Electrical Coupling between Interneurons with Different Excitable Properties in the Stratum Lacunosum-Moleculare of the Juvenile CA1 Rat Hippocampus

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Electrical coupling among GABAergic interneurons is believed to play an essential role in shaping synchronized brain network activity related to cognition and behavior. We have studied the rules governing the electrical coupling between hippocampal interneurons located in stratum lacunosum-moleculare of the CA1 hippocampus. The most frequently recorded interneuron subtype had short multipolar dendrites and a dense local axonal arborization, typical of neurogliaform cells. Electrical excitability in this class of interneurons was heterogeneous. Although injection of small current steps evoked late spiking, larger steps triggered different types of firing patterns. Trains of action potentials ranged from clearly adapting to highly irregular, with clustered or mostly regular spikes. Electrotonic and action potentials could be propagated to the coupled cells; the coupling coefficient for electrotonic signals was 0.035, which compared with 0.005 for action potentials. Electrical coupling was reversibly blocked by application of carbenoxolone. Multiple simultaneous recordings indicated that interneurons with similar and different firing patterns were electrically coupled. This visual impression was quantitatively confirmed by principal component analysis applied to variables related to membrane excitability. In fact, the probability of finding electrically coupled neurons in our sample was not dependent on the excitable properties of the cells tested and was $\sim 0.34$. The presence of diffuse electrical coupling among hippocampal interneurons of stratum lacunosum-moleculare with different excitability is a novel finding with important implications. For example, the promiscuity of electrical connections may endow inhibitory networks with a large degree of flexibility and regulate the computational power of the hippocampus during different synchronized states.

Key words: interneuron; GABA; gap junction; connexin; pannexin; hippocampus
interneurons and spiny stellate cells (Venance et al., 2000), between fast-spiking cells and pyramidal neurons (Meyer et al., 2002), and between layer 1 late-spiking and non-late-spiking cells (Chu et al., 2003) may occur, albeit much less frequently. Recent work by Simon et al. (2005) in the neocortex has suggested that the electrical coupling of neurogliaform interneurons is very promiscuous and may act as a bridge between different types of interneurons. However, in the hippocampus, no promiscuous electrical coupling of neurogliaform interneurons was reported by Price et al. (2005), despite the clear observation of promiscuous chemical connectivity.

Here, we have addressed this issue by studying the electrical coupling of hippocampal interneurons defined by their firing pattern properties. Our results indicate that both electrically homogenous and heterogeneous interneurons of stratum lacunosum-moleculare can form gap-junction-mediated networks.

Materials and Methods

Slice preparation. The procedure used to obtain slices is similar to the one described by Aradi and Maccaferri (2004). Briefly, 12- to 21-d-old Sprague Dawley rats were deeply anesthetized with isoflurane and decapitated in accordance with National Institutes of Health and institutional protocols. The brain was removed quickly, and slices of 300–400 μm thickness were cut using a vibrating microtome (Leica, Nussloch, Germany) and placed into ice-cold oxygenated solution, which contained the following (in mM): 2.5 KCl, 1.25 NaH2PO4, 28 NaHCO3, 0.5 CaCl2, 7 MgSO4, 7 glucose, 234 sucrose, 1 ascorbic acid, and 3 pyruvic acid. Slices were then incubated in oxygenated artificial CSF (ACSF) in a holding chamber at a temperature of 32–34°C for 30 min and then stored at room temperature until recording. The recording temperature was 32–34°C. The ACSF contained 130 mM NaCl, 3.5 mM KCl, 1.25 mM NaH2PO4, 1 mM MgSO4, 1.5 mM CaCl2, 24 mM NaHCO3, and 10 mM glucose and was saturated with a mixture of 95% O2 and 5% CO2 at pH 7.4. Interneurons were identified visually using an upright microscope (Olympus Optical, Tokyo, Japan) equipped with a 40X differential interference contrast (DIC) objective coupled to an infrared (IR) video camera system (Dage-MTI, Michigan City, IN). Neurons were selected in stratum lacunosum-moleculare, preferably close to the hippocampal fissure.

