Synaptic kainate receptors tune oriens-lacunosum moleculare interneurons to operate at theta frequency.
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GABAergic interneurons of the hippocampus play an important role in the generation of behaviorally relevant network oscillations. Among this heterogeneous neuronal population, somatostatin (SOM)-positive oriens-lacunosum moleculare (O-LM) interneurons are remarkable because they are tuned to operate at theta frequencies (6–10 Hz) in vitro and in vivo. Recent studies show that a high proportion of glutamatergic synapses that impinge on O-LM interneurons are mediated by kainate receptors (KA-Rs). In the present study, we thus tested the hypothesis that KA-Rs transmit afferent inputs in O-LM neurons during synaptic stimulation at theta frequency. We combined multibeam two-photon calcium imaging in hippocampal slices from SOM–enhanced green fluorescent protein (EGFP) mice, to record the activity of SOM cells as well as hundreds of neurons simultaneously, and targeted electrophysiological recordings and morphological analysis to describe the morphofunctional features of particular cells. We report that EGFP-positive O-LM neurons are the only subtype of interneuron that reliably follows synaptic stimulation of the alveus in the theta frequency range. Electrophysiological recordings revealed the crucial contribution of KA-Rs to the firing activity and to the glutamatergic response to theta stimuli in O-LM cells compared with other cell types. The reliable activation of O-LM cells in the theta frequency range did not simply result from the longer kinetics of KA-R-mediated postsynaptic events ($EPSP_{KA}$) but presumably from a specific interaction between $EPSP_{KA}$ and their intrinsic active membrane properties. Such preferential processing of excitatory inputs via KA-Rs by distally projecting GABAergic microcircuits could provide a key role in theta band frequency oscillations.

Key words: kainate; O-LM; theta; imaging; interneuron; network

Introduction

GABAergic cells control the generation of network oscillations (Cobb et al., 1995; Freund and Buzsáki, 1996; Whittington et al., 1997; McBain and Pisah, 2001; Klausberger et al., 2003; Whittington and Traub, 2003; Somogyi and Klausberger, 2005). A central issue is to understand how different GABAergic circuits integrate ongoing activity to generate behaviorally relevant oscillations. Recent studies suggested that the integrative properties of interneurons are strongly influenced by their synaptic inputs (Pouille and Scanziani, 2004; Oren et al., 2006). Glutamatergic synaptic transmission is principally mediated by AMPA receptors, but kainate receptors (KA-Rs) also play a central role (Castillo et al., 1997; Vignes and Collingridge, 1997; Cossart et al., 1998, 2002; Frerking et al., 1998; Bureau et al., 1999) although in a manner that is remarkably cell specific (Ben Ari and Cossart, 2003). Various types of interneurons express KA-Rs (Cossart et al., 1998, 2002; Frerking et al., 1998; Mulle et al., 2000; Ali, 2003; Lerma, 2003); however, the distribution of KA-R-operated synaptic transmission among GABAergic cells is still unknown. This is important in view of the slower kinetics of KA-versus AMPA-R-mediated EPSCs (Freerking et al., 1998; Cossart et al., 2002), which could enable the generation of oscillations with different dominating frequencies (Freerking and Ohlinger-Frerking, 2002). Therefore, the KA-R subtype is well suited to mediate cell-specific temporal information processing in interneurons, but its functional distribution and precise conditions of activation remain to be determined.

We addressed this issue using multibeam two-photon microscopy, on-line analysis, and electrophysiology, to record electrically evoked network dynamics and describe the distribution of active cells (Cossart et al., 2005). We concentrated on CA1 somatostatin-expressing stratum oriens interneurons and specifically on oriens-lacunosum moleculare (O-LM) cells, the predominant horizontal cell type in this layer (Ali and Thomson, 1998; Maccarferri, 2005). These neurons participate in hippocampal theta activity in vivo (Buzsáki, 2002; Klausberger et al., 2003) and in vitro (Pike et al., 2000; Gillies et al., 2002; Hajos et al., 2004; Gloveli et al., 2005). They also receive a large input mediated by KA-Rs (Cossart et al., 2002). To target this subpopulation, we used transgenic mice expressing enhanced green fluorescent protein (EGFP mice) in a subset of somatostatin-containing interneurons [EGFP-positive (EGFP$^+$)] that mostly comprise O-LM cells in the CA1 stratum oriens (Oliva et al., 2000). We report that O-LM neurons reliably follow theta stimulation (TS) of the
alveus, an effect selectively mediated by KA-R synaptic activation. Stratum oriens interneurons that did not follow theta stimulation via KA-R activation were perisomatic (O-P), bistratified (O-Bi), or septum/back-projecting (O-S/BP) cells but not O-LM neurons. Electrophysiological recordings from EGFP-positive/O-LM neurons showed that postsynaptic KA-Rs contribute to action potential firing at 10 Hz during theta stimulations. Short-term facilitation or summation of the postsynaptic response in O-LM cells were not observed in the theta frequency range. The reliable activation of O-LM cells by theta frequency stimulations did not simply result from the longer kinetics of KA-R-operated synaptic transmission but presumably from a specific interaction between the postsynaptic response and intrinsic membrane properties. These findings should support the recruitment of O-LM cells during hippocampal theta activity and provide a synaptic framework for the mechanisms driving theta rhythms.

Materials and Methods

Slice preparation and two-photon imaging. Transverse hippocampal slices (350 μm thick) were prepared from 12- to 17-d-old transgenic mice expressing EGFP in somatostatin-containing neurons (Oliva et al., 2000) using a Microm (Walldorf, Germany) tissue slicer (Microtome HM 650V) with ice-cold oxygenated modified artificial CSF (ACSF) (0.5 mM CaCl2 and 7 mM MgSO4, and in which NaCl was replaced by an equimolar concentration of choline). Slices were then transferred for rest (~1 h) at room temperature in oxygenated regular ACSF containing the following (in mM): 126 NaCl, 3.5 KCl, 1.2 NaH2PO4, 26 NaHCO3, 1.3 MgCl2, 2.0 CaCl2, and 10 D-glucose, pH 7.4. For AM loading, slices were incubated in a small vial containing 2.5 ml of oxygenated ACSF with 25 μg of a 1 mM fura-2 AM solution (in 100% DMSO; Invitrogen, Carlsbad, CA) for 20–30 min. Slices were incubated in the dark, and the incubation solution was maintained at 35–37°C. Experiments were performed at 30–32°C with regular ACSF and continuously aerated with 95% O2/5% CO2. Imaging was performed with a multibeam two-photon LASER scanning system (Trimscope; La Vision Biotec, Bielefeld, Germany) coupled to an Olympus Optical (Hamburg, Germany) microscope. This system is based on a patented beam splitter that splits up the incoming femtosecond LASER beam (provided by a titanium:sapphire LASER source; Chameleon; Coherent, Santa Clara, CA), into 64 beamlets, which are scanned simultaneously in the slice. This results in 64 times higher image acquisition rates compared with conventional multiphoton scanning microscopes. Images were acquired through a CCD camera (Imager 3QE; La Vision Biotec). Acquisition frequency was between 150 and 210 ms per frame. Slices were imaged using a low-magnification, high numerical aperture (NA) objective (20×, NA 0.95; Olympus Optical). The size of the imaged field was typically ~430 × 380 μm2. Calcium and GFP fluorescence signals were separated by the excitation wavelength of the LASER source (780 and 900 nm, respectively).

