the subiculum and deep layers of the presubiculum, parasubiculum, and EC. We found that similar to the CA1 region, each output population of the hippocampal–entorhinal network produces an organized, short-lived, fast oscillation of their discharging neurons concurrent with CA1 ripples.

MATERIALS AND METHODS

Animals and surgery. Twenty-four adult Sprague–Dawley rats were used in the following experiments. For surgery, rats were anesthetized with a ketamine cocktail (4 ml/kg) consisting of 25 mg/ml ketamine, 1.3 mg/ml xylazine, and 0.25 mg/ml acepromazine. After a midline scalp incision, two to four burr holes were drilled in the skull over the hippocampal and retrohippocampal cortices. Two or three sets of four 50 µm tungsten wires were positioned into the dorsal hippocampus/subicular regions anterior-posterior (AP), −2.5, −5.0, −7.5 from bregma; medial-lateral (ML), 1.5, 2.5, 4.0; dorsal-ventral (DV), 2.0−4.0 from the skull and above retrohippocampal regions (AP, −7.5−9.0; ML, 3.0−5.0). One or more sets were chronically fixed, whereas the other was attached to a drive wire and a single machine screw. The latter allowed for optimal, postsurgical positioning of electrodes. One or two single tungsten microelectrodes (0.5–3.0 MΩ) mounted to similar drives were positioned over either or both retrohippocampal areas (AP, −7.5−9.0; ML, 3.0−5.0). These mounts allowed for the slow passage of the microelectrode through the retrohippocampal region. In two animals, a 16-channel silicon probe (100 µm tip separations) (Bragin et al., 1995) was positioned via a movable microdrive into the EC. A pair of 150 µm wires were also positioned in the angular bundle (AP, 7.2; ML, 4.2; DV, 4.0) or CA3 region (AP, 4.0; ML, 4.0; DV, 4.0) for stimulation. Two stainless steel watch screws driven into the bone above the cerebellum served as indifferent and ground electrodes. Two or more additional support screws were positioned, and the entire ensemble was secured to the skull with dental acrylic. All electrodes—indifferent, ground, and stimulating—were attached to male pins that were secured in a rectangular 3 × 4 pin array and secured with dental acrylic.

Recording. Bioelectrical activity was recorded in the freely behaving rats during movement, awake immobility, or distinct sleep stages. The headstage of the animal (male pins) was connected to sixteen MOSFET-input operational amplifiers mounted in a female connector. This direct amplification at the headstage serves to eliminate cable movement artifacts (Buzsáki et al., 1989). An attached cable fed into a rotating swivel (Biela, Irvine, CA) allowed for the free rotation of the recording cable and movement of the rodent within a standard plexiglass home cage. An amplifier system (Grass Neurodata Acquisition System, Quincy, MA) and an analog-to-digital hardware/software system (RC Electronics, Santa Barbara, CA) run on a PC computer allowed for direct visualization and storage of electrical activity. Wide band signals (1 Hz–5 kHz) were sampled at 10 kHz (100 µsec) and stored on optical disks.

After optimization of hippocampal microelectrodes for detection of ripples (200 Hz oscillations) within stratum pyramidale of CA1 and sharp waves within stratum radiatum of CA1, the tungsten microelectrode or multielectrode 50 µm wires were lowered through retrohippocampal structures. Discriminable units and/or prominent oscillations were recorded during both sharp waves (awake immobility) and θ states (locomotor activity and paradoxical sleep); 100–400 (400 msec) epochs triggered by the occurrence of a sharp wave and/or ripple were recorded. When possible (depending on the stability of unit recording), additional continuous epochs (30–90 sec) were recorded during both sharp waves and θ states. When prominent retrohippocampal oscillations or a prominent retrohippocampal sharp potential was observed, additional epochs triggered by the occurrence of these events were recorded. After completion of a single pass of the movable microelectrode(s), rats were anesthetized with pentobarbital and perfused with the electrode in situ.