Electrophysiological recordings. Simultaneous double or triple recordings were performed in current-clamp mode, using two identical Multiclamp 700A amplifiers and the pClamp9 program package (Axon Instruments, Union City, CA). Patch electrodes were filled with an internal solution containing 125 mM K-methyl-
sulfate, 4 mM NaCl, 4 mM MgATP, 0.3 mM NaGTP, 16 mM KHCO₃, and 0.5% biocytin, saturated with 95% O₂ and 5% CO₂ to a pH of 7.3 and 285–295 mOsm. The junction potential associated with this solution was experimentally estimated (Neher, 1992) to be −8.5 mV and was not corrected. When filled with this solution, the resistance of the electrode was −5 MΩ. To reduce electrode capacitance, the tip of the pipettes was coated with beeswax (Fisher Scientific, Pittsburgh, PA). In addition, electrode capacitance was compensated through the amplifier. Membrane resting potential was estimated from voltage-clamp recordings immediately after breakthrough. Following Ohm’s law $V = IR$, the holding current recorded at −60 mV was used to calculate the resting potentials of the cells. Input resistance was estimated by a 10 mV (250 ms duration) negative voltage step. The advantage of this method is that it allowed the determination of a “resting potential” also from cells that were spontaneously active. Then, after a 10–20 pA current clamp by applying 10–20 pA current steps. We tested for the presence of gap junctions by repetitively applying a 645 ms current step of −100 pA (≥30 sweeps). After averaging, the coupling coefficient for current injection [direct current (DC) coupling] was calculated as the ratio of voltage deflection in the noninjected cell to the voltage deflection in the injected cell measured at the end of the stimulus. Cells were considered coupled if they had a DC coupling value higher than 0.001 and the shape of the electrotonic response was recognizable in the recording from the noninjected cell. To facilitate the recognition of such shape in cells with weaker coupling, a step of at least +100 pA was used. Spike coupling coefficients were calculated similarly.

**Chemicals.** 3β-Hydroxy-11-oxoolean-12-en-30-oic acid 3-hemisuccinate (carbenoxolone) was obtained from Sigma (St. Louis, MO) as a disodium salt and was dissolved directly into the ACSF to a final concentration of 100 μM. Occasionally, brief sonication (1–2 min) was used to facilitate the solution of the drug.

**Data analysis.** Data were filtered at 3 kHz and digitized at 20 kHz using a Digidata 1322A analog-to-digital board. Analysis was performed using pClamp (Axon Instruments), Origin (Microcal, Northampton, MA), Excel (Microsoft, Seattle, WA), Whole-Cell Program (courtesy of Dr. J. Dempster, University of Strathclyde, Glasgow, UK), MVSP (Kovach Computing Services, Pentraeth, Isle of Anglesey, UK), and Prism software (GraphPad, San Diego, CA) packages.

**Principal component analysis.** We applied principal component analysis (PCA) (Joliffe, 1986) to five variables related to membrane excitability measured during firing evoked by a 80 pA suprathreshold, 645-ms-long current step. AHP timing was measured at threshold current levels. The latency between the action potential and afterhyperpolarization peak (AP) to the modulation of spike amplitudes, and the latency between the action potential and afterhyperpolarization peaks (AHP timing) to the shape of the AHP. The first four variables were measured during firing evoked by a 80 pA suprathreshold, 645-ms-long current step. AHP timing was measured at threshold current levels. The different variables were transformed in Z-scores to account for the different units of measurement. As a result of PCA, we were able to plot individual cases in the three-dimensional space. The MVSP software (Kovach Computing Services) scaled the component loadings to unity so that the sum squares of an eigenvector equaled 1, and the component scores were scaled so that the sum of squares equaled the eigenvalue of the corresponding component. We calculated the eigenvectors that accounted for 95% of the variance.

**Figure 3.** Propagation of DC signals via electrical coupling. **A,** Left, DIC-IR image of two interneurons. Right, Simultaneous double current-clamp recording proving an electrical connection. $V_1$ and $V_2$ mark the voltage traces recorded from the two different neurons, and $I_1$ and $I_2$ show the timing of injection of −100 pA current steps in the two different cells, respectively. The insets show the voltage responses at a magnified scale. **B,** Left, Distribution plot of the DC coupling coefficients for 73 pairs of cells tested bidirectionally. Right, Summary graph of the ratio of the DC coupling measured bidirectionally in the same pairs. Notice the symmetrical distribution centered on 1.

