Factors Underlying Bursting Behavior in a Network of Cultured Hippocampal Neurons Exposed to Zero Magnesium

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Submitted 5 June 2003; accepted in final form 25 September 2003

Mangan, Patrick S. and Jaideep Kapur. Factors underlying bursting behavior in a network of cultured hippocampal neurons exposed to zero magnesium. J Neurophysiol 91: 946–957, 2004. First published October 8, 2003; 10.1152/jn.00547.2003. Factors contributing to reduced magnesium-induced neuronal action potential bursting were investigated in primary hippocampal cell culture at high and low culture density. In nominally zero external magnesium medium, pyramidal neurons from high-density cultures produced recurrent spontaneous action potential bursts superimposed on prolonged depolarizations. These bursts were partially attenuated by the NMDA receptor antagonist d-APV. Pharmacological analysis of miniature excitatory postsynaptic currents (EPSCs) revealed 2 components: one sensitive to d-APV and another to the AMPA receptor antagonist DNQX. The components were kinetically distinct. Participation of NMDA receptors in reduced magnesium-induced synaptic events was supported by the localization of the NR1 subunit of the NMDA receptor with the presynaptic vesicular protein synaptophysin. Presynaptically, zero magnesium induced a significant increase in EPSC frequency likely attributable to increased neuronal hyperexcitability induced by reduced membrane surface charge screening. Mean quantal content was significantly increased in zero magnesium. Cells from low-density cultures did not exhibit action potential bursting in zero magnesium but did show increased EPSC frequency. Low-density neurons had less synaptophysin immunofluorescence and fewer active synapses as determined by FM1-43 analysis. These results demonstrate that multiple factors are involved in network bursting. Increased probability of transmitter release presynaptically, enhanced NMDA receptor-mediated excitability postsynaptically, and extent of neuronal interconnectivity contribute to initiation and maintenance of elevated network excitability.

INTRODUCTION

Fast excitatory neurotransmission in the hippocampus is mediated primarily by glutamate acting on 3 types of ionotropic receptors named for their preferred agonists: kainate, α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA), and N-methyl-D-aspartate (NMDA) (Dingledine et al. 1990). All 3 types of receptor incorporate ion channels which, when activated, become cation permeable. Excitatory glutamatergic synaptic interactions are maintained in in vitro preparations of hippocampal slice (Parfitt and Madison 1993) and pyramidal cell culture. Hippocampal neuronal cultures establish dense neuritic networks with multiple synaptic contacts (Banker and Cowan 1977; Fletcher et al. 1991; Hoch and Dingledine 1986; Muller and Seifert 1982). AMPA and NMDA receptors are increasingly expressed and localized to synaptic sites with time (Bekkers and Stevens 1989; Rao et al. 1998). These anatomic studies have been bolstered by physiological findings of functionally intact excitatory synaptic interactions (Cummings et al. 1996; Wilcox et al. 1994).

Over the course of several years, study of physiological and pathophysiological aspects of glutamatergic neurotransmission has led to the development of techniques by which the excitability level of synaptic interactions can be modulated. One such technique involves the manipulation of divalent ions. The role of calcium in exocytotic neurotransmitter release has been well documented (see Poage and Meriney 2002 for recent review). Reduced extracellular calcium has been used for many years as a standard tool for inhibiting synaptic transmission. Magnesium also has marked effects on synaptic interactions. Dodge and Rahamimoff (1967) first demonstrated that quantal release at the frog neuromuscular junction decreased steeply as Mg\(^{2+}\) concentration increased. In the hippocampus, elevated magnesium has been found to attenuate or abolish the prolonged neuronal depolarizations and synchronized action potential bursts emblematic of epileptiform activity (Schwartzkroin and Prince 1978). Conversely, reducing magnesium below physiological levels induces enhanced excitability in both hippocampal slice (Walther et al. 1986) and cultures (Sombati and DeLorenzo 1995).

The mechanism of reduced magnesium enhancement of excitatory neurotransmission has been presumed to be related to the ions’ physiological blockade of NMDA receptor channels. It has long been recognized that physiological magnesium provides a voltage-dependent steric blockade of the cation permeable pore of NMDA receptors and thus renders them largely inactive at normal membrane potentials (Coan and Collingridge 1985; Collingridge et al. 1983; Hablitz and Langmoen 1986; Mayer and Westbrook 1984). Several studies have demonstrated that synchronized spontaneous neuronal activity induced by eliminating extracellular magnesium is reduced or abolished by NMDA receptor antagonists (Albowitz et al. 1997; Coan and Collingridge 1987; Collingridge et al. 1988; Gulyas-Kovács et al. 2002; Gutierrez et al. 1999; Tancredi et al. 1990). These studies suggested that increased excitability induced by magnesium reduction was primarily a postsynaptic phenomenon, caused by increased flux of depolarizing ions, mostly calcium, through NMDA receptor channels.

Other studies have suggested additional excitatory mechanisms associated with extracellular magnesium reduction. Magnesium reduction (vs. removal) induced enhanced excitatory responses in the CA1 region of the hippocampus that were not reversed by NMDA receptor antagonists (Hamon et
al. 1987). A careful study by Mody and colleagues (1987) suggested that reducing magnesium may produce a reduction in the electric field across the neuronal membrane by reduced surface charge screening causing what is, in essence, a membrane depolarization.