Data processing and analysis. Unit activity and field potentials were filtered digitally (120 dB/octave: unit, band pass 0.5–5.0 kHz; high-frequency ripples, band pass 100–400 Hz) and analyzed off-line on a 486/33 or an IBM RS 6000 computer or both. Putative single units were verified by the absence of spikes ≥1 msec (typically ≥3–5× baseline amplitude) in autocorrelograms, reflecting the refractory period. Remaining unit activity (units ≥2× baseline, with interspike intervals <1 msec) were considered multunit. Ripple peaks were detected after offline filtering (100–400 Hz), using a peak detection algorithm.

Single and multunit activity were cross-correlated with local ripple peaks and local ripple peaks with CA1 ripples, using the ripple peaks as the zero reference point. Local field averages were obtained by averaging wide-band and filtered signals, using ripple peaks or unit pulses as the zero reference.

Histology. Tissue was processed using either thionin stain or a modified silver method that allows for direct visualization of damaged neurons (Gallyas et al., 1990). The latter technique allowed for more direct visualization, and thus localization, of neurons at the electrode tip.

RESULTS

Coupling of CA1 ripples along the long axis of the hippocampus

Transient high frequency oscillations (ripples) at ~200 Hz were observed as electrodes traversed near the CA1 pyramidal cell layer coincident with hippocampal sharp waves. CA1 ripples are defined as a series of 5–15 oscillatory waves of varying amplitude/duration with a peak-to-peak time of ~5 msec (Figs. 1, 2). Our measurements along the longitudinal axis of the hippocampus indicate that several cycles (three to five) of the dynamically developing CA1 ripples are coherent along this dimension for distances of 4–5 mm (n = 6; Fig. 1E). It is important to note that the waxing and waning of CA1 ripples is a localized phenomenon, which is likely to represent the local development of synchrony among subpopulations of CA1 interneurons. Figure 2 illustrates that although distant sites within the dorsal CA1 region are on average coherent over several cycles of a ripple, ripples at specific sites can occur quite independently. Although the absence of a prominent ripple in the contralateral hippocampus was not unusual, occurring 30–50% (Fig. 2C), the absence of a prominent ipsilateral ripple was rare (<5%; Fig. 2B). Ripples developed in both hemispheres of the hippocampus virtually simultaneously, with cross-correlograms demonstrating peak occurrence near the zero-reference ripple; however, there was no cycle-by-cycle synchrony between ripples in the contralateral hippocampus (Fig. 1F).

Coupling of retrohippocampal neurons to local field, and CA1 ripples

While examining the relationship between CA1 ripples and the discharge of retrohippocampal neurons, we observed prominent local field oscillations within the subiculum and deep layer presubiculum, parasubiculum, and entorhinal cortices. We examined the relationship between these retrohippocampal ripples and unit discharge as well as the relationship of the retrohippocampal ripples to CA1 ripples. Given that principle neurons discharge only on some limited number of local ripple events, ripple–ripple cross-correlograms yield a more accurate reflection of population synchrony between CA1 neurons and retrohippocampal neurons. At all sites within the hippocampal–entorhinal output network where high-frequency ripples could be observed, single and mul-
tiunit activity was correlated to the negative phase of the local field oscillation (Figs. 3, 4, 5), although the degree of modulation for multiple cycles of the oscillation varied from site to site. This variation is likely to depend on the position of the recording electrode within the synaptic field generating the ripple, which may not be optimal for recording well isolated units, as well the number of times a neuron discharges during any given ripple. Thus, for example, the subicular neuron illustrated in Figure 3 seems better modulated because this neuron tended to fire more than once, often exhibiting multiple spikes, on any given ripple with which it discharged.

Figure 3 illustrates that ripples in the ipsilateral dorsal subiculum were coherent with CA1 ripples for multiple cycles (n = 4; Fig. 3B), as was observed at sites across the dorsal CA1 region. Ripples at contralateral subicular sites occurred concurrently but were not coherent on a cycle-by-cycle basis (Fig. 3B). Both single (n = 13) and multiunit subicular unit activity were phase-related to the negative peaks of these local high-frequency oscillations (Fig. 3).

