Intrinsic Cornu Ammonis Area 1 Theta-Nested Gamma Oscillations Induced by Optogenetic Theta Frequency Stimulation

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Gamma oscillations (30–120 Hz) are thought to be important for various cognitive functions, including perception and working memory, and disruption of these oscillations has been implicated in brain disorders, such as schizophrenia and Alzheimer’s disease. The cornu ammonis area 1 (CA1) of the hippocampus receives gamma frequency inputs from upstream regions (cornu ammonis area 3 and medial entorhinal cortex) and generates itself a faster gamma oscillation. The exact nature and origin of the intrinsic CA1 gamma oscillation is still under debate. Here, we expressed channelrhodopsin-2 under the CaMKIIα/H9251 promoter in mice and prepared hippocampal slices to produce a model of intrinsic CA1 gamma oscillations. Sinusoidal optical stimulation of CA1 at theta frequency was found to induce robust theta-nested gamma oscillations with a temporal and spatial profile similar to CA1 gamma in vivo. The results suggest the presence of a single gamma rhythm generator with a frequency range of 65–75 Hz at 32°C. Pharmacological analysis found that the oscillations depended on both AMPA and GABA_A receptors. Cell-attached and whole-cell recordings revealed that excitatory neuron firing slightly preceded interneuron firing within each gamma cycle, suggesting that this intrinsic CA1 gamma oscillation is generated with a pyramidal–interneuron circuit mechanism.

Key words: circuit; hippocampus; interneuron; mouse; network; oscillation

Introduction

The timing of neural cell firing is thought to be essential for successful information coding within the brain. Gamma oscillations, rhythmic activity measured in the local field potential (LFP) with a frequency range of 30–120 Hz, synchronize the spiking activity of neural populations to a specific phase of the gamma wave, thus segregating cell firing into the different gamma cycles (Buzsáki and Wang, 2012). Gamma oscillations often co-occur with theta oscillations (4–12 Hz) to which they are phase-amplitude coupled, such that the gamma oscillations have a larger amplitude at a consistent specific phase of the theta cycle (Buzsáki et al., 1983). This complex temporal control of neural activity is thought to be important for cognitive tasks, including navigation, sensory association, and working memory. Disruption of gamma oscillations has been implicated in a diverse range of diseases, such as schizophrenia (Uhlhaas and Singer, 2010) and Alzheimer’s disease (Goutagny and Krantic, 2013). It is therefore important to understand the mechanisms underlying gamma oscillations.

The cornu ammonis area 1 (CA1) of the hippocampus exhibits multiple types of gamma oscillations in vivo, which can be segregated based on their spectral, temporal, and spatial profiles. A gamma oscillation in the range of 30–80 Hz originates in cornu ammonis
area 3 (CA3) and propagates along the Schaffer collaterals to the stratum radiatum (SR) of the CA1 (Ciscisvari et al., 2003). In the medial entorhinal cortex (mEC), a faster gamma oscillation is generated in the range of 60–120 Hz and propagates to the terminal region of mEC projections in the CA1, the stratum lacunosum-moleculare (SLM; Colgin et al., 2009). Both of these afferent gamma oscillations can also be recorded from the stratum pyramidale (SP), the location of the cell bodies of CA1 pyramidal neurons, as well as from their respective input layers. Using independent component analysis, a third gamma oscillation generated in the SP was recently identified in the CA1 (Schomberg et al., 2014). This locally generated gamma oscillation has a similar frequency (>100 Hz) to mEC gamma, but it has a distinct temporal and spatial profile, suggesting the presence of a gamma generator in the CA1 that is independent of the afferent gamma generators.

In vitro preparations have been put to good use to reveal the mechanisms underlying the current generators for hippocampal oscillations (for review, see Butler and Paulsen, 2015). CA3 gamma oscillations can be induced in acute hippocampal slices by application of the cholinergenic agonist carbachol (Fisahn et al., 1998) and mEC gamma oscillations have been induced by optogenetic stimulation of stellate cells in the mEC (Pastoll et al., 2013). In the CA1, strong excitation by tetanic electrical stimulation (Whittington et al., 1995) or high concentrations of kainate (Traub et al., 2003) can give rise to local CA1 gamma. More recently, it was found that when the connections between CA1 and CA3 are severed, carbachol induces CA1 gamma oscillations (Pietersen et al., 2014). However, CA1 gamma oscillations in more physiological conditions, such as in the presence of theta oscillations, have not yet been reported in the intact hippocampal slice.