However, modeling studies in the hippocampus have suggested that enhanced excitability, although necessary, is not sufficient for the occurrence of bursting in a network of pyramidal neurons. A model network simulation of hippocampal bursting demonstrated that bursting can occur, even in the presence of neuronal inhibition, only if connectivity between neurons in the circuit is adequately dense. Low connectivity prevents the development of bursting (Traub et al. 1984, 1987).

We have used electrophysiological and immunocytochemical techniques to examine the factors affecting low magnesium-induced bursting in networks of cultured hippocampal pyramidal neurons. Specifically, we sought to identify the contributions of presynaptic and postsynaptic processes to enhanced network excitability and to test model-driven assumptions of adequate connectivity required for synchronized bursting.

METHODS

Treatment of animals followed guidelines set by the University of Virginia Health Sciences Center Animal Research Committee. All efforts were made to minimize animal stress and discomfort.

Hippocampal cultures

Neuronal hippocampal/glia cocultures were prepared according to the method of Goslin et al. (1998). Neurons and glia were cultured on separate surfaces then combined to form a tissue-culture “sandwich.” This approach allowed the preparation of relatively low density hippocampal cultures while still allowing access of hippocampal neurons to glia-derived neurotrophic substances. Each segment of the preparation is summarized.

Preparation of coverslips

FisherBrand coverslips (12-545-86-ID; 25 mm) were placed in 10% nitric acid for ≥18 h. Coverslips were then rinsed in 6 changes of distilled water (20 min each rinse) and sterilized with dry heat (225°C for 6 h). After cooling, coverslips were coated with poly-lysine (1 mg/ml in borate buffer), placed in a 37°C incubator overnight, then rinsed twice with sterile distilled water (30 min each rinse). Coverslips were then layered with minimum essential medium (MEM) plus 10% horse serum (HS; see Goslin et al. for detailed constituents) and placed in a 37°C incubator. Fresh MEM/HS was added the day of the coculture procedure (see following text).

Primary glial culture preparation

Gliaal cells were prepared 10 days before coculturing with hippocampal neurons. Before glial isolation, 3 drops of sterile, melted polylysine-coated coverslips were applied at the vertices of an equilateral triangle in several sterile 60-mm tissue-culture dishes. The parafilm provided the spacing necessary to keep the hippocampal neuron-containing coverslips from directly contacting glial cells during coculturing (see following text). Alternatively, spacing was provided by longitudinally cut Teflon O-rings (3 mm width, 22 mm ID; Small Parts, Miami Lakes, FL). In a laminar flow hood, neonatal Sprague-Dawley rat pups were decapitated after being placed on ice for 2–3 min. Brains were removed into cold HEPES-buffered Hank’s balanced salt solution (HEPES/HBSS). Cerebral hemispheres were isolated and the meninges removed. Excess HEPES/HBSS was removed and the tissue chopped as finely as possible with scissors. The tissue was placed in 12 ml fresh HEPES/HBSS to which 1.5 ml 2.5% trypsin and 1.5 ml DNase (1%) were added. The tissue was incubated at 37°C for 15 min with continuous slow speed stirring. The supernatant was passed through sterile nylon mesh (212 μm) and diluted with an equal volume of 10% HS in MEM. Cell suspension was centrifuged at 800–1,000 rpm for 5 min. Pellets were resuspended in 10 ml MEM/10% HS. Cells were counted with a hemocytometer. The yield was generally 6 × 10^6·1 × 10^7 cells per brain. Cells were diluted to approximately 3,000–5,000 cells/ml in 10% HS/MEM. 3 ml was placed in each of several 60-mm tissue-culture dishes. Medium was completely replaced the day after plating; thereafter, medium was replaced every 4 days with 10% HS/MEM.

Hippocampal neuronal culture preparation

 Cultures were prepared from E-18 Sprague-Dawley rat fetuses. Fetuses were decapitated and brains removed and placed in HEPES/HBSS. Hippocampi were removed under a dissecting microscope and collected in a small petri dish in HEPES/HBSS. Hippocampi from a single litter were placed in a 15-ml centrifuge tube; HEPES/HBSS was aspirated off and replaced with 5 ml of 0.25% trypsin. The preparation was incubated at 37°C for 15 min. Trypsin solution was replaced with 5 ml HEPES/HBSS. Rinsing with HEPES/HBSS was repeated twice more at 5-min intervals. Hippocampi were triturated until no fragments of tissue remained. Cell density was determined with a hemocytometer and was usually approximately 5 × 10^5 cells/hippocampus. Two densities of hippocampal neurons were prepared. For high-density cultures, 100,000 cells were added to each of the dishes containing the polylysine-coated coverslips prepared previously; for low-density cultures, 10,000 cells were added. After 3–4 h, coverslips were transferred to dishes containing glial cell monolayers in serum-free MEM with N2 supplement. Coverslips were turned so that hippocampal neurons faced glial cells and placed on parafilm or Teflon spacers; spacing was about 2 mm between the cell layers. N2 supplement (1 ml) was added every 10 days.

Electrophysiology

Patch electrodes were pulled from 1.5 mm (OD) × 1.1 mm (ID) borosilicate micropipettes on a horizontal Flaming-Brown micropipette puller (model P-97, Sutter Instruments) using a 2-stage pull protocol. Electrode resistances were 5–8 MΩ. Electrode tips were filled with an internal stock recording solution consisted of (in mM) CsCl 153.3, MgCl2 1.0, HEPES 10.0, and ethylene glycol-bis(β-aminoethyl ether) N,N,N′,N′-tetraacetic acid (EGTA) 5.0, pH 7.3 (with sterile-filtered CsOH), osmolarity 275–280 mOsm. Internal solution was sterile filtered before use. The electrode shank solution contained an ATP regeneration consisting of 3 mM ATP (disodium salt), 19 mM phosphocreatine, and 50 μM creatine phosphokinase.