**Figure 4.** Electrical coupling is eliminated by carbenoxolone (100 μM). **A,** The voltage response to the injection of a −100 pA current step was monitored in the presynaptic ($V_{1p}$) and postsynaptic ($V_{2p}$) cell in control conditions (1) and during the application (2) and washout (3) of the drug. Traces are the averages of 50 sweeps; recordings from the presynaptic and postsynaptic neuron are marked as $V_1$ and $V_2$, respectively. The time course of the voltage response from the presynaptic ($V_{1p}$) and postsynaptic ($V_{2p}$) neurons are shown in the top and bottom scatter plots, respectively. The black bar indicates drug application. **B,** Summary plot of the effect of carbenoxolone on the normalized DC coupling coefficient (DC coupling) in n = 8 experiments. Notice that the effect was partially reversible.
value. Euclidean distances between pairs of points (Ed) were calculated using an extension of the Pythagoras theorem following the formula $Ed = \sqrt{(\Delta x)^2 + (\Delta y)^2 + (\Delta z)^2}$, where $x$, $y$, and $z$ are the three principal components.

Data are expressed as mean ± SEM. Parametric and nonparametric tests were used, as detailed in the text, as required.

Visualization of recorded cells and reconstruction. Methods were similar to the ones described by Maccanelli and Dingledine (2002). Briefly, slices were fixed for 1–10 d in a 4% paraformaldehyde PBS solution at 4°C. Endogenous peroxidase activity was removed by incubating the slices in 10% methanol/1% H$_2$O$_2$ PBS solution. Biocytin staining was processed using an avidin-HRP reaction (Vectastain ABC Elite kit; Vector Laboratories, Burlingame, CA), and axon visualization was improved using a PBS solution containing NiNH$_4$SO$_4$ (1%) and CoCl$_2$ (1%). Slices were not resectioned but directly mounted on the slide using an aqueous mounting medium (Vectorshield; Vector Laboratories). Alternatively, they were first dehydrated and then mounted on the slide using a toluene solution (Permount; Fisher Scientific, Fair Lawn, NJ). Slices were observed at 40× magnification and photographed using the AxioVision software package (Zeiss, Oberkochen, Germany).

Results

Heterogeneity of intrinsic excitability of CA1 stratum lacunosum-moleculare interneurons

Forty-five individual single-cell recordings from interneurons located in stratum lacunosum-moleculare of the CA1 hippocampus were used to study their anatomical heterogeneity and intrinsic excitability (Fig. 1). The most common cell type encountered ($n = 11$) had short multipolar dendrites, mostly restricted to stratum lacunosum-moleculare, and was approximately centered on a dense local axonal arborization (Fig. 1A–D). Thirteen additional interneurons with a similar dendritic structure were recovered, but the axon could not be visualized. In 12 more cases, interneurons with long dendrites spanning through stratum lacunosum-moleculare, stratum radiatum, and occasionally reaching stratum pyramidale were recovered (Fig. 1E,F). However, their dendritic structure was quite variable and much less stereotyped than the previous cell group. The axon was either not visible or was cut at the surface of the slice before it could branch profusely. Therefore, a precise identification of these cells was not possible. It is very likely that many interneuron subtypes were variably represented in this cell group. Indeed, the dendrites of five cells were not obviously organized along a preferential axis, whereas in five cells, the predominant axis along which the dendrites was oriented was approximately parallel to stratum pyramidale; in the remaining two cases, it was perpendicular to it. The typical axonal branching mostly restricted to stratum radiatum allowed the identification of an additional cell as a Schaffer-associated interneuron (Vida et al., 1998; Cope et al., 2002). In four more cases, the axon was the structure more clearly recovered and was located mostly within stratum lacunosum-moleculare. Last, in four neurons, not enough structures were recovered for any type of meaningful description.