Here, we asked whether sinusoidally modulated theta-frequency optogenetic stimulation of the CA1 is sufficient to generate gamma oscillations in acute hippocampal slices. Expressing the optogenetic activator channelrhodopsin-2 (ChR2) under the CaMKIIα promoter in excitatory neurons granted us high temporal and spatial control over the pyramidal cells of the CA1. We found that sinusoidally modulated stimulation in the theta frequency range induced robust gamma oscillations, which were phase-amplitude coupled to the theta stimulation. Both the frequency and theta coupling were analogous to those seen for CA1 gamma oscillations in vivo. The oscillations were resilient to changes in the input to the CA1 region and pharmacological analysis found that the oscillations depended on both AMPA and GABA<sub>α</sub> receptors. Cell-attached and whole-cell recordings revealed that excitatory neuron firing slightly preceded interneuron firing within each gamma cycle, confirming a pyramidal-interneuron circuit mechanism of gamma generation.

Materials and Methods

Mice. Transgenic mice expressing the optogenetic activator ChR2 under the CaMKIIα promoter were created by crossing LoxP-ChR2(H134R)-EYFP<sup>7/4</sup> mice on a 129S6 background (Jackson Laboratories, stock #012569; http://jaxmice.jax.org/strain/012569.html) with CaMKIIα-Cre<sup>7/4</sup> mice on a C57BL/6 background (Jackson Laboratories, stock #005359; http://jaxmice.jax.org/strain/005359.html). This resulted in heterozygous mice expressing both CaMKIIα-Cre and LoxP-ChR2(H134R)-EYFP, thus expressing ChR2 selectively in CaMKIIα cells, including pyramidal cells in the CA1 of the hippocampus. All animal care and experimental procedures were conducted in accordance with the United Kingdom Animals (Scientific Procedures) Act of 1986. All mice used were between 4 and 8 weeks old, and both genders were used.

In vivo recordings. Adult mice of the C57BL/6 strain (n = 4) were anesthetized with 2 g/kg urethane (Sigma-Aldrich; diluted 20% w/v in saline) and supplemented with 0.2 g/kg urethane as needed (approximately twice per 4 h session). Mice were then placed in a stereotaxic frame and the skin and skull above the hippocampus removed. An extracellular recording electrode was then slowly inserted into the CA1 SP of the right hippocampus. Tail pinches were used to induce activity and recordings taken immediately after tail pinching were used for analysis. The recording site was confirmed post hoc by passing a 3 mA current for 2 s through the recording electrode, making acute slices from the brain, and then identifying the lesion site under a microscope.

Slice preparation. The transgenic mice were deeply anesthetized with 4% (v/v) isoflurane at 1.4 L·min<sup>−1</sup> and decapitated. The brain was then quickly removed and horizontal (unless otherwise stated) hippocampal slices with a thickness of 350–400 μm were prepared in cold (0–3°C), oxygenated (95% O<sub>2</sub>, 5% CO<sub>2</sub>) artificial CSF (aCSF) containing the following: 126 mM NaCl, 3 mM KCl, 1.25 mM NaH<sub>2</sub>PO<sub>4</sub>, 2 mM MgSO<sub>4</sub>, 2 mM CaCl<sub>2</sub>, 26.4 mM NaHCO<sub>3</sub>, and 10 mM glucose, pH 7.2. Slices were stored in a submerged-style storage chamber at room temperature (22–24°C) for ≥50 min before use. For minislice preparation, slices were gently removed from the recording chamber and a cut was made under a microscope. The slice was then remounted in the recording chamber and left to recover for 10 min.

Multielectrode recordings. Multielectrode probes were coated in poly-D-lysine for ≥1 min to aid with slice adhesion and then rinsed with aCSF. Probes contained 64 electrodes 50 μm in size, structured in an even 8 × 8 grid with an interpolateral distance of 150 μm (Panasonic MED-P155, Titan Biosciences). Slices were mounted to incorporate as much of the CA1 as possible. Slices were then transferred to the recording unit, where they were maintained in a submerged condition, continually superfused with aCSF at a rate of 3–4 ml·min<sup>−1</sup> and heated to 27–29°C. The slices were left for ≥10 min before recording commenced at a 20 kHz acquisition rate using the Panasonic MED64 system (Titan Biosciences).