Electrical stimulation. Axons of CA1 pyramidal cells were stimulated via a custom-made bipolar NiCh electrode (model 762000; A-M Systems, Sequim, WA) placed ~100 μm from the imaged region in the alveus (intensity of 5–10 μA, 50 μs). A Grass Instrument (Quinacy, MA) stimulation unit was used (model SIU5B). The TS protocol included 10 stimuli delivered at a frequency of 10 Hz every 30 s. Action potential firing triggered by antidromic activation resulting from the direct stimulation of the cell membrane by the electrode was discarded from the analysis. Antidromically evoked calcium events, i.e., events that were not blocked by antagonists of ionotropic GABAergic and glutamatergic transmission, were also systematically discarded.

Analysis. Analysis was performed with custom-made software in Matlab (MathWorks, Natick, MA) written by D. Aronov (Massachusetts Institute of Technology, Cambridge, MA). We developed a program aimed at the automatic identification of loaded hippocampal cells and at measuring their fluorescence as a function of time. This program is an improved version of the previously designed software for cortical slices analysis (Cossart et al., 2003). We encountered two major problems for an automatic identification of cells in an image: (1) variations in background fluorescence, which precluded the use of a uniform threshold value throughout the image, and (2) the inability of a simple thresholding procedure to separate nearby cells. The latter problem was especially severe in the dense pyramidal cell layer of the hippocampus. As before (Cossart et al., 2003), we solved the problem of background variations by normalizing each pixel by the average fluorescence in its vicinity. We also convolved the image with a two-dimensional Gaussian (σ = 6 μm), which emphasized circular neuronal shapes and partially separated nearby neurons. A threshold (usually top 10 percentile of the overall pixel fluorescence distribution) was applied to the image to separate cell contours from the background. To complete the separation of nearby cells, we measured a circularity threshold for every contour, defined as $c = P^2/(4\pi A)$, where $P$ is the perimeter and $A$ is the area of a contour. High values of $c$ (usually $>4$) identified highly noncircular shapes usually indicative of unseparated cell contours. Local fluorescence maxima were identified within such contours, and the contours were separated into a corresponding number of concave shapes. All image processing was performed on time averaged records of the calcium signal. The calcium signal of each cell was the average fluorescence within the contour of that cell, measured as a function of time. Signal processing algorithms from the MiniAnalysis software (Synaptosoft, Decatur, GA) were used to automatically detect the onsets and offsets (time of half-amplitude decay) of calcium signals within the traces of individual cells. Calcium changes below 5% ΔF/ΔF0 were discarded because calibration of the imaging system showed that, on average, a single action potential will produce a fluorescence change always above this value (average ΔF/ΔF0, 6.5 ± 0.3%). The entire procedure could be performed on-line sufficiently quickly to identify cells for targeted patch-clamp recordings. However, fluorescence traces were also visually scanned off-line to correct for the detection of false positives. A significant calcium response to a TS is an event that passed the threshold for detection within a 400 ms time window after the time of the stimulus. The analysis was performed separately for each hippocampal layer, with the borders between layers drawn manually.

Electrophysiology. Slices from EGFP mice were prepared as for imaging except that a few recordings were also performed in 1-month-old animals ($n = 3$). Interneurons recorded for miniature glutamatergic activity analysis were visualized by infrared video microscopy using an upright Zeiss (Le Pecq, France) microscope equipped with a 40× water immersion objective. Fluorescent interneurons were identified with a UV lamp (xenon, excitation filter 470/40) and recorded using a pipette solution containing: (1) in voltage-clamp mode, 120 mS Cs-glutamate, 10 mM MgCl2, 0.1 mM CaCl2, 1 mM EGTA, 5 mM Na-glutamate, triphosphate, 10 mM HEPES, and 0.5% bicocytin; (2) in current-clamp mode, 130 mM K-methylsulfate, 5 mM KCl, 5 mM NaCl, 10 mM HEPES, 2.5 mM Mg-ATP, 0.3 mM GTP, and 0.5% bicocytin. The osmolarity was 265–275 mOsm, pH 7.3. Miniature currents had to exceed a minimum level of 4–8 μA. Uncompensated series resistances were, on average, 14 MΩ.

Signals were fed to an EPC10 (HEKA Elektronik, Heidelberg, Germany), filtered (3 kHz), digitized (10 kHz) with a Labmaster interface card to a personal computer, and acquired using Axoscope 7.0 software (Molecular Devices, Union City, CA). Data was analyzed with MiniAnalysis 5.1 program (Synaptosoft). Spontaneous glutamatergic currents were measured at ~60 mV and miniature EPSC (mEPSCs) were recorded in the presence of TTX (1 μM), bicuculline (10 μM), and n-AVP (40 μM). Addition of GYKI 52466 [4-(8-mercapto-9H-1,3-dioxolo[4,5-h][2,3]benzodiazepin-5-yl)-benzenemethy]hydrochloride (100 μM) or 2,3-dihydroxy-6-nitro-7-sulfonyl-benzoic[quinoxaline (NBQX) (1 μM) blocked AMPA-R-mediated-mEPSC (mEPSCAMP), and further addition of CNXQ (50 μM) blocked all mEPSCs. Single and averaged events (200 events per cell) were fully characterized: rise times (~10–90%), amplitudes, and decay time constants were calculated.

Kinetic analysis. Identification of KA-R-mediated-mEPSC (mEPSC KA) and mEPSCAMP was based on their kinetics profile (Cossart et al., 2002). Briefly, spontaneous mEPSCAMP were recorded in the presence of 1 μM TTX, 10 μM bicuculline, and 40 μM n-AVP were recorded from CA1 stratum oriens interneurons, and each single event was fully characterized by the following parameters: rise time (10–90%), amplitude, charge (area under EPSCs), and decay time constants. In control conditions, two types of mEPSCs could be distinguished based on the time course of their decay:
fast and slow. After 10 min application of AMPA-R antagonists (100 μM GYKI 52466 or 1 μM NBQX), fast events were abolished and only KA-R-mediated events with a slower time course and a small amplitude remained. Slow events were blocked by SYM 2081 [(25,4R,6R)-2-methylglutamic acid], a functional antagonist for KA-Rs that required 15 min drug application. Decay times of slow events in control and mEPSC<sub>KA</sub> left after blocking mEPSC<sub>AMPA</sub> had the same kinetics (n = 17 interneurons; one-way ANOVA, p > 0.05), demonstrating that slow kinetics events were KA-R mediated. To separate fast mEPSC<sub>AMPA</sub> from slow mEPSC<sub>KA</sub>, a “kinetic limit” was determined. Events were individually fitted, and decay times were plotted versus rise times and versus amplitudes. Fast mEPSC<sub>AMPA</sub> and slow mEPSC<sub>KA</sub> clustered in two distinct areas of these scattered plots. Limits to segregate between the two populations of events were as follows: mEPSC<sub>AMPA</sub> rise-time <1 ms and decay time <4 ms; mEPSC<sub>KA</sub> rise time >1 ms and decay time >4 ms.

**Pharmacology.** Bicuculline, NBQX, n-APV, CNQX, GYKI 52466, and SYM 2081 were purchased from Tocris (Ellisville, MO), TTX was from Latoxan (Valence, France), and CX546 [1-(1,4-benzodioxan-6-ylcarbonyl)piperidine] was from Sigma-Aldrich (Lyon, France).