Given that different cell types are present in stratum lacunosum-moleculare (Williams et al., 1994; Vida et al., 1998), it was not surprising that a large variety of firing patterns could be observed in response to suprathreshold depolarizing current injections via the recording pipette. Cells were held at approximately −67 mV, close to their average resting potentials (see Materials and Methods). However, heterogeneity of the firing patterns was not only observed across interneurons with different anatomical properties but also within interneurons with similar dendritic and axonal distributions. For example, four interneurons with a similar morphology, but different firing patterns, are shown in Figure 1A–D. Trains of action potentials could range from clearly adapting to highly irregular with clustered or mostly regular spikes. The amplitude of the spike could remain mostly constant or be subjected to strong modulation during the train (Fig. 1, compare A, B). These cells were typically recovered close to the border with the dentate gyrus. The short multipolar dendritic architecture and dense local axon distribution is very similar to the previously described neuroglialform interneurons in the hippocampus and neocortex (Kawaguchi, 1995; Hestrin and Armstrong, 1996; Vida et al., 1998; Tamas et al., 2003). Two examples of the firing patterns of cells with longer dendrites are shown in Figure 1, E and F. In these cases, the axon reached a considerable distance before branching into collaterals and being cut off at the surface of the slice, preventing a proper classification. In conclusion, different firing patterns can be recorded in stratum lacunosum-moleculare interneurons. Also, similar excitatory properties are not necessarily associated with a specific cell type defined by its axonal and dendritic morphology.

Next, we tried to solidify these initial observations into a quantitative result by measuring parameters that are related to firing patterns such as (1) mean ISI, (2) CV ISI, (3) ratio ISI, (4)
Electrical coupling between interneurons and propagation of electrotonic responses and action potentials

Next, we tested for the presence of electrotonic coupling in pairs and triplets of simultaneously recorded cells (Fig. 3). We quantified the strength of electrotonic coupling by injecting a −100 pA current pulse in one cell of the pair (or triplet), as shown in Figure 3A, and by calculating the ratio of the voltage response recorded in the other neuron(s), which was not subjected to any pulse injection (the distribution of the DC coupling obtained from averaged sweeps is shown in Fig. 3B). The average DC coupling calculated from 146 connections was 0.035 ± 0.002. In all cases, coupling was found to be bidirectional, and the distribution of the ratio of the DC coupling in the two opposite directions was centered on 1. The mean ratio was 1.2 ± 0.1 (n = 73).

Application of the gap-junction uncoupler carbamoloxone at 100 μM (Fig. 4) reduced the DC coupling to 3.1 ± 1.5% (p < 0.05, paired t test) of control in 13 connections tested in separate paired recordings. In eight of such cases, after the DC coupling coefficient decreased to 1.5 ± 2.5% of control in the presence of carbamoloxone, it was possible to partially wash out the effect of the drug and restore the DC coupling coefficient to 36.4 ± 8.2% of control (p < 0.05, repeated-measures ANOVA with Bonferroni’s post hoc test). Thus, our results indicate that electrotonic coupling is mediated by gap junctions.

The propagation of electrical signals via gap junctions depends on the frequency components of the presynaptic electrical waveform (Galarreta and Hestrin, 1999). Therefore, we also measured the postsynaptic response to presynaptic spikes (Fig. 5). The spike-coupling coefficient was defined as the ratio of the amplitude of the averaged postsynaptic spikelet to the averaged presynaptic spike. From a total of 111 electrical connections, the
calculated spike coupling coefficient was $0.005 \pm 0.001$. As shown by the cumulative distributions superimposed in Figure 5B, DC signals propagate with a much higher efficiency when compared with action potentials. However, despite the reduced efficiency, spike coupling calculated from averaged sweeps in individual pairs of neurons was significantly correlated to the level of DC coupling, as shown in Figure 5C ($r^2 = 0.17; p < 0.05$). The average ratio of the spike coupling over the DC coupling coefficient in our sample is shown by the slope of the regression line (0.078). The low $r^2$ coefficient and the scatter of the data are probably the results of spike amplitude modulation during trains of action potentials in particular sets of interneurons. As shown in the example of Figure 5D, although the amplitude and duration of the action potential waveforms could greatly vary during the response to a current step injection, the spikelet amplitudes observed on the postsynaptic cell varied much less. Indeed, the spike-coupling coefficient was modulated along different action potentials in the train, probably reflecting the different frequency components of the presynaptic signals. Another possibility that may explain the low $r^2$ coefficient is that different type of interneurons may express specific dendritic voltage-dependent conductances, which may differentially affect action potential propagation to the gap-junction site(s). In conclusion, these results indicate that interneurons in stratum lacunosum-moleculare are electrically coupled via gap junctions and that various electrical waveforms can be propagated across the connected neurons.