Before electrophysiological experimentation, coverslips containing hippocampal neurons were removed from culture medium and placed in a 30 × 10-mm polystyrene culture dish containing external recording consisting of NaCl (142), CaCl2 (1.0), CsCl (8.1), MgCl2 (2.1), glucose (10.0), and HEPES (10.0), pH 7.4 (NaOH). The osmolarity was adjusted to 320–325 mOsm with sucrose. External recording solution was sterile filtered before use. For nominally zero magnesium experiments, MgCl2 was replaced by an osmotically equivalent concentration of NaCl. Experiments in hippocampal slices have implicated elevated external cesium in the induction of spontaneous recurrent bursting activity in CA3 and dentate gyrus but only in the presence of the calcium channel blocker cadmium (Xiong and Stringer 2001). To determine whether the elevated cesium in our external solution contributed to neuronal bursting in culture and to replicate an external solution used previously by Sombati and De Lorenzo (1995) in examining the electrophysiological activity of cultured neurons exposed to zero magnesium, KCl (2.5 mM) replaced...
CsCl in some experiments. NaCl was increased to 145 mM in these experiments. We observed no difference in the frequency of recurrent bursting in these 2 external solutions; results obtained under the 2 conditions were combined. Cultured neurons were viewed on the stage of an inverted Nikon microscope. Microelectrodes were interfaced with the amplifier by an Axon Instruments CV-4 headstage and maneuvered into recording position with a PCS 5000 series micro-manipulator (Burleigh Instruments, Fishers, NY).

Whole cell recordings were made at room temperature with an Axopatch 1-D patch clamp amplifier (Axon Instruments, Union City, CA) and low-pass filtered at 2 kHz with an 8-pole Bessel filter before digitization. Data were recorded to a personal computer with Axoscope 7.0 data acquisition software using a Digidata 1200 interface (Axon Instruments). Voltage-clamp recordings of excitatory postsynaptic currents (EPSCs) were digitized at 10–20 kHz; current-clamp recordings of neuronal membrane potential were digitized at 20 kHz. EPSCs were analyzed using MiniAnalysis software (Synaptosoft, Decatur, GA).

**Immunocytochemistry**

Coverslips used for NMDA receptor subunit NR1, AMPA receptor subunit GluR1, or synaptophysin immunostaining were fixed with 4% paraformaldehyde (PFA)/4% sucrose in phosphate-buffered saline (PBS) for 20–30 min at room temperature. Rao and Craig (1998) reported that NR1 receptor subunit staining required methanol fixation; however, we were able to attain satisfactory staining with PFA. Neurons were permeabilized with 0.25% Triton X-100 in PBS for 5 min. Coverslips were washed twice with PBS, blocked with 10% rabbit serum in PBS for 30 min, and exposed to primary antibodies in 3% rabbit serum in PBS.

The mouse monoclonal antibody 54.1 to NR1 (PharMingen, San Diego, CA) was used at a concentration of 2 μg/ml and rabbit polyclonal antibody to GluR1 (Chemicon International, Temecula, CA) was used at 1 μg/ml. Either mouse anti-synaptophysin (Chemicon; for use in double-labeling experiments with GluR1) or rabbit anti-synaptophysin (Zymed Laboratories, San Francisco, CA; for use with NR1) was used at a concentration of 2.5 μg/ml.

Primary antibodies were visualized with fluorochrome-conjugated secondary antibodies (3–4 μg/ml, Molecular Probes, Eugene, OR). The fluorochromes used were Alexa Fluor 488 (GluR1) and Alexa Fluor 594 (NR1). Synaptophysin primary antibody was labeled with secondary antibodies conjugated to either Alexa Fluor 488 or Alexa Fluor 594. Coverslips were mounted in glycerol. Fluorescent images were captured on a Nikon inverted Eclipse TE 200 microscope equipped with a Photometrics CoolSnap cf digital camera using a 40× oil immersion objective (1.0 numerical aperture). Images were captured using Metamorph imaging software and prepared for printing with AdobePhotoshop 6.0.

For whole cell visualization, neurons were first filled with Biotin (Vector Laboratories; 0.5 mg/ml in internal pipette solution), then incubated with streptavidin conjugated with Alexa Fluor 350 (Molecular Probes).

**FM1-43 visualization**

Cells were loaded by exposure to 10 mM potassium external solution (osmolarity maintained by concomitant decrease in sodium concentration) with 8 μM FM1-43 for 10 min. Cells were then washed 3–4 times with normal external solution. For destaining and visualization, cells were exposed to nominally zero-magnesium external.

**Data analysis**

All data are presented as means ± SE unless otherwise noted in the text.

**RESULTS**

**Neuronal properties**

Morphologically identified pyramidal-shaped neurons in culture for 13–17 days were studied by the whole cell patch-clamp technique. In initial experiments, pyramidal cells and their processes were filled with biocytin during electrophysiological recording and visualized with fluorescent dye after fixation. Large- and small-diameter dendrites emerged from the apex and base of pyramidal neurons and branched into secondary and tertiary dendrites (Fig. 1). The complex dendritic arbor of pyramidal neurons extended several hundreds of microns from the cell soma. Previous studies have shown that nonpyramidal interneurons are morphologically distinct bipolar or multipolar neurons whose cell soma stains intensely with the γ-aminobutyric acid synthetic enzyme glutamic acid decarboxylase (GAD; Esclapez and Houser 1999). Passive and active membrane properties of these pyramidal neurons were examined in current clamp mode in magnesium-containing external medium to establish that neuronal properties conformed to those for healthy, viable cells. Mean resting membrane potential was −58.3 ± 2.4 mV (n = 23), in good agreement with previous studies on cultured pyramidal neurons (Coulter et al. 1992).