**Morphology.** Slices were processed for the detection of biocytin-filled neurons according to a previously established procedure (Cossett et al., 2006). Briefly, slices were fixed overnight at 4°C in a solution containing 4% paraformaldehyde in 0.1 M phosphate buffer (PB), pH 7.4. After fixation, slices were rinsed in PB, cryoprotected in sucrose (30% in PBS), and quickly frozen on dry ice. To neutralize endogenous peroxidase, slices were pretreated for 30 min in 1% H<sub>2</sub>O<sub>2</sub>. After several rinses in 0.01 M PBS, pH 7.4, slices were incubated for 24 h at room temperature, in 1:100 avidin–biotin peroxidase complex (Vector Laboratories, Burlingame, CA) diluted in PBS containing 0.3% Triton X-100. After 30 min rinses in PBS, slices were processed with 0.04% 3,3′-diaminobenzidine-tetrahydrochloride (Sigma, St. Louis, MO) and 0.006% H<sub>2</sub>O<sub>2</sub> diluted in PBS. Seventy-eight biocytin-filled interneurons were identified. Among these 78 interneurons, we could obtain both a complete characterization of mEPSCs/EPSPs and a complete axonal and dendritic labeling from 72 interneurons. Neurons were classified based on previously established morphological criteria. We classified as O-S/BP interneurons with axonal branches invading the CA3 region (stratum oriens and/or stratum radiatum) and possibly the dentate gyrus and/or axonal branches traveling through the alveus and reaching the fimbria (see Figs. 3, 7). This population is likely to comprise at least two different cell types (i.e., septum and BP interneurons) that essentially differ by their septal innervation. We decided to group them because both were shown to innervate specifically GABAergic interneurons in the hippocampus (Gulyas et al., 2005) and because they are hardly morphologically distinguishable in hippocampal slices: they display an axonal arborization that crosses subfield boundaries and sparsely spiny dendrites (a property that distinguishes them from O-LM cells with a cut axon). These interneurons were reconstructed for morphometric analysis with a computer-assisted system (Neurolucida; MicroBrightField, Williston, VT) attached to a Nikon (Tokyo, Japan) microscope.

**Results**

**CA1 stratum oriens EGFP-positive cells are activated by theta stimulation**

We used multibeam two-photon imaging in hippocampal slices from EGFP mice, loaded with a calcium indicator to determine the network dynamics evoked by repetitive (30 s interval) electrical TS (train of 10 stimuli at 10 Hz; see Materials and Methods). Stimulation electrodes were placed in the alveus ∼100 μm away from the imaged area (Fig. 1). Activity was monitored in hundreds of neurons simultaneously, spanning across all CA1 hippocampal layers. We performed automated signal processing with custom-made software to obtain raster plots describing the activity of all neurons imaged as a function of time (Figs. 1, 2). A 5% DFF amplitude threshold was chosen for calcium signal detection (more than five times larger than the noise SD) because calibration of the imaging system showed that a single action potential produced, on average, a fluorescence change always above this value (Fig. 1) (see Materials and Methods). Detected calcium events were indeed dependent on action potential firing because they were...
Figure 2. EGFP-positive neurons preferentially respond to theta stimulation.

A1. Raster plot of the activity from a movie; filled arrows mark the TS stimulation times. Each row represents the activity within a cell, and each dot represents the onset of a calcium transient. Antidromically activated neurons were discarded. A2. Contour plot of the neurons that fluorescence traces are shown in A3 (black filled). Green filled contour indicates a EGFP + neuron. Scale bar, 100 μm. A3. Representative calcium fluorescence traces during a TS protocol in control conditions (green, EGFP + neurons; black, EGFP − neurons). Note the evoked responses to every stimulus in the EGFP − neuron. o, Stratum oriens; t, stratum radiatum; p, stratum pyramidale. B1. Targeted cell-attached recordings from an EGFP + cell in response to 10 alveus stimulations in control ACSF (2 consecutive trains of stimulation separated by 30 s; open arrows mark each stimulation within a TS train). Note that each stimulation within the train drove the cell to fire an action potential (1 spike is indicated by a large filled arrow; enlarged in B2). Average spike probability as a function of stimulus number is plotted on the bottom histogram. Error bars indicate SEMs. B2. Representative superimposed cell-attached recordings of action potentials evoked in response to a train of 10 consecutive alveus stimulations in an EGFP + cell (10 Hz). Latency histogram of the recorded cell is plotted below. C. Bar graphs of the mean calcium response probability (fraction of trains within a stimulation protocol that evoked a suprathreshold calcium response; antidromic stimulation was discarded) in response to TS protocol in EGFP + (dark green; n = 21) and EGFP − (black; n = 884) cells. Note that mean action potential probability (fraction of stimuli evoking an action potential) assessed by targeted cell-attached recordings in EGFP + cells (n = 10; light green) is similar to the calcium response probability measured in the same cell population. The calcium response probability was significantly higher in EGFP + cells compared with EGFP − cells (Mann–Whitney test, ***p < 0.001).

KA-Rs are major contributors to the spontaneous glutamatergic currents received by O-LM cells

As a first step, we quantified the contribution of KA-Rs to glutamatergic transmission in different cell types without pharmacological manipulation. We performed whole-cell voltage-clamp recordings of glutamatergic activity in 13 EGFP + and 33 EGFP − interneurons in the CA1 stratum oriens using biocytin-filled electrodes. Six CA1 stratum oriens interneurons were also recorded in wild-type mice (Table 1).

EPSCka can be distinguished from EPSCampa solely on the basis of their kinetics profile (Cossart et al., 2002; Epstein et al., 2005). We thus determined the kinetic properties of EPSCs and focused on action potential-independent mEPSCs, which represent quantal events that are particularly suited for kinetic analysis (Cossart et al., 2002; Epstein et al., 2005). Miniature EPSCs were recorded in the presence of the sodium channel blocker TTX (1 μM) at the reversal potential for GABAergic currents (V_hold = −60 mV). The rise and decay time constants were calculated for each miniature event. mEPSCs amplitude, frequency, and charge were also quantified (Table 2). We included only interneurons that could be fully morphologically identified (Fig. 3) (n = 40): O-LM