Patch-clamp recordings. Single-cell recordings were performed in slices, using a Multiclamp 700B amplifier and CV-7B headstages (Molecular Devices). Signals were acquired at 20 kHz (4 kHz Bessel filter) with custom Matlab software controlling a 16-bit National Instruments analog board. Slices were superfused with aCSF at 27–29°C. Before patching, a borsalicic glass electrode (3–5 MΩ) filled with 10 MΩ HEPES-buffered aCSF for LFP recording was positioned in the SP close to the cell that was patched. All pyramidal neurons recorded from were in the SP and all interneurons from which recordings were made were in either the SP or the stratum oriens (SO).

Three different patch-clamp configurations were used to verify the input and output of individual cells during theta-nested gamma oscillations. To examine the output of cells, spikes from single neurons were recorded in cell-attached voltage-clamp mode followed by whole-cell current-clamp mode. First, a borsalicic glass pipette (3–5 MΩ) filled with 10 MΩ HEPES-buffered aCSF at 27–29°C. The slices were left for ≥10 min before recording commenced at a 20 kHz acquisition rate using the Panasonic MED64 system (Titan Biosciences).

To examine the input to cells, IPSCs and EPSCs were recorded in whole-cell voltage-clamp mode with the following intracellular solution: 105 mM K gluconate, 30 mM KCl, 10 mM HEPES, 1 mM EGTA, 10 mM phosphocreatine Na<sub>2</sub>, 4 mM ATP Mg, and 0.3 mM GTP Na (−10 mV liquid junction potential correction). To examine the input to cells, IPSCs and EPSCs were recorded in whole-cell voltage-clamp mode with the following intracellular solution: 130 mM Cs methanesulfonate, 3 mM CsCl, 10 mM HEPES, 1 mM EGTA, 10 mM phosphocreatine Na<sub>2</sub>, 4 mM ATP Mg, 0.3 mM GTP Na, and 5 mM QX314, pH 7.2 (balanced with CsOH; −8 mV liquid junction potential correction). During optical stimulation, IPSCs were recorded at −10 mV and EPSCs at −70 mV. Series resistance was always <35 MΩ and compensated up to 90%.

Neurobiotin (0.5%) was also added to the intracellular solution. Neurobiotin allowed cells to be stained by Alexa Fluor 633 streptavidin (Life Technologies) and have their morphologies confirmed. The cells were left in whole-cell mode for ≥15 min to allow a sufficient amount of neurobiotin to enter the cell. After a successful cell recording had taken place, the slice was transferred to a 4% paraformaldehyde solution and left overnight. The next day, the slice was thoroughly washed in PBS solution and then left in Alexa
Fluo 633 streptavidin (Life Technologies) for 3 h. The slice was then washed once again using PBS solution and fixed on a microscope slide. The slides were then left in darkness at 4°C for ≥24 h before being imaged.

**Imaging.** A Leica SP2 confocal microscope was used to image the cells that had been recorded. The following fluorescence excitation and emission wavelengths were used to visualize the slices: DAPI, 405 and 420–460 nm; enhanced yellow fluorescent protein (eYFP), 514 and 530–560 nm; Alexa Fluor 633, 633 and 640–780 nm. Cells were first visualized at 10× resolution and this was then increased to 63× if a putative interneuron was identified for further confirmation of the cell type. For each cell, a minimum 2-μm-resolution z-stack was acquired, which was used for 3D reconstructions with the software Vaa3D (Peng et al., 2010, 2014). Cells were classified as pyramidal neurons when the stereotypical apical dendrite was observed, and interneurons when only local projections were identified.

**Light delivery.** A 473 nm diode-pumped solid-state laser (Ciel, Laser Quantum) or a 470 nm digital micromirror device (Polyгон, Mightyet) was used to excite ChR2. Light was delivered to the top of the slice through the objective of an Olympus BX51 microscope. The light, which was used to excite ChR2. Light was delivered to the top of the slice were identified.

**Data analysis and statistics.** Data were imported into Igor Pro (WaveMetrics) or Matlab (Mathworks) was used to control the light sources, modulating the light sinusoidally at a frequency of 5 Hz and synchronizing the light stimulation with the recordings. In all cases, the maximum light intensity (100%) was <2.5 mW·mm<sup>-2</sup>.