High-frequency ripples were also observed near the deep layers of the presubiculum. Figure 4 illustrates the relation of presubicular ripples to an ipsilateral CA1 ripple. Note the qualitative decrement in cycle-by-cycle coherence in the ripple–ripple cross-correlogram (Fig. 4B), although the peak of the cross-correlogram indicates that these dynamically developing events occur virtually simultaneously in both structures. The peaks of ripple–ripple cross-correlograms did not indicate any significant (>5 msec or a single cycle) lag in the occurrence of presubicular ripples as compared with CA1 ripples (Fig. 4B). Single presubicular (n = 8) and multiunit neurons recorded at all sites within the deepest layers near the overlying white matter were modulated powerfully by local high-frequency oscillations (Fig. 4A,D), discharging on the negative peak of the local oscillation.

Entorhinal ripples were more variable and exhibited fewer oscillatory waves per ripple (Figs. 5A, 6). Although an occasional entorhinal ripple preceded the occurrence of CA1 ripples, the majority (~95%) occurred either virtually simultaneously (~±5 msec) or were delayed (>5–30 msec peak-amplitude wave to peak-amplitude wave) as compared with a dorsal CA1 site (Figs. 5A,D; 6A,D).

Entorhinal and parasubicular single and multiunit activity were phase-related to local ripples, although the degree of oscillatory
modulation was clearly less than at CA1, subicular, or presubicu-
lar sites (Fig. 5A,D). Entorhinal neurons were phase-related to the
negative peak of the local oscillation \( n = 17 \), with several
neurons \( n = 6 \) exhibiting two or three oscillatory peaks in their
cross-correlogram (Fig. 5D). It also was observed that the peak
amplitude entorhinal oscillations were not the most prominent
within layers V–VI, but were better expressed in the most super-
facial aspect of layers V, IV, and III.

Entorhinal ripples, observed as electrodes penetrated layers
V–VI and entered the broad expanse of layer III, were associated
with a prominent negative-going wave that we refer to as en-
trhinal sharp waves (Figs. 6, 7). This slow potential reversed in
polarity near layer II, suggesting that it represents a synchronized
field EPSP in the apical dendritic zone of layer V–VI neurons.
This entorhinal sharp wave is similar to that described previously
by Paré and colleagues (1995) in the cat. These authors also
observed that deep layer EC neurons discharge regularly in asso-
ciation with these events, whereas superficial neurons discharged
much less frequently.

Figure 7 illustrates the relationship between entorhinal ripples
and their associated entorhinal sharp waves to the hippocampal
CA1 ripple. Correlations were observed between the occurrence
of these entorhinal sharp waves and CA1 ripples in all animals
(Figs. 6, 7); however, there was an obvious variability in degree of
concurrence of these field events. This variation may represent
differences in the topographical innervation of the EC from CA1.
The topography of CA1 to the EC is such that the septal CA1 and
subiculum project more to the lateral aspect of the EC, whereas
the temporal CA1 projects to the more medial aspect of the EC
(Amaral and Witter, 1995). We have a limited sample over the
rostral-caudal, medial-lateral dimension, and although Figure
illustrates a prominent relationship between dorsal CA1 and
the caudal-medial extent of the EC, additional studies will be
needed to define the topography of the observed physiological
connectivity.

**DISCUSSION**

The present study demonstrates novel findings that integrate the
ripple-related discharge of CA1 neurons during hippocampal
sharp waves with electrophysiological events occurring through-
out the output networks of the hippocampal–entorhinal axis.
First, during hippocampal sharp waves, local field oscillations are
synchronous throughout the dorsal CA1 region, as well as in the
subicular and deep layers of the presubiculum, parasubiculum, and
entorhinal cortices. Second, throughout the hippocampal–ento-
rhinal output network, discharge of neurons synchronized to the
negative peak of locally developing ripples in each region. Third,
entorhinal ripples are associated with a negative-going entorhinal
sharp wave that reverses in polarity near the border of layers
II–III. This latter phenomenon may reflect the synchronized de-
polarization of the apical dendritic field of layer V–VI neurons
by the hippocampal sharp wave-related discharge of CA1, subiculum,
and presubiculum neurons that synapse in this region.