blocked by the sodium channel blocker TTX (1 μM; n = 3 movies, 259 cells; data not shown). Evoked calcium events could be triggered by either an EPSP or antidromic activation producing action potential firing. Antidromic activation resulted from the direct electrical stimulation of the cell membrane and therefore was not blocked by antagonists of ionotropic synaptic transmission. Antidromically evoked calcium events were discarded from the analysis (see Materials and Methods). We quantified the response of imaged neurons to TS protocols by calculating a “calcium response probability” per imaged neuron that is the fraction of TS trains within the course of the experiment that evoked a detected calcium event in the cell (i.e., a calcium signal above the 5% DF/F detection threshold). Cells that could never be activated during the TS protocol were discarded. Such calcium response probability thus indirectly quantifies the fraction of TS trains evoking at least one action potential. We found that, in control conditions, calcium response probability was 100 ± 0% in all EGFP + cells (n = 21) and 27 ± 7% in EGFP-negative (EGFP −) cells (n = 884 neurons, 7 movies; p < 0.001) (Fig. 2). Response probability in EGFP − cells was not different between layers (39 ± 7% in stratum pyramidale vs 24 ± 11% in stratum oriens and 28 ± 11% in stratum radiatum; p > 0.1, Mann–Whitney test; n = 8 movies) but was always significantly lower than in EGFP + cells, most of which were located in stratum oriens. We next performed targeted cell attached recordings from EGFP + cells to measure action potential firing in these neurons. In keeping with imaging experiments, EGFP + cells fired at least one action potential in response to each TS train (99 ± 1% of trains evoked at least one action potential; n = 98 trains, 10 EGFP + neurons) (Fig. 2). Furthermore, these experiments showed that EGFP + cells also reliably followed every single stimulation within trains because spike probability for the first stimulus was 77 ± 25% and remained stable during the train (spike probability for the fifth stimulus was 84 ± 22%; n = 8 EGFP + neurons; p = 0.53, Mann–Whitney test). Spike latency (measured as the delay between the stimulus artifact and the action potential onset) was 11.2 ± 1.9 ms (n = 10 EGFP + cells) (Fig. 2). The spike jitter was relatively large, suggesting that the response was driven by an EPSP with slow kinetics (1.7 ms jitter; n = 8 EGFP + cells) (Pouille and Scanziani, 2004). We conclude that CA1 stratum oriens EGFP + neurons respond to synaptic stimulations at theta frequency. This property distinguishes them from other CA1 neurons. Because KA-Rs represent a significant contribution to the glutamatergic influx received by CA1 stratum oriens interneurons (Cossart et al., 1998, 2002), we next asked whether KA-Rs could mediate such selective synaptically evoked response.
cells (n = 20); O-P cells (n = 11) (Harris et al., 1985; Kosaka et al., 1985; Maccaferri et al., 2000; McBain and Fisahn, 2001), O-Bi cells (n = 5) (Sik et al., 1995), and O-S/BP interneurons (n = 4) (Sik et al., 1995; Gulyas et al., 2003) (see Materials and Methods). As detailed in our previous studies (Cossart et al., 2002; Epsztein et al., 2005), mEPSC_{KA} were identified in each recording based on their kinetics and pharmacological properties (Fig. 3) (see Materials and Methods) to determine their relative contribution to the global miniature glutamatergic influx for each cell type. We found that the decay time constants of averaged mEPSCs in O-LM cells were slower (6.1 ± 0.2 ms; n = 20; p < 0.001) than those in O-P (3.6 ± 0.5 ms; n = 11) and O-Bi (2.8 ± 0.3 ms; n = 5) interneurons. In O-S/BP interneurons, there was a larger variability of kinetics between cells (5.4 ± 1.3 ms; n = 4). The difference in decay times was unlikely to result from different KA-R or AMPA-R subunit compositions between interneuron types because decay times for mEPSC_{KA} or mEPSC_{AMPA} were not significantly different between cell types (Table 2).

We then determined the relative contribution of KA-R-mediated synaptic currents in different types of interneurons. We found that mEPSC_{KA} accounted for a large majority of the total quantal glutamatergic influx in the O-LM cell type as opposed to other neurons. Thus, mEPSC_{KA} represented 88 ± 2% (n = 20) (Fig. 3) of the total quantal glutamatergic activity on O-LM neurons but less than half (43 ± 6%, n = 11 and 38 ± 6%, n = 5) (Fig. 3) of the activity in O-P and O-Bi interneurons, respectively. In O-S/BP neurons, there was also less contribution of KA-R to miniature activity (57 ± 7%; n = 4) although the values were more variable most likely attributable to the weaker reliability of the morphological characterization of this cell type (see above).

We conclude that synaptic KA-Rs are differentially distributed among CA1 stratum oriens interneurons and represent the major source of the ongoing glutamatergic influx received by O-LM cells.

A selective processing of postsynaptic glutamatergic responses evoked at theta via KA-Rs in O-LM cells

We next asked whether the differential distribution of KA-Rs among interneuron subtypes could account for the reliable activation of EGFP + cells during TS protocols. We first isolated KA-R-mediated responses in the presence of NMDA and GABAA receptor antagonists (control conditions: response probability in EGFP + and EGFP − cells was not significantly different from drug-free saline), relying on their resistance to AMPA-R antagonists (GYKI 52466 at 100 μM or NBQX at 1 μM) (Paternain et al., 1995; Bureau et al., 1999) (see Materials and Methods) and blockade by the mixed AMPA/KA-R antagonist CNQX (50 μM). Imaging experiments indicated that, in the presence of AMPA-R antagonists, calcium response probability was significantly higher in EGFP + than EGFP − neurons (58 ± 17 vs 9 ± 6%, respectively; n = 8 EGFP + cells, n = 884 neurons imaged; p < 0.05). Furthermore, blockade of AMPA-Rs significantly reduced response probability in EGFP − cells relative to control in all layers (reduction to 33 ± 4% of control; p < 0.05) (Fig. 4), whereas probability in EGFP + cells was also reduced but not significantly (to 65 ± 18% of control; p > 0.05) (Fig. 4). However, relative changes in probability resulting from AMPA-R blockade were not significantly different when comparing EGFP + and EGFP − cells (p = 0.3, Mann–Whitney test). Indeed, two subpopulations of EGFP + neurons could be distinguished in conditions in which synaptic excitation was provided only by KA-Rs because calcium response probability remained unchanged relative to control in half of EGFP + cells (n = 8 EGFP + cells). The amplitude of the evoked calcium transient in KA-R-responsive EGFP + neurons was reduced, but not significantly, in the presence of blockers, to 72 ± 29% of control values (n = 8 EGFP + cells; p > 0.05), suggesting that fewer action potentials were triggered when blocking AMPA receptors and decreasing network excitability or when KA-Rs were partially blocked by AMPA-R antagonists (Castillo et al., 1997). Evoked calcium responses in GYKI-responsive EGFP + neurons were fully suppressed when KA-Rs were blocked by further adding CNQX (50 μM) to the perfusion saline. Therefore, synaptically evoked responses presenting a KA-R pharmacology (i.e., resistant to AMPA, NMDA, and GABAA blockers but suppressed by CNQX) were selectively present in the EGFP + cell population.

Second, to determine the involvement of KA-Rs in more physiological conditions, we used a functional antagonist for KA-Rs (SYM 2081 at 10 μM) (Zhou et al., 1997; Paternain et al., 1995; DeVries, 2000; Cossart et al., 2002; Epsztein et al., 2005) that did not affect the global level of spontaneous activity in the network (fraction of active cells in control, 46 ± 5 vs 40 ± 6% after SYM 2081; n = 899 imaged neurons; data not shown; p > 0.05). The calcium response probability was not significantly affected by SYM 2081 in EGFP − neurons from all hippocampal layers (98 ± 5% of control in the presence of SYM 2081; n = 899 neurons; p > 0.05) (Fig. 4). In contrast, in the EGFP + cell population, calcium response probability was significantly decreased in the presence of SYM 2081 to 53 ± 14% of control after SYM 2081; n = 11 neurons; p < 0.05) (Fig. 4). This further confirmed that approximately half of the EGFP + cell population responds to the TS through the activation of KA-Rs (KA-responsive EGFP + cells: EGFP + neurons with >80% calcium response probability in GYKI 52466 and <20% probability in SYM 2081, i.e., 51% of EGFP + cells), whereas the other half responds through AMPA-Rs (AMPA-responsive EGFP + cells: EGFP + neurons with <20% calcium response probability in GYKI 52466 and >80% probability in SYM 2081, i.e., 41% of EGFP + cells; n = 25).