Neuronal input resistance was measured in a sampling of cells (n = 7) by injecting 2-s current pulses (−1.0 to 0.3 nA in 0.1-nA increments). R_in was given by the slope of the plot of current versus voltage change (Mangan and Bertram 1998). The R_in was 71.3 ± 4.2 MS, higher than that recorded in pyramidal neurons with sharp microelectrodes in our previous studies (Mangan and Bertram 1998; Mangan and Lothman 1996; Rempe et al. 1995), but in accord with other studies using patch electrodes (Spruston and Johnston 1992; Spruston et al. 1994). Action potential (AP) characteristics were also determined as an indication of active, voltage-dependent membrane function. Membrane potential was varied by current injection. Threshold for AP firing was −59.4 ± 2.2 mV (n = 24). Holding the membrane potential at −58 mV by current injection (usually <100 pA) resulted in occasional spontane-
rembrane potential (RMP) (57.6 ± 1002). Membrane potential was held at −58 mV by DC current injection. B: recording from a neuron 15 days in culture. Magnesium in the external solution was replaced by equiosmolar Na+. Membrane potential was held at −58 mV. C: pharmacology of pyramidal cell bursting in 0 [Mg2+]o. Top trace: recording from a pyramidal cell 14 days in culture exposed to 0 [Mg2+]o. Middle trace: recording from the same cell 25 min after addition of 40 μM D-APV, an NMDA receptor antagonist. Bursting activity is attenuated but not abolished. Bottom trace: neuronal excitatory activity ceases 15 min after further addition of 20 μM DNQX, an AMPA receptor antagonist. Recordings in A, B, and C are from different cells.

ous overshooting APs with amplitude 77.2 ± 3.4 mV. These AP properties were in accordance with previous findings (Rempe et al. 1995). Rarely (4 of 36 neurons tested), depolarization (to about −50 mV) resulted in prolonged action potential bursts. In 3/4 cells, these bursts were not recurrent; in one cell bursting continued with an irregular interburst interval for the duration of the depolarization (about 3 min).

In nominally Mg2+-free external medium (0 [Mg2+]o), most cells exhibited recurrent sustained bursting consisting of multiple (≥3) APs with interspike intervals of <200 ms superimposed on a prolonged (500–2,000 ms) depolarization (Fig. 2B). Two pieces of evidence suggested that spike bursting was network-driven rather than an endogenous neuronal property. First, similar activity was absent from cells in low-density cultures (see Fig. 7 below). Second, hyperpolarization to as much as −90 mV by DC current injection often did not terminate bursting. Bursting occurred in 38/43 cells examined and was sustained for the length of the recording (≥20 min). Exposure to 0 [Mg2+]o did not significantly alter resting membrane potential (RMP) (−57.6 ± 3.1 mV; a period of 2–10 s occurred between bursts in all cells; the mean RMP was calculated from this interburst interval), Rm (67.7 ± 3.3 MS; n = 7), or action potential height recorded at a holding potential of −58 mV (79.6 ± 2.1 mV; n = 21). However, the AP firing threshold was shifted significantly to −77.7 ± 4.5 mV after 0 [Mg2+]o exposure. Additionally, treatment resulted in an altered pattern of AP firing. Instead of individually occurring spikes, APs in 0 [Mg2+]o tended to group in discrete repeating epochs (Fig. 2B). Bursting behavior in 0 [Mg2+]o contrasted with action potential firing in Normal Mag where 32/36 cells did not burst (Fig. 2A).

Role of NMDA receptors

Removal of magnesium enhances excitatory activity in neuronal networks by abolishing the voltage-dependent blockade of NMDA-type glutamate receptors (Ascher et al. 1988; Mody et al. 1988; Tancredi et al. 1988; Westbrook 1994). We tested whether 0 [Mg2+]o-induced bursts could be blocked by the NMDA receptor antagonist D-2-amino-5-phosphonovaleric acid (D-APV; Fig. 2C). A neuron 17 DIV exhibited recurrent spontaneous action potential bursts in 0 [Mg2+]o external solution. Addition of D-APV (40 μM) reduced the mean number of action potentials per burst and increased the interburst interval but did not return the cell to the intermittent firing of single action potentials observed in normal magnesium. Only on further addition of the AMPA receptor blocker dinitroquinoxaline (DNQX; 20 μM) was bursting behavior eliminated. This observation was repeated in 14 cells from 8 separate culture preparations (each culture prepared from different embryos). In normal external magnesium, synaptically driven action potential firing was abolished by DNQX alone (data not shown); cells were still capable of spiking if depolarized by DC current injection. These studies demonstrated that APV only partially attenuated 0 [Mg2+]o-induced bursting.