Thus, in association with hippocampal sharp waves, highly
organized discharges occur throughout the entire output network
of the hippocampal–entorhinal axis. These organized discharges
bring tens of thousands of neurons within these interconnected
networks together so that they discharge in discrete subsets on
each of several high-frequency oscillatory waves. This coordinated
population output is likely to have a potent impact on many
forebrain targets.

**Temporal structure in hippocampal–entorhinal neuronal networks: a role for interneurons**

Our observations demonstrate that in association with hippocam-
pal sharp waves, there are locally developing field oscillations
throughout the output network of the hippocampal–entorhinal
axis. The extracellular recorded oscillatory fields may reflect
synchronous membrane oscillations in pyramidal neurons caused
by rhythmic IPSPs. Previous findings demonstrate that the 200 Hz
field oscillation observed within CA1 reflects synchronized IPSPs
in the perisomatic region of CA1 neurons (Ylinen et al., 1995).
Networks of interneurons can achieve rhythmic, synchronous pop-
ulation discharges, which then exert a hyperpolarizing oscillation
on the membrane potential of pyramidal neurons (Michelson and
Wong, 1994; Whittington et al., 1995; Traub et al., in press). Such
oscillations, occurring in the context of a depolarizing input, can
impose a periodic fluctuation in the pyramidal cell membrane

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**Figure 2.** Ripples at various positions with the CA1 region can emerge independently. **A**, **B**, **C**, Single 200 msec traces recorded concurrently in the ipsilateral (5 mmdistance from reference CA1 ripple) and contralateral CA1 pyramidal cell region. Ripples typically occur concurrently at various sites within CA1 (**A**), but the occurrence, amplitude, and duration can be independent at various sites within CA1 (**B**, **C**).
Figure 3. CA1 ripple and associated neuronal events in the ipsilateral and contralateral subiculum. A, Single 400 msec sweep with ripple doublets recorded at three sites and subicular unit. B, Cross-correlograms of ipsilateral (black) and contralateral (gray) subicular ripples to the peak of the CA1 reference ripple (n = 224 CA1 ripples). Note the prominent wave-by-wave coherence on the ipsilateral side and its absence in the contralateral subiculum. Insets in B illustrate position of electrodes in the dorsal subiculum (figures as from Swanson, 1992). C, Averaged subicular ripple (n = 193) and its relation to single (black) and multunit (gray) activity recorded at the same site. D, Zero reference was negative peak of the local subicular ripple. Inset in D, autocorrelogram of single unit.
Figure 4. CA1 ripple and associated neuronal events in the deep layers of the ipsilateral presubiculum. A, Single 400 msec sweeps recorded from CA1 and the ipsilateral presubiculum. Top trace illustrates the discharge of a single (*) and multiunit presubiculum neurons. B, Cross-correlogram of ipsilateral presubiculum ripple to the peak of the CA1 reference ripple (n = 257 CA1 ripples). Inset in B illustrates position of electrode in the dorsal presubiculum. C, Averaged presubiculum ripple (n = 213). D, Cross-correlograms of single (black) and multiunit (gray) activity and presubiculum ripples; zero reference was negative peak of local presubiculum ripple. Inset in D, Autocorrelogram of single unit.
Figure 5. CA1 ripple and associated neuronal events in the deep layers of the ipsilateral entorhinal cortex. A, Single 400 msec sweeps with concurrent ripples in CA1 and the ipsilateral EC. Top trace, discharge of entorhinal neurons at the site where the entorhinal ripple was recorded. B, Cross-correlograms of the ipsilateral entorhinal ripple with the peak of CA1 ripple as zero reference (n = 211). Note the absence of ripple-related modulation in the cross-correlogram. Inset in B illustrates position of electrodes in the rostral EC. C, Averaged entorhinal ripple (n = 213). D, Cross-correlograms of single (black) and multi-unit (gray) activity and entorhinal ripples; zero reference was negative peak of entorhinal ripple. Inset in D, Autocorrelogram of single unit.
close to, but below, discharge threshold. Although CA1 interneurons discharge near 200 Hz, CA1 pyramidal cells typically discharge once or not at all during a ripple. The discharge of CA1 pyramidal cells is thus highly constrained, despite a massive depolarization from the CA3 input.