### Table 1. Morphological distribution of EGFP + and EGFP − CA1 stratum oriens interneurons

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<tr>
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<th>Wild type</th>
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<tr>
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<td>46</td>
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O-Tri, Stratum oriens interneurons with a trilaminar axonal arborization; O-0, stratum oriens interneurons with an axon innervating only the stratum oriens.

### Table 2. Properties of mEPSCs, mEPSC_{KA}, mEPSC_{AMPA}, and fraction of mEPSC_{KA} in CA1 stratum oriens interneurons

<table>
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97 × 103 mm (600 × 600 dpi).
To further confirm that 10 Hz stimulation of the alveus specifically triggered a suprathreshold KA-R-mediated synaptic response in a subpopulation of somatostatin-containing neurons, we performed current-clamp recordings in EGFP⁺ neurons. In the presence of AMPA, NMDA, and GABA_A receptor antagonists, an EPSP_KA could be evoked by TS in a majority of EGFP⁺ cells (71%) with an amplitude decreased to 37 ± 10% of control values (n = 14 EGFP⁺ cells). Such EPSPs could lead to action potential firing with a latency comparable with that obtained in cell-attached recordings (10.1 ± 2.3 ms; n = 5 cells) (see Fig. 7). We specifically performed cell-attached recordings to quantify KA-R-mediated action potential probability in more reliable experimental conditions. A KA-R-mediated action potential firing could be evoked in approximately half of EGFP⁺ neurons (40%), and action potential probability was 68 ± 21% (n = 10 EGFP⁺ cells) (Fig. 5). Both current-clamp and cell-attached responses were blocked by further addition of CNQX (50 μM), validating that they were mediated by KA-Rs. To further confirm the role of KA-Rs in evoked glutamatergic responses in EGFP⁺ neurons, we next performed current-clamp and cell-attached recordings in these cells in the presence of the KA-R functional antagonist SYM 2081 (see Materials and Methods). Current-clamp recordings in EGFP⁺ cells revealed that application of SYM 2081 reduced the amplitude of the evoked EPSP to 44 ± 11% of control (n = 12; p < 0.05). The effect of SYM 2081 on EPSP amplitude was dependent on the EGFP⁺ cell type as revealed by post hoc morphological identification of recorded neurons. Indeed, application of SYM 2081 completely abolished the evoked EPSP in EGFP⁺/O-LM neurons (to 10 ± 10% of control; n = 5) (see Fig. 7), whereas the response was relatively preserved (to 73 ± 9% of control; n = 7; p < 0.01) (see Fig. 7) in EGFP⁺/non-O-LM neurons such as O-S/BP interneurons. Remaining responses were fully blocked by further addition of GYKI 52466 (100 μM), D-APV (40 μM), and bicuculline (10 μM). Finally, application of SYM 2081 abolished evoked action potential firing recorded in current-clamp mode in 66% of EGFP⁺ neurons (n = 3 EGFP⁺ cells) (see Fig. 7).

We conclude that KA-R activation selectively provides a subpopulation of EGFP⁺ cells that includes O-LM cells, with the property to follow reliably theta stimulation protocols. Finally, to further

**Figure 3.** Differential contribution of KA-Rs to glutamatergic transmission in different types of stratum oriens interneurons. A1, Left, Neurolucida three-dimensional reconstruction of a biocytin-filled EGFP⁺/O-LM (green). The axon is indicated in color, o, Stratum oriens; p, stratum pyramidale; r, stratum radiatum; lm, stratum lacunosum moleculare. Scale bar, 100 μm. Inset, Averaged mEPSCs from an O-LM cell (color trace, control conditions, i.e., in the presence of TTX at 1 μM). Note that plots overlap in the O-LM cell. Right, Cumulative probability plot of the distribution of the averaged fraction of mEPSCs from an O-LM cell (color trace, control conditions, i.e., in the presence of TTX at 1 μM). Note that plots overlap in the O-LM cell. B, Cumulative probability plots of the distribution of the averaged fraction of mEPSCs from an O-LM cell (color trace, control conditions, i.e., in the presence of TTX at 1 μM). Note that plots overlap in the O-LM cell. B, Cumulative probability plots of the distribution of the averaged fraction of mEPSCs from an O-LM cell (color trace, control conditions, i.e., in the presence of TTX at 1 μM). Note that plots overlap in the O-LM cell.
determine the cell type specificity of KA-R-responsive cells, we filled cells with biocytin at the end of the imaging experiments and histologically processed the slices. We found that all KA-R-responsive stratum oriens interneurons belonged to the O-LM cell type \((n = 4)\) (Fig. 6), whereas AMPA-R-responsive stratum oriens EGFP\(^+\) interneurons were O-S/BP neurons \((n = 5)\) (Fig. 6). Interestingly, we recorded from a few EGFP\(^+\) cells that followed TS through KA-Rs and found that they all corresponded to O-LM neurons \((n = 3)\). This also confirms that EGFP labels only a subset of O-LM cells as reported previously (Oliva et al., 2000). However, such EGFP\(^+\)/O-LM neurons are likely to constitute a minority of imaged stratum oriens EGFP\(^+\) neurons because the overall response probability to TS of EGFP\(^+\) cells in this region was not significantly different from that in other layers. Finally, we also recorded from EGFP\(^-\) interneurons that did follow TS but not through KA-Rs; these cells were either O-P or O-Bi interneurons \((n = 2)\) (Fig. 6). We conclude that KA-R activation selectively provides O-LM cells with the ability to reliably follow theta stimulation protocols. We next asked for the cellular mechanisms underlying such specificity.

Efficient EPSP\(_{KA}\)–spike coupling in O-LM cells stimulated at theta frequency

Several presynaptic and postsynaptic mechanisms could drive O-LM cells to fire at 10 Hz in response to afferent theta inputs, including, as previously reported, a short-term facilitation of the glutamatergic drive impinging on this cell type (Ali and Thomson, 1998; Maccarroni, 2005) or a summation of slow EPSP\(_{KA}\)s (Castillo et al., 1997; Cossart et al., 1998; Frerking et al., 1998; Frerking and Ohliger-Frerking, 2002).

To discriminate between these hypotheses, because both phenomenon are frequency dependent, we recorded evoked EPSPs in EGFP\(^+\) neurons during a sequence of trains of 10 stimuli at 1, 3, 10, 50, and 100 Hz, repeated once every 30 s. Frequency facilitation was assessed by measuring the amplitude ratio of the second and the last \((10th)\) EPSP relative to the first one \((\text{EPSP}_{2-1} \text{ and } \text{EPSP}_{10-1})\) across all frequencies while EPSP summation was visually inspected. At 10 Hz, the response of EGFP\(^+/\)O-LM neurons was, on average, stable between stimulation trials \((10 \text{ Hz}; \text{EPSP}_{2-1} 130 \pm 19\% ; \text{EPSP}_{10-1} 130 \pm 19\% ; n = 5; p > 0.05)\) (Fig. 7). We found a significant facilitation of the response only for stimulation rates above 50 Hz in EGFP\(^+/\)O-LM cells, whereas short-term facilitation could already be observed at 10 Hz in non-O-LM/EGFP\(^+\) cells, including O-S/BP neurons \((\text{EPSP}_{2-1} 221 \pm 19\%; \text{EPSP}_{10-1} 172 \pm 19\%; n = 4; p < 0.05)\) (Fig. 7).