One possible explanation for the failure of D-APV to completely block 0 [Mg2+]o-induced bursts was that NMDA receptors were not present at synapses in cultured hippocampal neurons and these did not participate in excitatory neurotransmission. Alternately, 40 μM D-APV may have been insufficient to block NMDA receptor currents in the 0 [Mg2+]o condition. Finally, it is possible that multiple mechanisms enhance synaptic transmission in 0 [Mg2+]o medium. To evaluate these possibilities, voltage-clamp recordings of spontaneous excitatory postsynaptic currents (sEPSCs) were performed at −50 mV in the presence of 5 μM bicuculline methiodide to block GABAA receptor-mediated events. In some experiments, 1 μM tetrodotoxin (TTX) was used to eliminate spontaneous release of multiple neurotransmitter vesicles and produce single postsynaptic quantal events [i.e., miniature EPSCs (mEPSCs)].

Figure 3 shows an averaged mEPSC consisting of ≥1,000 events from 10 cells under 0 [Mg2+]o conditions. The averaged

FIG. 2. Current clamp recordings from cultured hippocampal pyramidal neurons in normal and nominally zero external magnesium (0 [Mg2+]o). A: recording from a pyramidal cell 16 days in culture. External medium contained 2.1 mM Mg2+. Membrane potential was held at −58 mV by DC current injection. B: recording from a neuron 15 days in culture. Magnesium in the external solution was replaced by equiosmolar Na+. Membrane potential was held at −58 mV. C: pharmacology of pyramidal cell bursting in 0 [Mg2+]o. Top trace: recording from a pyramidal cell 14 days in culture exposed to 0 [Mg2+]o. Middle trace: recording from the same cell 25 min after addition of 40 μM D-APV, an NMDA receptor antagonist. Bursting activity is attenuated but not abolished. Bottom trace: neuronal excitatory activity ceases 15 min after further addition of 20 μM DNQX, an AMPA receptor antagonist. Recordings in A, B, and C are from different cells.

FIG. 3. Pharmacology of miniature excitatory postsynaptic currents (mEPSCs) in pyramidal cells exposed to 0 [Mg2+]o. Left-most trace: averaged mEPSC with biexponential decay kinetics. Exposure to D-APV (top arrow) produced an averaged mEPSC with a faster 10–90% rise time (1.2 ± 0.3 ms vs. 2.2 ± 0.3 ms) and a monoeXponential decay with τ decay of 33.3 ± 2.2 ms. If exposed to DNQX (instead of D-APV; bottom arrow), the resulting mEPSCs have a much slower rise (4.4 ± 0.9 ms) and τ decay (64.1 ± 3.7 ms).
event had a mean 10–90% rise time of 2.2 ± 0.3 ms and a biexponential decay with τ₁ and τ₂ of 16.2 ± 1.4 and 80.8 ± 3.2 ms, respectively. After addition of d-APV (Fig. 3, top arrow), neurons produced mEPSCs with mean rise of 1.2 ± 0.3 ms and a monoexponential decay with τ of 33.3 ± 2.2 ms (5 cells, 612 events). If, instead of blocking NMDA receptor-mediated events, TTX-treated cells were exposed to the AMPA receptor blocker DNQX (Fig. 3, bottom arrow), the resulting mEPSCs events, TTX-treated cells were exposed to the AMPA receptor blocker DNQX (Fig. 3, bottom arrow), the resulting mEPSCs were much slower kinetically, with a rise time of 4.4 ± 0.9 ms and decay τ of 64.1 ± 3.7 ms (373 events, 5 cells). These experiments suggested that NMDA receptors were present and active at synapses in the 0 [Mg²⁺]₀ condition. They also suggested that all NMDA receptor-mediated currents were blocked by 40 μM d-APV.

We further confirmed the presence of NMDA receptors at synapses on pyramidal neurons in culture at 13–17 days by immunohistochemical staining. The obligate NMDA receptor subunit NR1 was examined in relation to the presynaptic vesicular protein synaptophysin. In the cell illustrated in Fig. 4, synaptophysin clusters (green fluorescence) were evident in all cellular processes within the field of view (left panels; bottom panel is a higher magnification view of the boxed area in the top panel). NR1 staining was more discrete (red fluorescence) but was widespread in this neuron (middle panels). Overlay of synaptophysin and NR1 fluorescence revealed that most NR1 clusters colocalized with synaptophysin (right panels, orange clusters), suggesting a postsynaptic locus for NMDA receptors at this stage of neuronal development.

Presynaptic effects of reduced magnesium

In addition to unblocking NMDA receptors, lowered magnesium can induce increased excitability by presynaptic mechanisms. For example, magnesium displaces calcium at the presynaptic membrane surface layer (McLaughlin et al. 1978), thus reducing the amount of calcium available for entry into presynaptic terminals. Removal of magnesium would thus enhance presynaptic release by facilitating influx of calcium into the presynaptic terminal. We therefore examined the effect of magnesium reduction on the frequency of transmitter release. Figure 5, A and B illustrates sEPSCs recorded in normal and reduced magnesium. sEPSC frequency was markedly increased in the neuron recorded in 0 [Mg²⁺]₀; 90% of events occurred with interevent intervals <800 ms (Fig. 5C). In normal magnesium, the comparable interval was 5,200 ms (Fig. 5C). The shift in interevent interval for the cell pictured in Fig. 5 was significant [Kolmogorov–Smirnov 2-sample test; d = 0.58, P < 0.001]. Analysis of 8 cells each in normal and reduced magnesium revealed a similar increase in frequency. In each cell, the 90% interevent interval cutoff value decreased by ≥3-fold.