Given the heterogeneity of electrical excitability in stratum lacunosum-moleculare interneurons, we next analyzed the firing pattern of coupled cells to test the hypothesis that interneurons with different intrinsic excitability could be electrically coupled.

**Electrical coupling between interneurons with similar and different intrinsic excitability**

When we compared the firing pattern of electrically coupled neurons, we found evidence for coupling between interneurons with similar and different firing patterns. Figure 6 shows two examples of electrical connectivity in the case of interneurons displaying mostly regular or highly irregular and “stuttering” firing patterns. The examples selected had a strong DC coefficient coupling, and spikelets were clearly visible in single sweeps. Although these recordings provide direct evidence that electrically similar interneurons of hippocampal CA1 stratum lacunosum-moleculare are coupled, we also found evidence for coupling between neurons with different excitability properties. Figure 7 shows an example of a simultaneous recording between two morphologically similar interneurons with different firing patterns. This connection was also strong, and spikelets could be clearly observed in single sweeps. Additional experiments taking advantage of simultaneous recordings from three neurons allowed us to test electrical couplings in multiple combinations. Figure 8 shows two examples in which interneurons with different firing patterns were variably interconnected.

Although visual inspection of the firing pattern of connected cells clearly indicated that electrically different interneurons were often coupled, we reasoned that it was crucial to objectively quantify this degree of connectivity as a function of the relative diversity of the excitable properties of the coupled neurons.

To address this point, we performed PCA on the five variables (mean ISI, CV ISI, ratio ISI, ratio AP, and AHP timing) measured from the firing pattern responses of 323 neurons that had been tested for electrical coupling in double or triple recordings (Fig. 9A, B; factor loadings are shown in C). The first three factors accounted for 74.7% of the total variance (eigen values: factor 1, 1.640; factor 2, 1.137; factor 3, 0.959; and cumulative percentage: factor 1, 32.8%; factor 2, 55.5%; factor 3, 74.7%). We decided to examine the degree of dissimilarity between the electrical properties of the neurons of the database as Euclidean distances and to relate the measured distances to the presence or absence of electrical coupling. The average Euclidean distance between cells that were not electrically coupled was $0.103 \pm 0.014 (n = 137$ connection tested), which compared with a mean value of $0.100 \pm 0.006 (n = 71; p > 0.05$, Mann–Whitney test) in neurons with verified electrical connections. However, the mean value of the Euclidean distance does not give information about how this measurement was distributed. We considered the possibility that, despite the similar mean value, a difference in the shape of the distribution of the Euclidean distances could affect the probability of finding connections because of the different occurrence of specific a
range of values in the two datasets. However, when we compared the two cumulative distributions, they were not statistically different (Kolmogorov Smirnoff test, \( p > 0.05 \)) (Fig. 9D).

In conclusion, the results of this analysis indicate that electrical coupling in our sample did not depend on the relative similarity of the excitable properties of the pair tested. We further quantified the probability of finding a connection as a function of the Euclidean distance of the tested pairs. The probability was calculated for Euclidean distances binned as shown in Figure 10. As expected, no correlation was found (\( F \) test, \( p > 0.05 \)).

In conclusion, the probability of finding an electrically coupled pair between two stratum lacunosum-moleculare interneurons in our sample, independently of their firing pattern, was relatively high, resulting in a value of 0.34.

Discussion

The main finding of this work is that hippocampal interneurons with different intrinsic excitability are electrically coupled. Our conclusion is based on electrophysiological and pharmacological evidence acquired from single and multiple recordings. Overall, our results indicate that the organizing principles based on the similar-to-similar coupling rule described in many inhibitory networks are not followed by every class of interneuron in the hippocampus.