Accordingly, EPSP\(_{KA}\)s, recorded in the presence of GYKI 52466, neither facilitated nor summated at 10 Hz \((\text{EPSP}_{2-1} 141 \pm 14\% ; \text{EPSP}_{10-1} 221 \pm 19\% ; n = 4; p < 0.05)\). This also confirms that EGFP labeling provides a minority of imaged stratum oriens EGFP\(^+\) neurons because the overall response probability to TS of EGFP\(^+\) cells in this region was not significantly different from that in other layers. Finally, we also recorded from EGFP\(^-\) interneurons that did follow TS but not through KA-Rs; these cells were either O-P or O-Bi interneurons \((n = 2)\) (Fig. 6). We conclude that KA-R activation selectively provides O-LM cells with the ability to reliably follow theta stimulation protocols. We next asked for the cellular mechanisms underlying such specificity.

![Figure 4](image_url) A selective processing of theta stimulation influx via KA-Rs in a subpopulation of EGFP\(^+\) cells. A, Fluorescence traces in response to the TS protocol from representative examples of the two populations of EGFP\(^+\) neurons (green, EGFP\(^+\) I; blue, EGFP\(^+\) II) in bicuculline at 10 \(\mu M\) and D-APV at 40 \(\mu M\) (control), when adding a KA-R antagonist (NBQX at 1 \(\mu M\)) and the AMPA/KA-R antagonist (CNQX at 50 \(\mu M\)); arrows mark the stimuli. Note that the EGFP\(^+\) I neuron (green) responds to all stimuli in the presence of the AMPA-R antagonist, whereas the EGFP\(^+\) II neuron (blue) responds only to the first stimulus. B, Same as in A but in a EGFP\(^+\) I, II, and EGFP\(^-\) interneuron (black), in drug-free ACSF (control), when adding a KA-R antagonist (SYM 2081 at 10 \(\mu M\)) and the AMPA/KA-R antagonist (CNQX at 50 \(\mu M\)). Note that the response in the EGFP\(^+\) I neuron (green) was significantly reduced in the presence of SYM 2081, although it was not significantly affected in both other cell types. C, Bar histograms show the averaged changes in calcium response probability to the 10 Hz stimulation relative to control conditions (D-APV at 40 \(\mu M\) and bicuculline at 10 \(\mu M\)), in KA-R-mediated conditions (i.e., in the presence of GYKI52466 at 100 \(\mu M\) or NBQX at 1 \(\mu M\); filled bars), and in the presence of a KA-R antagonist (SYM 2081 at 10 \(\mu M\); open bars) in EGFP\(^-\) (black) and EGFP\(^+\) (green) cells. *p < 0.01 when comparing relative change with control and between populations. D1, Graph illustrates the averaged amplitudes of the calcium responses evoked by nine successive TS protocols as a function of time (i.e., TS trial number) in KA-R-responsive EGFP\(^+\) cells (EGFP\(^+\) I, green) and in AMPA-R-responsive EGFP\(^-\) neurons (EGFP\(^+\) II, blue). Error bars indicate SEMs. D2, Representative examples of the calcium responses evoked by the first and the fourth TS trial in an O-LM cell (green) and an O-S/BP cell (blue). Arrows indicate TS time.
across time for KA-responsive EGFP^+ neurons than for AMPA-responsive EGFP^+ neurons.

Because modeling studies predict that the frequency of network oscillations depends in part on the time course of synaptic currents in interneurons (Traub et al., 1996; Wang and Buzsáki, 1996; Fuchs et al., 2001), we next hypothesized that the specific EPSP_{\alpha} kinetics could partly provide the O-LM interneuron type with their selective activation at theta. In keeping with this hypothesis, the decay time constants of averaged EPSPs in O-LM cells were indeed slower (40 ± 9 ms; n = 5; p < 0.05) than those in EGFP^+/non-O-LM cells (22 ± 3 ms; n = 8; p < 0.05), whereas input resistance was comparable in both cell types (139 ± 11 and 145 ± 10 MΩ, respectively; p > 0.1). There was no statistical difference in the evoked EPSP decay time between control conditions and after adding GYKI 52466 in O-LM neurons (38 ± 7 ms in GYKI 52466; n = 9 cells; p > 0.05) (Fig. 7). We indirectly tested the hypothesis that slower EPSPs could drive any cell type to fire at theta using a compound (CX546 at 200 μM) that, as reported previously (Pouille and Scanziani, 2004; Xia and Arai, 2005), slows the deactivation and desensitization of AMPA-Rs. Responses to TS were monitored at the network level using calcium imaging and at the cell level using current-clamp recordings. CA1 was surgically isolated from CA3 to avoid contamination from polysynaptic events that could be generated in the latter region, and control conditions were recorded in the presence of d-APV (40 μM) only in the ACSF. We confirmed that CX546 application significantly increased the decay time constant of averaged evoked EPSPs in CA1 pyramidal cells and stratum oriens interneurons (to 166 ± 19 and 208 ± 51% of control values, respectively; n = 5 pyramidal cells and 7 interneurons; p < 0.05) (Fig. 8). Current-clamp recordings at resting membrane potential in the presence of CX546 showed that slower EPSPs tended to summate (Fig. 8). This could increase the probability of action potential firing because 48 ± 13 and 49 ± 17% of TS trains could evoke at least one spike in the presence of CX546 in pyramidal cells (n = 5) and interneurons (n = 7), respectively, compared with 6 ± 6 and 10 ± 5% trains in control. However, firing was not reliably evoked by each stimulation (as measured in O-LM cells) because spike probability per train was 13 ± 4 and 30 ± 20% in the presence of CX546 in pyramidal cells (n = 5) and interneurons (n = 7) compared with 1 ± 2 and 2 ± 1% in control conditions. In keeping with this observation, we observed a small but not significant increase in the amplitude of the evoked calcium response to the TS in the presence of CX546 (to 125 ± 12% of control values; n = 7 movies, 1245 cells; p > 0.5, Mann–Whitney test) but no change in the fraction of cells displaying a 100% calcium response probability (18 ± 3% in control vs 16 ± 4% in CX546; n = 7 movies, 1245 cells; p > 0.5, Mann–Whitney test) (Fig. 8A). We conclude that EPSP kinetics alone cannot account for the ability of the O-LM cells to reliably follow theta stimulation and that intrinsic properties specific to this particular cell type (Maccalferri and McBain, 1996; Martina et al., 2000; Pike et al., 2000; Saraga et al., 2003; Gloveli et al., 2005; Maccalferri, 2005; Lawrence et al., 2006a,b,c) are likely to be involved in the reliable EPSP_{\alpha}–spike coupling at 10 Hz. Future studies are required to investigate the nature of this coupling.

In summary, these results show both at the cellular and network level a specific and reliable processing of theta glutamatergic afferent inputs by KA-Rs in O-LM interneurons.

Discussion

Using fast two-photon imaging to analyze electrically evoked calcium dynamics from hundreds of cells, we show that O-LM cells

EPSP_{10\%-", 122 ± 14%; n = 7). Therefore, KA-Rs mediate a constant and reliable postsynaptic response when activated at 10 Hz in O-LM cells without significant facilitation or summation, which are observed at higher stimulation rates (Fig. 7). Interestingly, because O-S/BP neurons followed TS through a facilitating AMPA-R-mediated response, we hypothesized that these properties would not enable this cell type to fire in a steady way across different trains of stimulation. To test this hypothesis, we compared the stability of the calcium response between successive TS trials in KA-responsive and AMPA-responsive EGFP^+ neurons and found that the amplitude of the evoked calcium event progressively declined in AMPA- but not KA-responsive EGFP^+ neurons (Fig. 4D). The decrease started being significant for the third trial. This indicated that the response to TS was more stable across time for KA-responsive EGFP^+ neurons than for AMPA-responsive EGFP^+ neurons.