We suggest that the locally developing field oscillations throughout the hippocampal–entorhinal output network develop in a manner analogous to that observed in CA1. Thus, feedforward excitation simultaneously drives low-threshold networks of interneurons that discharge at a high frequency, whereas pyramidal cells discharge in population bursts: the population burst driven by feed-forward excitation, but concurrently constrained by the influence of the local interneuron barrage. The high degree of transient synchrony observed throughout the network is likely to reflect the anatomical arrangement of (1) feed-forward excitatory projections and/or (2) long-range inhibitory projections. Coupling across regions may be achieved by GABAergic interneurons with wide-ranging axon collaterals (Sik et al., 1995) and/or gap junctions, and such interneuronal “supernetworks” may co-operatively entrain large populations of pyramidal cells in the hippocampal–entorhinal axis (Buzsáki and Chrobak, 1995).

Fast oscillations within the hippocampal/retrohippocampal output pathways during sharp waves: coordinated activity in a distributed neural network

The temporally coordinated activity of neurons within spatially distributed networks is an important functional unit within the nervous system (Hebb, 1949; Konorski, 1949). Rhythmic field potentials reflect temporal frameworks for synchronizing neural activity, i.e., for bringing neuroanatomically distributed and sparsely connected neurons together in time (Buzsáki et al., 1983; Steriade et al., 1990, 1993; Gray, 1994). Although sharp waves are aperiodic, they are associated with a high-frequency oscillation within the entire hippocampal–entorhinal output network. Synchronized neural discharges occur within this large network during these events. It is important to note that synchronized oscillations and unit discharges do not occur throughout the entire network with each and every sharp wave; rather they emerge within varying topographical locations and involve varying subpopulations of neurons. The degree of synchronization between any two sites within the network varies from event to event and is likely to reflect anatomical/synaptic connectivity. Clearly, the degree of synchronization across the dorsal CA1 and adjacent sub-
Figure 7. CA1 ripple and associated high-frequency oscillations in the EC. A, Single 400 msec sweeps with concurrent ripples in the CA1 region and both hemispheres of the EC. Note the time lag and lower frequency of the entorhinal oscillations. B, Averaged extracellular fields (wide band) in the hippocampus and ipsilateral EC triggered by the negative peak of CA1 ripples \((n = 243)\). Top trace is average extracellular fields in stratum oriens. Bottom traces are averaged entorhinal sharp waves illustrating reversal near the border of layers II–III. C, Average extracellular fields in the same EC locations triggered by the negative peak of the entorhinal high-frequency oscillation. Top and bottom traces (1–5 kHz) are layers III and II sites as shown in B, respectively. Middle traces are the same sites filtered (100–400 Hz) for ripples. Note reversal of high-frequency oscillation. D, Relation of contralateral entorhinal ripples (black) and ipsilateral CA1 ripples (gray) to the negative peak of the ipsilateral entorhinal ripple (same zero reference as C).
icular region, over many ripple events, is much greater within any given hemisphere as compared with the contralateral hemisphere (Figs. 1–3) or ipsilateral retrohippocampal sites (Figs. 4–6). As suggested above, the degree of synchronization within/across regions may reflect the anatomical interconnectivity of interneuronal networks. In this regard, it is important to appreciate the extensive interconnectivity of GABAergic interneurons within the CA1 region (Sik et al., 1995), the lack of substantial interconnections among CA1 pyramidal neurons (Christian and Dudek, 1988; Thomson and Radpour, 1991), and the lack of any significant contralateral projections from either CA1 interneurons or pyramidal neurons (Amaral and Witter, 1995).

Bringing neurons within the hippocampal–retrohippocampal network together in time on such a short time-scale provides a potent means of enhancing their impact on common postsynaptic targets. If convergent activation of neurons is related to potentiating synaptic efficacy (Bliss and Lømo, 1973; Bliss and Collinridge, 1993), then the convergent activity of the hippocampal–entorhinal output network is a potent, endogenous, means for modifying the synaptic connectivity within this network and in its anatomical targets throughout the forebrain.