These data demonstrated an increased frequency of synaptic currents recorded postsynaptically, likely attributable in part to increased neuronal circuit excitability. However, they did not clarify whether magnesium removal also affected presynaptic glutamate release mechanisms. We first examined synaptic activity before and after the application of 1 μM TTX to assess the effect of 0 [Mg²⁺]₀ on action potential–dependent transmitter release. Figure 6, A and B shows examples of sEPSC (no TTX) and mEPSC (after TTX application) in cells 16 DIV in normal and 0 [Mg²⁺]₀ conditions. For the cellular activity shown, mEPSC frequency was significantly increased in the 0 [Mg²⁺]₀ condition (Fig. 6C; Kolmogorov–Smirnov 2-sample test; d = 0.26, P < 0.001). The number of mEPSCs recorded in a 4-min interval was 188 in normal magnesium and 331 in 0 [Mg²⁺]₀. In 6 pairs of cells exposed or not exposed to magnesium, the number of mEPSCs was 57.6 ± 12.9% higher in 0 [Mg²⁺]₀ conditions versus normal magnesium. This result indicates that action potential–independent neurotransmitter release was enhanced in the 0 [Mg²⁺]₀ condition.

We also examined whether quantal content, and thus the amplitude, of sEPSCs was altered by 0 [Mg²⁺]₀. Although no formal quantal analysis was performed, the mean number of neurotransmitter quanta released per presynaptic action potential was calculated by the direct method in control and 0

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**FIG. 4.** Hippocampal neuron 14 days in culture labeled with immunofluorescent antibodies to synaptophysin (green) and the NR1 subunit of the NMDA receptor. Labeling of each is extensive proximal and distal to the cell body within the field of view (top row, left and center). Overlay of the 2 images reveals colocalization of synaptophysin and NR1 (top right, orange punta). Box outlined in the top left photo is enlarged in the bottom row of images.
The preceding experiments demonstrated that 0 [Mg\(^{2+}\)]\(_{o}\) increased the strength of autapses (recurrent synapses from a neuron onto itself) or that of interconnections between pyramidal neurons. In the past, recurrent bursting was demonstrated in microcultures containing as few as 2 excitatory hippocampal neurons (Segal and Furshpan 1990). However, these results may not be applicable to the culture preparation used in our experiments in that cells were exposed to blockers of synaptic activity (kynurenic and elevated Mg\(^{2+}\)) for several weeks and exhibited bursting behavior only when cultures were washed free of these blockers. Chronic NMDA receptor blockade has been demonstrated to markedly increase clustering of NMDA receptors and promote their shift to a more synaptic distribution (Rao and Craig 1997). In contrast to this microculture model, the interconnectivity of multiple neurons was found necessary for recurrent bursting induced by penicillin, picrotoxin, or 4-amino pyridine in hippocampal slices (Hablitz 1984; Schwartzkroin and Prince 1978). To test whether interconnections among hippocampal neurons were necessary for recurrent bursting, they were grown at low density (10\(^3\) neurons/coverslip). Responses to control and 0 [Mg\(^{2+}\)]\(_{o}\) conditions were compared with those grown at high density (10\(^4\) neurons/coverslip) after 13–17 days in culture. Figure 7 shows current- and voltage-clamp recordings from pyramidal neurons cultured from the same embryonic hippocampal cells that were divided into low- and high-density groups. In low-density cultures, neurons were able to generate spontaneous action potentials. Exposure to 0 [Mg\(^{2+}\)]\(_{o}\) did not result in recurrent bursting (0/18 cells) but did increase the frequency of excitatory postsynaptic currents (Fig. 7A). sEPSC frequency increased significantly with 0 [Mg\(^{2+}\)]\(_{o}\) treatment in 9/13 cells (Fig. 7, B and C; Kolmogorov–Smirnov 2-sample test; \(d = 0.31, P = 0.04\)).

Although network excitability was not altered in 0 [Mg\(^{2+}\)]\(_{o}\) in low-density cultures, alterations in nonaction potential–dependent presynaptic glutamate release probability should still have been enhanced. We examined this possibility by comparing mEPSC frequency in low-density cultures under normal and 0 [Mg\(^{2+}\)]\(_{o}\) conditions. Figure 8A shows an example of mEPSCs recorded from cells 14 DIV. In normal magnesium, mEPSCs were infrequent; in 0 [Mg\(^{2+}\)]\(_{o}\), mEPSC frequency increased significantly. Eighty percent of events occurred with an interevent interval less than about 4,000 ms; in 0 [Mg\(^{2+}\)]\(_{o}\), the interval was significantly shorter at 1,550 ms (Kolmogorov–Smirnov test; \(d = 0.35, P < 0.001\)).

The finding that neurons from low-density cultures could not be induced to burst suggested that they form fewer active synapses that those cultured more densely. This was tested directly. The presence of synaptic contacts in cells from low- and high-density cultures was compared qualitatively by staining for synaptophysin distribution. In neurons from high-density culture pictured in Fig. 9A, synaptophysin clusters were ubiquitous on all neuronal processes. In several regions, individual puncta could not be distinguished because of their high density. In contrast, synaptophysin distribution was more sparse in the low-density example; individual synaptophysin puncta were easily distinguished on all processes, suggesting that neurons in low-density culture make and receive fewer synaptic contacts.