Figure 5. Targeted cell-attached recordings show that EGFP^+ cells can respond to theta stimulation through KA-Rs. A, Cell-attached recordings from an EGFP^+ cell in response to 10 alveus stimulations at 10 Hz (open arrows) in the presence of bicuculline (Bic; 10 μM) and d-APV (40 μM; green). Action potentials (large arrows, time-locked spikes) were still evoked when the AMPA-R antagonist GYKI 52466 (100 μM; blue) was added but not in the presence of the KA-R antagonist SYM 2081 (10 μM; pink) or the AMPA/KA-R antagonist CNQX (50 μM; gray). B, Peristimulus time histograms (same recordings as in A; dashed line indicates time of stimulation) constructed from 10 series of 10 alveus stimulations at 10 Hz. C, Bar graphs of averaged spike probability in cell-attached recordings from EGFP^+ cells in control (n = 10), in d-APV plus bicuculline (n = 10), and when adding GYKI 52466 (n = 4) and SYM 2081 (n = 3). Error bars indicate SEMs.
are the only stratum oriens interneuronal subtype that processes glutamatergic inputs exclusively through KA-Rs. Relying on the imaging data, we performed electrophysiological recordings showing unequivocally that EGFP+/O-LM neurons are selectively activated by theta stimuli by means of KA-R-mediated EPSPs. Functionally, we show that this unique postsynaptic property endows O-LM cells with the ability to follow reliably input stimulation at theta frequency.

Somatostatin-containing interneurons are activated by theta frequency stimulation protocols

In imaging experiments, we found that applying a TS protocol to the alveus resulted in the preferential activation of stratum oriens EGFP+ cells as opposed to pyramidal or other interneurons in all layers of the CA1 region. The detected calcium response resulted from the opening of voltage-gated calcium channels attributable to synaptic activation of intracellular calcium accumulation [such as calcium release from intracellular stores (Rozas et al., 2003)]; (2) evoked calcium signals were abolished by blocking synaptic transmission or action potential firing; and (3) the TS protocol evoked action potential firing and EPSPs in EGFP+ cells.

An increase in membrane excitability attributable to the slow afterhyperpolarization currents (I_{AHP}) by synaptic activation of KA-Rs could also contribute to the calcium signal (Cherubini et al., 1990; Melyan et al., 2002; Ruiz et al., 2005). However, such effect is unlikely because (1) it should also involve CA1 pyramidal cells (Ruiz et al., 2005), (2) increased excitability would be abolished within the time window of our TS protocol (Ruiz et al., 2005), (3) current-clamp recordings do not indicate an increase in the firing of interneurons after repeated stimuli trials, and (4) the calcium response amplitude remained unchanged during TS in EGFP+/O-LM cells, indicating that the number of spikes evoked was constant (Smetters et al., 1999). We conclude that the TS protocol used here induced synapse-driven action potential firing in EGFP+ cells.

Stratum oriens somatostatin-containing neurons comprise O-LM (Katona et al., 1999), O-S/BP (Gulyas et al., 2003), and O-Bi (Maccarelli et al., 2000; Martina et al., 2000; Losonczy et al., 2002) neurons. Most biocytin-filled EGFP+ cells in our sample data were O-LM cells (Oliva et al., 2000) but also included O-S/BP interneurons (Maccarelli et al., 2000; Losonczy et al., 2002). The distinction between O-S/BP and O-LM cells was unequivocal because, even with a cut axon, an O-LM cell presents a characteristic spiny dendritic arbor. We have not isolated the type of fibers activated during
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stimulation. However, because the stimulation electrode was placed in the alveus and direct activation of dendrites was excluded, it is likely that Schaffer or CA1 collaterals were stimulated. For O-LM cells, the latter alternative is more probable because these cells are exclusively innervated by CA1 collaterals (Wittner et al., 2006) and because direct antidromic stimulation of CA1 pyramidal cells could be verified while imaging.

Cellular basis for the selective activation of O-LM/EGFP+ interneurons at theta stimulation frequencies

O-LM cells are remarkably tuned to operate at theta frequencies spontaneously (Maccalferri and McBain, 1996), during current injections (Pike et al., 2000), pharmacologically induced oscillations in vitro (Hajos et al., 2004; Gloveli et al., 2005), as well as during local theta oscillations in vivo (Klausberger et al., 2003). We established the uniqueness of this property in the CA1 region by imaging all neurons simultaneously and morphologically identifying reliably responsive cells. Although intrinsic conductances, muscarinic receptor activation, and slow membrane time constants (Maccalferri and McBain, 1996; Ali and Thomson, 1998; Martina et al., 2000; Pike et al., 2000; Losonczy et al., 2002; Saraga et al., 2003; Pouille and Scanziani, 2004; Gloveli et al., 2005; Maccalferri, 2005; Lawrence et al., 2006a,b,c) are also involved in the intrinsic resonance peak of O-LM cells at theta frequencies, the activation of postsynaptic KA-Rs is instrumental to their ability to follow theta synaptic stimulation. First, the selective blockade of KA-Rs by SYM 2081 (Zhou et al., 1997; Li et al., 1999; De Vries, 2000; Coissart et al., 2002; Epstein et al., 2005) specifically blocks the calcium response to TS in O-LM/EGFP+ neurons. Second, TS reliably evoked a GYKI 52466-resistant/CNQX-blocked calcium response in O-LM cells. Third, activation of KA-Rs in EGFP+ neurons drives postsynaptic depolarization and action potential firing; both responses were blocked by SYM 2081 only in O-LM/EGFP+ neurons. Fourth, almost 90% of mEPSCs recorded in O-LM cells were KA-R mediated; voltage-clamp experiments provide a quantification of the involvement of KA-Rs in the absence of pharmacological manipulation. It is difficult to isolate a KA-R-mediated process based solely on pharmacological agents because specific antagonists for AMPA-Rs also partially block KA-Rs (Castillo et al., 1997), whereas the functional KA-R antagonist SYM 2081 works in a use-dependent manner (De Vries, 2000). Finally, current-clamp recordings showed that blocking AMPA-Rs did not affect the decay of the EPSPs in O-LM cells.