Interneurons and stratum lacunosum-moleculare network

In the CA1 subfield of the hippocampus, stratum lacunosum-moleculare receives excitatory input from several extra hippocampal regions, including the nucleus reuniens of the thalamus (Wouterlood et al., 1990), the inferotemporal cortex (Yukie and Iwai, 1988), and the direct input originating from layer III of the entorhinal cortex [temporoammonic (TA) pathway] (Steward and Scoville, 1976). Of these three sources of information, the TA projection is thought to be critical for the bidirectional modulation of the Schaffer collateral input leading to spiking activity and synaptic plasticity (Remondes and Schuman, 2002) in CA1 pyramidal cells, which are the output neurons of the hippocampus. These cellular and network basic mechanisms are likely to be involved in the consolidation of spatial long-term memory, which is impaired after lesion of the TA pathway (Remondes and Schuman, 2004).

Several studies have also shown that inhibitory interneurons in stratum lacunosum-moleculare are synaptically activated in a feedforward manner by the TA input or the Schaffer collateral (Kunkel et al., 1988; Lacaille and Schwartzkroin, 1988; Khazipov et al., 1995; Morin et al., 1996) and release GABA on postsynaptic pyramidal cells, thus generating IPSPs (Lacaille and Schwartzkroin, 1988; Empson and Heinemann, 1995; Vida et al., 1998; Bertrand and Lacaille, 2001). Different types of stratum lacunosum-moleculare interneurons have been shown to target specific postsynaptic domains, which are associated with specific layers receiving excitatory input from different sources (Vida et al., 1998). Nevertheless, despite the potential importance of stratum lacunosum-moleculare interneurons for the regulation of a circuit involved in higher cognitive functions, some essential data on their network organization are still missing. In particular, little is known, at present, about their electrical coupling.

Electrical heterogeneity of stratum lacunosum-moleculare interneurons

When we examined the anatomy of the recorded interneurons, we realized that the firing pattern was different not only across different anatomical types but also within a group of small multipolar interneurons displaying a stereotyped morphology. In particular, the morphological appearance of these latter interneurons was consistent with the previously reported description of neurogliaform cells. The firing pattern of the “late-spiking”
the contrary, the occasional stuttering firing pattern observed by our recordings illustrated in Figures 6Kawaguchi (1995) and Tamas et al. (2003) were very similar to neurogliaform cells of neocortical layer II/III reported by Vida et al. (1998). Our results are also consistent with recent work in the primate layer II/III prefrontal cortex showing distinct firing patterns in anatomically identified neurogliaform interneurons (Krimmer et al., 2005).

In conclusion, although some classes of interneurons are known to be associated with a stereotypical firing pattern (Maccari and Lacaille, 2003), this may not be a universal case, as reported previously by others (Mott et al., 1997; Parra et al., 1998; Gupta et al., 2000).

Coupling between electrically heterogeneous interneurons

In the hippocampus, immunocytochemical and ultrastructural studies have long suggested the existence of gap junctions between basket cells and interneurons with horizontal dendrites in stratum oriens/aluves (Kosaka and Hama, 1985; Yamamoto et al., 1989; Fukuda and Kosaka, 2000). Recent electrophysiological work has directly demonstrated electrotonic coupling among the broad class of horizontal stratum oriens interneurons (Zhang et al., 2004). However, many different cell types with a variety of physiological properties are included under this denomination (Maccari and Lacaille, 2000; Maccari, 2005), and the relationship between cellular excitability and connectivity was not explored in that study. Thus, the question whether only interneurons with similar excitable membranes could be coupled was left unresolved. A recent report by Price et al. (2005) has provided evidence for homologous electric coupling between anatomically identified neurogliaform cells in the CA1 stratum lacunosum-moleculare. However, the degree of similarity of the firing pattern of the connected neurons was not studied in detail.

This work is the first demonstration of an electrically coupled network in the hippocampus that is also formed by elements with different excitability. Electrical coupling between cells with similar intrinsic excitability has been clearly shown to facilitate the synchronization of inhibitory networks (Garrella and Hestrin, 1999; Gibson et al., 1999; Beierlein et al., 2002). On the contrary, the consequences of gap-junction-mediated coupling between neurons with different intrinsic excitability are complex to predict and may yield different results depending on the strength of the connection (Perez-Velazquez et al., 2001). Theoretical studies have proposed that electric coupling between neurons with different intrinsic properties can lead to antiphase or asynchronous firing (Sherman and Rinzel, 1992; Chow and Kopell, 2000; Lewis and Rinzel, 2003; Pfeuty et al., 2003).