Pharmacology. Stock solutions (1000×) of the drugs were made using the following solvents: dimethyl sulfoxide (DMSO) for the GluK1-subunit-containing kainate receptor antagonist UBP-302, the GABA<sub>A</sub> receptor antagonist (+)-bicuculline, the AMPA and kainate receptor antagonist 2,3-dihydroxy-6-nitro-7-sulfamoyl-benzo[f]quinoline-2,3-dione (NBQX) and the allosteric AMPA receptor modulator cyclotrizide (CTZ); ethanol for the barbiturate (±)-thiopental; 1.1 molar equivalent NaOH for the metabotropic glutamate (mGlu) receptor antagonist (RS)-α-methyl-4-carboxyphenylglycine (MCPG); H<sub>2</sub>O for the muscarinic receptor antagonist atropine, the N-methyl-D-aspartate (NMDA) receptor antagonist N-methyl-α-aspartate (NMDA) receptor antagonist N-methyl-D-aspartate (NMDA) receptor antagonist AP5, and the AMPA receptor antagonist GYKI-53655. These stock solutions were then frozen, defrosted on the morning of the relevant experiment, and diluted 1000× in aCSF. After a 20 min incubation period, a recording of activity was taken during stimulation with blue light. The relevant drug solution was then bath applied and another recording taken 10 min later. All drugs and reagents were purchased from either Tocris Bioscience or Sigma-Aldrich.

**Results**

**Optogenetic stimulation of the CA1 activates a single intrinsic gamma oscillation**

When recording the LFP in the SP from the CA1 of a urethane-anesthetized mouse, a tail pinch consistently induced network activity that persisted for a few seconds afterward (Fig. 1A, top). From the PSD, this network activity consisted of two main components: a large amplitude theta oscillation and a smaller amplitude gamma component spanning the majority of the gamma range (Fig. 1B).

When only the theta cycles containing fast gamma cycles (>50 Hz) were analyzed, the oscillations were phase-locked to the trough of the theta cycle measured in the LFP (Fig. 1C).

By expressing ChR2 in the CA1 and preparing acute hippocampal slices, we were able to induce CA1 gamma oscillations in vitro in response to stimulation with a 2.5 mW·mm<sup>-2</sup> blue light sinusoidally modulated at a frequency of 5 Hz (Fig. 1A, bottom). In contrast to the in vivo condition, these oscillations had a narrow, specific frequency range (52 ± 1 Hz at 27–28°C, n = 106 slices; Fig. 1B), which was consistent between all slices and animals tested. As was the case for in vivo gamma oscillations, the in vitro oscillations were phase–amplitude coupled to the theta stimulation, occurring at the peak of each sinusoidal stimulation cycle (the trough of the theta activity recorded in the LFP; Fig. 1C).

The oscillations were stable over time (Fig. 1D) and persisted for as long as the light stimulation continued, immediately stopping when the light stimulation did (Fig. 1E). The oscillations could be induced using slices prepared using all three different angles tested (horizontal, coronal, and longitudinal), with no difference in the power (Fig. 1F) or frequency (Fig. 1G) between the different slicing angles. When the temperature of the superfusion solution was increased, the in vitro gamma oscillations decreased in power by 78 ± 3% from room temperature (22–24°C) to 32°C (Fig. 1H). The frequency of the oscillations was found to have the opposite relationship with temperature, increasing from 35 ± 1
from an 8 Hz gamma across the CA1, we conducted a CSD analysis. Recording this was presumably due to the high metabolic strain that oscil-

32°C and this was accompanied by a severe reduction in both excitatory and inhibitory currents in CA1 cells (data not shown).

Hz at room temperature (22–24°C) to 62 ± 4 Hz at 32°C (n = 11; Fig. 1). Increasing the intensity of light stimulation failed to revert the changes in frequency, demonstrating this effect was not caused by a temperature-dependent reduction in ChR2 kinetics (Fig. 1). There were no discernible oscillations present above 32°C and this was accompanied by a severe reduction in both excitatory and inhibitory currents in CA1 cells (data not shown). This was presumably due to the high metabolic strain that oscillations place on in vitro slices (Hajos et al., 2009).

To accurately localize the sinks and sources of the optogenetic gamma across the CA1, we conducted a CSD analysis. Recording from an 8 × 8 electrode grid allowed us to visualize activity across the majority of the CA1 during stimulation. The gamma oscillations were present in both the SP and SLM layers, with no activity observed in the intermediary SR layer. The oscillations spread ~400 μm into the nearby areas of the dentate gyrus and subiculum, where they could be detected in the LFP and decreased in magnitude with distance from the CA1 (data not shown). Much smaller theta-nested gamma oscillations could also be detected in the deep layers of the mEC during CA1 stimulation, though it is unclear whether these were due to volume conduction through the extracellular tissue or generated locally by CA1 projections that terminate in these layers of the mEC.