Figure 7. Frequency- and cell-type-dependent plasticity of the evoked EPSP in EGFP+ neurons. A1, Neuroluica reconstructions of two typical EGFP-positive neurons, one O-LM cell (left, green axon) and one O-S/BP interneuron (right, axon in blue). A2, Scale bar, 100 μm. A3, Current-clamp recordings at resting membrane potential (V = 0 pA), from the two cell types illustrated in A1, showing the EPSPs evoked by 10 successive electrical stimuli (×, 10 Hz) in control conditions (i.e., in D-APV at 40 μM and bicuculline at 10 μM) and in the presence of the functional antagonist for KA-Rs (SYM 2081 at 10 μM). Note that the EPSPs evoked in control in the EGFP+/O-LM cell were suprathreshold (arrow indicates an action potential), whereas spikes could not be evoked in the illustrated O-S/BP neuron. A3, Evoked EPSPs are blocked in the presence of SYM 2081 in EGFP+/O-LM cells (n = 5, green bar; * p < 0.05), indicating that they are KA-R mediated, whereas they are only partially affected in non-O-LM/EGFP+ cells (n = 5). EPSP amplitude is measured relative to control values. Error bars indicate SEMs. A4, Latency histogram of the spikes evoked in current clamp in the EGFP+/O-LM cell illustrated in A3. B1, Example of an evoked EPSP recorded in current clamp (V = 0 pA) in an O-LM/EGFP+ cell in the presence of D-APV (40 μM) and bicuculline (10 μM; green) and when adding GYKI 52466 (100 μM; dashed orange). Both traces overlap, indicating that the evoked EPSP in control conditions was KA-R mediated (EPSP KA). The EPSP KA was blocked after addition of CNQX (50 μM; black). B2, EPSP KA are not plastic for repetitive 10 Hz stimulation (left traces) but show a significant summation/facilitation for shorter stimulation intervals (50 Hz; right), as illustrated by the current-clamp recordings in the presence of AMPA-R, NMDA-R, and GABA-A antagonists (GYKI 52466 at 100 μM, D-APV at 40 μM, and bicuculline at 10 μM). * indicates the time of stimulation. C1, Graph illustrates paired-pulse ratio (fraction of the EPSP amplitudes evoked by 2 successive stimuli) as a function of stimulation frequency in EGFP+/O-LM cells (green) and in EGFP+/non-O-LM cells (blue). * p < 0.05, significant facilitation of the response. C2, Representative current-clamp recordings from EPSPs evoked by 10 Hz stimulation in the two EGFP+ populations mentioned in C1. Note the difference in EPSP kinetics in EGFP+/O-LM cells and non-O-LM cells. The open arrow indicates time of stimulation.
Therefore, we establish that KA-Rs transmit theta inputs selectively to O-LM cells. An almost exclusive processing of glutamatergic inputs through KA-Rs is in agreement with our previous study in rat slices (Cossart et al., 2002). Glutamate uncaging experiments have also established that most O-LM cells show KA-R-mediated currents, with a small minority expressing responses restricted to a few dendritic spots (Yang et al., 2006, 2007). AMPA-Rs also contributed to the uncaging response, but such results are difficult to interpret attributable to the likely contribution of extrasynaptic receptors under these experimental conditions. Unfortunately, most studies examining synaptic transmission onto O-LM cells did not differentiate between AMPA-R and KA-R contributions (Ali and Thomson, 1998; Losonczy et al., 2002; Pouille and Scanziani, 2004; Biro et al., 2005; Biro and Nusser, 2005) but reported slower EPSP kinetics in these cells (Ali and Thomson, 1998; Losonczy et al., 2002; Pouille and Scanziani, 2004).

We show that KA-Rs not only convey but also partly mediate the reliable activation of O-LM cells within this frequency range. Indeed, short-term plasticity of the response in O-LM cells is excluded because facilitation of the evoked EPSP was not observed between 1 and 10 Hz. Summation of the EPSP was not recorded either given the relatively fast kinetics of evoked EPSP$_{KA}$ in O-LM cells compared with the stimulation interval and with the slower kinetics of EPSP$_{KA}$ in other cell types such as CA3 pyramidal neurons (∼100 ms) (Castillo et al., 1997; Cossart et al., 2002; Frerking and Ohliger-Frerking, 2002). Facilitating/summing EPSCs in O-LM cells have been reported for stimulus intervals shorter than 100 ms, in agreement with our findings (Ali and Thomson, 1998; Losonczy et al., 2002; Pouille and Scanziani, 2004). Our results indicate that summation of glutamatergic inputs does not translate into reliable EPSP–spike coupling efficiency at theta frequency because prolonging the decay of the EPSP$_{AMPA}$ to mimic an EPSP$_{KA}$ using CX546 (Pouille and Scanziani, 2004) increased firing probability but did not activate non-O-LM/CA1 neurons as reliably as O-LM cells. Therefore, the efficient EPSP$_{KA}$–spike coupling for 10 Hz inputs is specific for the O-LM cell type, and the exact intrinsic membrane properties responsible for such property remain to be determined. The strong sodium channel density expressed by O-LM cells may facilitate KA-R-driven action potential generation (Martina et al., 2000). Furthermore, the combination of theta bandpass resonant properties, significantly longer AHPS, and slower membrane time constants (Gloveli et al., 2005) should also attribute O-LM cells with a greater propensity to fire in the theta frequency range.

In contrast to O-LM cells, calcium imaging and patch-clamp recordings revealed that non-O-LM/EGFP$^+$ neurons, such as O-S/BP cells, were activated by TS stimulation but that such response was not KA-R mediated. Indeed, evoked calcium events and EPSPs were blocked by GYKI 52466 and resistant to SYM 2081, in agreement with the fact that KA-Rs mediated half of the response to TS stimulation protocols (Castillo et al., 1997; Cossart et al., 1998). soma and CA1 stratum radiatum so-

Figure 8. EPSP kinetics alone do not determine the response to theta stimulation protocols. A, Contour maps indicate an example of the distribution of cells responding with 100% calcium response probability to TS protocols in the CA1 region (black filled contours) in control (o-APV at 40 μM; left) and in the presence of CX546 (200 μM; right) to slow the kinetics of EPSP$_{AMPA}$ α. Stratum oriens; p, stratum pyramidale; r, stratum radiatum. Scale bar, 100 μm. The fraction of cells responding reliably to all trains of stimulation is not increased in the presence of CX546. B1, Current-clamp recording at resting membrane potential (V$_{rest}$ of approximately −65 mV) of EPSPs evoked by a TS train in control (o-APV at 40 μM; top, black) and in the presence of CX546 (200 μM; middle, gray) in a CA1 stratum oriens interneuron. Open arrows indicate time of stimulation. Action potential firing (filled arrow, truncated) could be evoked in CX546 conditions. Summation of the EPSPs is clearly visible in the presence of CX546 (baseline level is indicated by dotted lines). Bottom traces are averaged evoked EPSPs during a TS in control (black) and in CX546 (gray) recorded at V$_{rest}$ in a CA1 stratum oriens interneuron. Note that CX546 significantly increased the EPSP decay times. Open arrows indicate stimulation. B2, Same as B1 but in a CA1 pyramidal cell. C, Bar histogram plots averaged values of evoked EPSP decay times, action potential probability per train (AP Prob/TS, i.e., averaged number of action potentials evoked within 1 train of 10 stimuli at 10 Hz), and action potential probability per TS protocol (AP Prob/TS, i.e., fraction of trains within the TS protocol evoking at least 1 action potential), for CA1 stratum oriens interneurons (left; n = 7) and pyramidal cells (right; n = 5) in control (o-APV at 40 μM; black) and when adding CX546 (200 μM; gray). Error bars indicate SEMs. *p < 0.05.
neuronal subtype that reliably processes glutamatergic inputs received with theta frequency primarily through postsynaptic KA-Rs. KA-R-mediated theta band activation in neurons is fully predictive of their O-LM phenotype. Intriguingly, the specific inhibition of glutamate receptor subtype 5-containing KA-Rs, the subtype principally expressed in CA1 stratum oriens interneurons (Cossart et al., 1998), was shown recently to affect hippocampal theta oscillations in freely moving animals (Huxter et al., 2007). This finding strongly suggests that the postsynaptic specificity of O-LM cells described here might be the cellular basis for the generation of theta oscillatory activity in behaving animals.

References