When the anatomy of the recorded cells was examined in

**Figure 9.** PCA analysis of excitability in electrically coupled versus uncoupled stratum lacunosum-moleculare interneurons. A, B, Three-dimensional summary plot showing the position along the three principal components of all interneurons tested for electrical connectivity (black symbols). The connections tested are represented by the lines between the symbols. To facilitate the visibility of most points, the scale excludes the two furthest points of the dataset (see lines exiting the borders of the plots). Combinations that were found electrically uncoupled are shown by the lines in A, whereas electrophysiologically verified combinations of coupled cells are represented by the lines in B. C, Factor loadings (correlations) for each variable are shown in histogram form (white, factor 1; gray, factor 2; and black, factor 3). D, The cumulative distributions of the Euclidean distances in the cases of unconnected (u) and connected (c) neurons are superimposed. Notice the similarity between the two curves.
relation to their firing pattern in individual recordings, neuroglial-like interneurons were the largest proportion of the cells recovered that could be identified as a consistent anatomical group. Therefore, it is likely that this cell type was also widely represented in multiple recordings from coupled cells that were not anatomically recovered. This observation suggests that neuroglialform cells with different excitable properties form a complex inhibitory network in the hippocampus. If the electrical coupling of neuroglialform cells with different excitable properties led to antiphase or asynchronous firing during oscillatory activity, then it could play an important role in ensuring tonic rather than phasic GABAergic release. Furthermore, modulation of the junctional conductance between different neuroglialform cell types could provide a large degree of flexibility in the regulation of GABAergic input during oscillations. Last, the possibility that neuroglialform interneurons are also coupled with different anatomical types should not be excluded. In fact, although neuroglialform interneurons were the most represented cell type recognized in our single recordings, a significant fraction of the recovered neurons had a dendritic structure clearly different from neuroglialform cells, and similar proportions of cell types are likely to have occurred during our multiple recordings. This possibility would be entirely consistent with the recent study of Simon et al. (2005) in the neocortex. Highly promiscuous electrical connectivity would have the potential to create a network of neurons exquisitely sensitive to the general level of activity in inhibitory networks. For example, direct regulation of blood vessels by neuroglialform neurons expressing neuronal nitric oxide synthase (NOS) and neuropeptide Y (NPY) has been reported in neocortical circuits (Cauli et al., 2004). Intriguingly, very recent work by Price et al. (2005) has confirmed that a high percentage of neuroglialform cells in the hippocampal stratum lacunosum-moleculare express the mRNA for both NOS and NPY. Therefore, diffuse electrical coupling might be an essential feature to sense the general level of activity in neuronal networks and translate it into a signal that can be used for the fine regulation of blood flow.

**Chemical versus electrical coupling of stratum lacunosum-moleculare interneurons**

Work by several groups has shown that electrical coupling in interneurons is often associated with chemical synaptic coupling (Galarreta and Hestrin, 1999; Gibson et al., 1999; Bartos et al., 2001; Blatow et al., 2003; Simon et al., 2005). We did not specifically test for this possibility. Our recording conditions were not optimized to detect GABAergic unitary IPSPs because we recorded cells at membrane potentials associated with a small driving force for GABA receptor-mediated events. Another potential explanation is that we did not test connections at very low frequency, which has been reported to be an essential experimental condition to trigger a postsynaptic response originating from neuroglialform interneurons (Tamas et al., 2003). In addition, in several of the recorded cells the axon appeared to be cut at the surface of the slice before it could generate a significant arborization. If this occurred to similar degrees during single and double recordings, then our results would be consistent with a dendritic location of the gap-junction-mediated connection. A final possibility could be the tonic suppression of GABA release by a presynaptic modulator. For example, work by Losonczy et al. (2003, 2004) has clearly shown that tonic activation of presynaptic receptors can strongly reduce release from GABAergic glutamatergic terminals in hippocampal slices. Price et al. (2005) have indeed shown recently that tonic GABA$_B$ receptor activity modulates GABAergic release from neuroglialform cells. Therefore, our lack of evidence for chemical connections remains inconclusive.

In summary, our results highlight a novel pattern of organization for electrically coupled networks in the hippocampus, which suggests that inhibitory circuits may be more complex than previously thought. This pattern of organization may provide an additional point of regulation for learning and memory.

**References**