Cycle averages were constructed from 48 theta cycles and both the SP and SLM gamma cycles were still clearly visible after this averaging, demonstrating the strong phase preference between theta cycles of the oscillations (Fig. 2C). The CSD showed large uniform sink and source bands across the transverse axis of the SP and the SLM, demonstrating that the entirety of the CA1 from which recordings were taken participated synchronously in the oscillations. This again suggested that when the CA1 receives theta input, a single gamma generator is activated that entrains all the CA1 lam, where they could be detected in the LFP and decreased in magnitude with distance from the CA1 (data not shown). Much smaller theta-nested gamma oscillations could also be detected in the deep layers of the mEC during CA1 stimulation, though it is unclear whether these were due to volume conduction through the extracellular tissue or generated locally by CA1 projections that terminate in these layers of the mEC.
To further ascertain that these oscillations were intrinsic to the CA1 and independent of afferent activity, CA1 minislices were prepared from acute hippocampal slices and the oscillations were recorded from both the SP and SLM layers simultaneously (Fig. 3A). A cut between the border of the CA1 and the subiculum, severing EC–CA1 connections, caused no significant change in the power (43 ± 19% increase, $p = 0.0658$, $n = 18$; Fig. 3B, C) or the frequency (55 ± 1 Hz compared with 58 ± 4 Hz before and after the cut respectively, $p = 0.4907$; Fig. 3B, D) of the oscillations recorded from the SP. The oscillations recorded concurrently from the SLM layer also demonstrated no significant change in the power (6 ± 13% increase, $p = 0.9698$, $n = 17$; Fig. 3B, C) or the frequency (54 ± 1 Hz compared with 55 ± 1 Hz before and after the cut respectively, $p = 0.1357$, $n = 17$; Fig. 3B, D) when this cut was made. As there is also a gamma generator in the CA3, a second cut was then made along the CA1–CA3 border. Again, this caused no significant change in the power (18 ± 9% increase, $p = 0.4641$, $n = 10$; Fig. 3B, C) or frequency of the oscillations (56 ± 1 Hz, $p = 0.6049$; Fig. 3B, D) recorded from the SP. This was also true of the power (19 ± 9% decrease, $p = 0.0328$, $n = 15$; Fig. 3B, D) and frequency (59 ± 4 Hz, $p = 0.3382$, $n = 15$; Fig. 3C, D) of the gamma oscillations recorded from the SLM after the second cut. These gamma oscillations were therefore independent of activity in the afferent regions.

In our transgenic mice, no eYFP expression was observed in CA3 pyramidal neurons, whereas pyramidal neurons in the EC did express eYFP. Accordingly, the EC projections terminating in the SLM could still be activated by light even after the cortex was severed from the hippocampus. Therefore, using a digital micro-mirror device to focus the light in a narrow band on individual layers of the CA1, we investigated the importance of the different CA1 layers for generating the gamma oscillations (Fig. 3E). The power of the oscillations was highest when the light was focused over the SP layer, with SO and SR stimulation inducing oscillations with a power of 78 ± 5% and 83 ± 4% respectively compared with stimulation of the SP ($p = 0.0019$ and 0.0055 respectively).
respective, \(n = 14\); Fig. 3F, G). Stimulation of the SLM, the location of EC projections, decreased the oscillations further still to 42 ± 6\% \((p = 0.0001, n = 14); \text{Fig. 3F, G}\). Furthermore, these changes in power were also observed in the concurrent oscillations recorded from the SLM (SO stimulation: 87 ± 5\%; SR: 85 ± 2\%; SLM: 44 ± 4\%; \(p = 0.0161, 0.0008, \text{and} \ 0.00001 \text{respectively,} \ n = 12); \text{Fig. 3G}\). There was no change in the frequency of the oscillations across all conditions tested (Fig. 3H). The SP layer was therefore the most important for generation of the gamma oscillations.

Therefore, theta-frequency activation of the CA1 is sufficient to generate gamma oscillations that are intrinsic to the CA1, which we refer to as CA1 gamma. Our next step was to examine the relationship between the theta input and the resulting gamma oscillations that are generated.

CA1 gamma is resilient to changes in the frequency and intensity of theta input

Using optogenetics to elicit gamma oscillations afforded us temporal control over the input to the CA1. We used this to ask whether different stimulation patterns could influence the properties of the resulting gamma oscillations.