Relevance of hippocampal–entorhinal population oscillations to cognitive/computational processes

We have suggested that formation and consolidation of memories within this network involves two distinct companion processes (Buzsáki, 1989; Chrobak and Buzsáki, in press). The initial transfer of information into the circuitry of the hippocampus is thought to be supported by θ-modulated γ oscillations within the entorhinal–dentate axis and at their targets within CA3–CA1 (Bragin et al., 1995; Chrobak and Buzsáki, 1995; in press). This on-line modification of preexistent network connectivity serves as a means of imprinting information on the output circuitry of the hippocampus. Free from the ongoing transfer of input into its circuitry, the CA3-CA1-retrohippocampal network then engages in synaptic modification of its own and target neocortical networks. The synchronization of the output circuitry associated with each sharp-wave burst is thought to play a fundamental role in a protracted consolidation process, “whereby medial temporal lobe structures direct the gradual establishment of memory representations in neocortex” (Squire and Alvarez, 1995; also see Marr, 1971). This general framework is compatible with a considerable volume of evidence documenting the temporal gradient of retrograde amnesia associated with temporal lobe insult (Ogden and Corkin, 1991; Squire, 1992; Cho et al., 1993; Kim et al., 1995). This literature demonstrates that the hippocampal–entorhinal substrates participate in a consolidation process that takes place after the acquisition of information and over a variable period of minutes, hours, days, weeks, and even many months transfers representations, or access to representations, to a neocortical substrate. This framework is also compatible with recent multunit recordings in freely behaving rats demonstrating correlated neural activity among pairs of neurons as a consequence of recent experience that is maintained during subsequent sleep periods (Wilson and McNaughton, 1994; Kudrimoti et al., 1995), as well as theoretical modeling studies that have examined the relative efficiency of this dual-process approach for modifying neural networks (McClelland et al., 1995).

We suggest that the major physiological role of this organized network burst is long-term alterations in synaptic efficacy. The synchronized 200 Hz discharge of the hippocampal–entorhinal output network (1) occurs within the appropriate anatomical pathway, so as to convey the outcome of hippocampal processing to neocortical networks, (2) occurs within a logical time domain, so as to be relevant to an “after the fact” consolidation process, and (3) may provide the prerequisite depolarizing force needed to produce synaptic modifications of neocortical networks.

Entorhinal sharp waves: the impact of the output network on the input (layers II–III) network

Cells within the deep layers of the presubiculum and parasubiculum appear continuous with the principal cell layer of the subiculum and the deep layers of the EC (Amaral and Witter, 1995). Our present findings support the continuity of this functionally integrated output network.

Previous findings demonstrated that the superficial layer (II–III) neurons within these regions do not seem to be influenced by hippocampal sharp waves (Chrobak and Buzsáki, 1994). Anatomical circuits exist for the activation of layer II–III neurons by the hippocampal–entorhinal output network, which would then allow for reentrant activation of the hippocampus via the perforant path; however, the physiological activity of this circuitry seems to be constrained powerfully by dominant inhibitory influences within the superficial layers (Finch et al., 1986, 1988; Jones and Heinemann, 1988; Bartesaghi et al., 1989; Jones and Lambert, 1990; Jones and Buhl, 1993; Chrobak and Buzsáki, 1995; Paré et al., 1995). This dominant inhibition, however, may provide mechanisms for a circumscribed activation of superficial layer neurons, and this possibility is under investigation. On the other hand, a general compromise of this inhibition would expose this circuitry to an unaccustomed excitatory barrage. Failure of inhibition at this critical juncture is likely to play a prominent role in the transition of physiological network bursting to pathophysiological reverberating epileptiform activity (Paré et al., 1992; Jones, 1993). We believe that understanding the basic physiology of this intricate neural machinery at the level of an operational system is essential for describing the neural substrate of memory formation as well as the pathological dysfunction of this substrate as manifest in temporal lobe epilepsy and Alzheimer’s dementia.

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