To examine active synaptic activity, neurons from high- and low-density cultures were treated with the fluorescent dye FM1-43, which is taken up in endocytosed synaptic vesicles
and is detectable on vesicle exocytosis, thus labeling synapses actively undergoing the exocytotic/endocytotic cycle (Cochilla et al. 1999). Figure 9B shows neurons from high- and low-density cultures exposed to FM1-43. The number of fluorescent puncta was markedly higher in the high-density neurons. Quantitative analysis of puncta in 5 cells each from high- and low-density cultures was performed by counting puncta in an area of about 30 × 30 μm. Puncta were counted in 2 areas per cell showing the greatest synaptic activity. In low-density neurons, the mean number of FM1-43 fluorescent puncta was 48.7 ± 5.1; for high-density neurons, activity was significantly higher, 177.2 ± 25.1 (P < 0.001, t-test). These data demonstrate that functional connectivity between cells, as measured by active synaptic contacts, is much reduced in low-density cultures.

**DISCUSSION**

The major findings of this study are 1) networks of cultured hippocampal neurons were capable of generating recurrent spontaneous action potential bursts very similar to those observed in intact hippocampal slice preparations when exposed to nominally zero magnesium; 2) postsynaptically, bursting behavior was attenuated, but not abolished, by D-APV, suggesting that abolition of magnesium blockade of NMDA receptor channels contributed to bursting onset and maintenance; 3) presynaptically, 0 [Mg2+]o caused an increased probability of glutamate release (analysis suggested a concomitant increase in mean quantal content); 4) reduced magnesium-induced alterations in pre- and postsynaptic neuronal properties were insufficient to induce bursting in lower-density cultures with sparse synaptic connections, indicating that bursting was a network, rather than intrinsic cellular, phenomenon. However, 0 [Mg2+]o did induce an increase in both sEPSC and mEPSC activity in low-density cultures.

**Cultured hippocampal neurons as a model for bursting**

Recently cell culture has become a tool for the study of synaptic interactions and plasticity (see Salter 2001 for review). Electrophysiological and morphological characterization has extended information on synapse function not amenable to other types of preparations (Gomperts et al. 2000; Stevens and Wesseling 1999). Cell culture has also been a
different cells.

valuable tool for investigation of pathological conditions relating to alteration of normal synaptic function (DeLorenzo et al. 1999), and culture preparations (Sombati and Avoli et al. 1986; Avoli et al. 1987), slice culture preparation results from synaptic interactions rather than intrinsic neuronal properties. However, our data do not rule out the latter. Bursting behavior occasionally occurred in normal external magnesium. It is unlikely that neuronal hyperexcitability in these cases was the result of hypoxia (Rubas et al. 2003) or pH alteration. Cultures included both CA1 and CA3 pyramidal neurons; the intrinsic pacemaking properties of the latter have been well described and modeled (Traub et al. 1991). It is possible that, given sufficient density and interconnectedness of a particularly CA3 pyramid cell-rich region of the culture, spontaneous activity could occur. Numerous studies have shown that network-driven recurrent hyperexcitable discharges in intact hippocampus originate in CA3 and propagate to CA1 (Hablitz 1984; Mody et al. 1988).

The importance of network interactions in enabling burst activity is supported by several lines of evidence. First, neuronal hyperpolarization of 10–20 mV from the resting potential did not eliminate bursting, although burst duration and frequency were reduced. Second, neurons from low-density cultures did not burst under either normal or 0 [Mg2+]o conditions. Third, pharmacological blockade of synaptic excitation eliminated bursts in 0 [Mg2+]o. Simultaneous dual recordings will be required to definitively address the question of whether neurons are receiving synchronized excitatory barrages.

Reduced magnesium bursting: postsynaptic mechanisms

Hippocampal hyperexcitability induced by magnesium reduction or removal has been noted for several years. The effect is sufficiently pronounced that magnesium deficiency has been used as a model of temporal lobe seizures since the late 1970s (Buck et al. 1978). Numerous reports have noted the dependency of reduced-magnesium bursting on functional NMDA receptor/channels (Albowitz et al. 1997; Collingridge et al. 1988; Gulyas-Kovacs et al. 2002; Gutierrez et al. 1999; Hamon et al. 1987; Quilichini et al. 2002; Tancredi et al. 1988). This is presumably attributable to the well-documented postsynaptic effect of magnesium in electrostatically blocking NMDA receptors/channels from the outside of the cell at the resting membrane potential (Dingledine et al. 1990). Our results support these previous studies. First, immunofluorescence studies showed NMDA receptors (the obligate NMDA receptor subunit NR1) were localized with synaptophysin in burst-capable cultures, indicating a synaptic locus. Second, magnesium removal was a requirement for bursting in nearly all experiments, whereas bursting was attenuated by the specific NMDA receptor antagonist d-APV. Thus NMDA receptor activation by reduced magnesium appears crucial to initiation and maintenance of neuronal bursting in our preparation.

However, our data suggest additional excitatory postsynaptic mechanisms may be operative. Excessive neuronal activity (bursts of high-frequency EPSCs) induced by reduced magnesium did not return to normal after blockade of NMDA receptors with d-APV. EPSC bursting still occurred, although interburst intervals were increased and burst duration was lessened. Blockade of AMPA-type glutamate receptors was necessary to eliminate activity. These observations suggest that increased AMPA receptor activity as well as activation of NMDA receptors may contribute to neuronal hyperexcitability in 0 [Mg2+]o. One possible explanation for this observation may be a recent hypothesis of the physiological substrate for long-term potentiation, a form of synaptic plasticity that may underlie learning and memory. Liao et al. (2001) demonstrated that activation of NMDA receptors in culture results in rapid recruitment (within minutes) of AMPA receptors to NMDA receptor synaptic sites with a parallel increase in mEPSC frequency. Similar enhancement of mEPSCs occurred after brief, focal application of glycine to activate NMDA receptors (Lu et al. 2001). Augmentation of excitatory transmission was exocytosis dependent and presumably involved cycling of AMPA receptors from remote locales to the site of NMDA activation (Liang and Huganir 2001; Luscher and Frerking 2001; Luscher et al. 1999). Such an influx of AMPA receptors to synaptic sites could result in the d-APV-resistant excitability we have observed.

Reduced magnesium bursting: presynaptic mechanisms

Although enhancement of NMDA receptor-mediated excitability has been demonstrated to play a role in reduced magne-
sium induced bursting, it is unlikely to play a significant role in the presynaptic alterations we have observed in zero magnesium: increased sEPSC frequency and an apparent increase in mean quantal content. These changes suggest that the probability of glutamate release was increased by magnesium reduction.

One possible mechanism was first proposed by Frankenhauser and Hodgkin (1957). They reasoned that reductions in divalent cations could increase neuronal excitability by decreasing charge screening at the membrane surface. This would cause a smaller portion of the transmembrane potential to fall across the lipid bilayer (McLaughlin et al. 1971), thus decreasing the electric field sensed by voltage-dependent conductances such as calcium channels in the synaptic terminal. Muller and Finkelstein (1974) later proposed a more magnesium-specific model. They hypothesized that increasing Mg\(^{2+}\) concentration displaces calcium in the surface charge layer adjacent to the negatively charged membrane, thus decreasing the amount of calcium available for influx into synaptic terminals. Reducing or eliminating Mg\(^{2+}\) would have the opposite effect, increasing the effective Ca\(^{2+}\) concentration at the mouth of presynaptic Ca\(^{2+}\) channels. This mechanism may explain the increase in intracellular calcium observed in acutely isolated CA1 pyramidal neurons in the presence of reduced extracellular magnesium, an influx mediated by voltage-dependent calcium channels (Zhang et al. 1996). Thus one presynaptic effect of reducing Mg\(^{2+}\) would be to increase the probability of release as proposed in the Katz model of neurotransmission (Katz 1971). The probability of release reflects whether the release site is occupied and the availability of transmitter (Dobrunz and Stevens 1997; Hanse and Gustafsson, 2002; Staley et al. 1998, 2001) but can be modulated by calcium and magnesium (Bouron 2001). Reduced magnesium may also cause increased glutamate release by facilitating calcium egress from intracellular stores by reduced magnesium inhibition of ryanodine receptors (Masumiya et al. 2001). Whether such a process occurs is problematical given that fluctuations in the extracellular magnesium concentration do not appear to alter intracellular levels (Zhang et al. 1996).

Such mechanisms could explain the marked increased in sEPSC frequency, reflecting increased presynaptic glutamate release, observed in reduced magnesium. It may also explain the calculated increase in mean quantal content. Although we did not attempt a formal quantal analysis for this study, we did use 2 different methods for determining quantal content. These methods did not agree as to the extent of the increase, but in both instances, the increase was significant.

**Reduced magnesium bursting: connectivity**

Physiological and modeling studies in the hippocampus have suggested that synchronized bursting in some systems of pyramidal neurons may require connectivity among a critical minimum of cells and sufficient excitatory synaptic strength to drive the network (Bains et al. 1999; Traub and Wong 1983; Traub et al. 1995). A model network simulation of hippocampal neuronal bursting demonstrated that bursting can occur, even in the presence of neuronal inhibition, only if connectivity between neurons in the circuit is adequately dense. Low connectivity prevents the development of bursting (Traub et al. 1987). In vitro studies in hippocampus and neocortex suggest that the relative paucity of excitatory synaptic connections...
likely contributes to neonate rodent seizure resistance (Swann and Hablitz 2000).

We tested this idea by subjecting low-density cultures to the same conditions that produced bursting behavior in high-density cultures. The contrast between the 2 preparations when subjected to $0 [\text{Mg}^{2+}]_o$ was evident in 2 regards. No multiple action potential bursts were observed in any low-density neuron tested regardless of the number of days grown in culture and immunofluorescent labeling for synaptophysin indicated many fewer potential synaptic sites. This latter finding was confirmed by labeling of synapses with FM1-43; the number of actively exocytosing synapses was significantly less in low-density cultures. Increases in both sEPSC and mEPSC frequency were evident in $0 [\text{Mg}^{2+}]_o$, indicating that presynaptic mechanisms for increasing glutamate release operate in low-density as well as high-density cultures.

The importance of connectivity in generating aberrant excitatory neuronal activity may be appreciated by an examination of animal models of temporal lobe seizures. The neuronal activity described in this study (multiple action potentials superimposed on prolonged depolarizations) is similar to that observed in these models. On a cellular level, seizure activity is manifested in synchronous excitatory activity among a large population of neurons (Dudek et al. 1999). A prominent anatomic feature in many animal models of chronic temporal lobe seizures is the sprouting of neuronal processes and subsequent formation of aberrant, presumably glutamate-mediated, excitatory synaptic interactions (Lehmann et al. 2001; Sutula et al. 1996; Wuarin and Dudek 1996), although sprouting is not a universal feature of these models (Swann et al. 2001). Recently, Lynch and Sutula (2000) demonstrated aberrant, apparently monosynaptic recurrent excitatory circuitry that likely underlies abnormal neuronal activity in this model.

ACKNOWLEDGMENTS

We thank M. Alietta and C. Gregory for expert glial and hippocampal culture preparations.

GRANTS

This research was funded by National Institute of Neurological Disorders and Stroke Grants NS-37192 to P. S. Mangan and NS-02081 and NS-40337 to J. Kapur.

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