During movement, the frequencies of both hippocampal theta oscillations (McFarland et al., 1975; Sławińska and Kasicki, 1998) and gamma oscillations (Ahmed and Mehta, 2012) increase with the running speed of the animal. Therefore, we asked whether a change in theta frequency influences the resulting gamma oscillations. As stimulation frequency was increased from 1 to 10 Hz, the power of CA1 gamma was highest in the middle of the frequency range, at 5 Hz stimulation (Fig. 4A). There was an almost linear decrease either side of this frequency, and the power decreased to 81 ± 7\% at 1 Hz stimulation frequency and to 68 ± 11\% at 10 Hz stimulation frequency (Fig. 4B; \(n = 11\)). Outside of this frequency range, gamma oscillations could still be induced across all stimulation frequencies tested (0.25–30 Hz), but the power and robustness of the oscillations were dramatically reduced. The frequency of CA1 gamma remained relatively constant throughout the different stimulation frequencies, increasing slightly with stimulation frequency, but by ≤10 Hz (Fig. 4C). Therefore, there is an optimum theta frequency range within which the CA1 oscillates with the strongest gamma power, and this range coincides with the middle of the theta range.

We then investigated the effect of the intensity of theta stimulation on CA1 (Fig. 4D). The power of the gamma oscillations measured from the SP increased with stimulation intensity between 20 and 80\% of maximum light intensity (1.9–2.1 mW·mm\(^{-2}\)), and plateaued between 80 and 100\% intensity (20\% compared with 80\%, 24 ± 4\%, \(n = 11\); Fig. 4E). This change in power was mirrored by the change in the amount of inward current in pyramidal neurons elicited by ChR2 across the different light intensities (Fig. 4D, E). Despite these large changes in power, the frequency of the gamma oscillations remained essentially constant across all the different light intensities tested (44 ± 1 Hz at 40% light intensity compared with 46 ± 2 Hz at 100% intensity; Fig. 4F).

To analyze the phase locking of the gamma oscillations to the theta stimulation, a CWT was calculated for each stimulation cycle (Fig. 4G). The phase locking of each individual gamma cycle was high within each slice but not between slices, with the gamma in each different slice having slightly different phase preferences relative to the theta stimulation (Fig. 4H). We then compared the relationship between stimulation intensity and theta–gamma phase–phase coupling by calculating the average \(r\) value for the three largest gamma oscillations recorded for each slice under the different stimulation intensities. There was an almost linear increase in the \(r\) value as the power of the oscillations increased.

Figure 3. CA1 gamma is independent of upstream regions. A, Slices were cut along two axes (left) to isolate the CA1 from the cortex (top row) and from both the cortex and CA3 (bottom row). Right, Representative activity from a single slice recorded from the SP for each condition. The bottom blue trace represents the light stimulation. Vertical scale bar, 50 µV; horizontal scale bar, 100 ms. B, PSD plot for the recordings shown in A and before any cut was made to the slice. C, Average change in the power of the gamma oscillations measured from both the SP (black) and SLM (gray), calculated as the area under the curve ≥ 15 Hz from the peak in the gamma range (30–120 Hz), between when slices were intact (left column) to when a cut was made to separate the CA1 from the cortex (middle column; \(n = 18\) for SP and \(n = 17\) for SLM), and when the second cut was made to isolate the CA1 from the CA3 in addition to the cortex (right column; \(n = 16\) for SP; \(n = 15\) for SLM). D, Same as in C but the peak frequency for the different conditions is plotted. For C and D there are no significant differences when the full slice condition was compared with either of the other two conditions using a paired \(t\) test with Bonferroni’s corrections (\(p\) value threshold of 0.025). E, A digital micromirror device was used to precisely deliver light to the slice. This allowed for the light to be concentrated on a thin band spanning a single layer of the CA1 (blue sections). F, Representative traces of activity recorded from the SP when the light stimulation was focused in a narrow band on the SO (left), the SP (second from left), the SR (second from right), and SLM (right). Horizontal scale bar, 100 ms; vertical scale bar, 50 µV. G, Average change in the power of the gamma oscillations measured from both the SP (black) and SLM (gray) calculated in the same manner as in C, depending on the layer that was stimulated with the light. The values are normalized to when the light was focused on the SP layer; \(n = 14\) for SP and \(n = 12\) for SLM recordings. *\(p < 0.0167\), **\(p < 0.0033\), and ***\(p < 0.0003\); paired \(t\) test with Bonferroni’s corrections. H, Same as in G but for the average frequency of the oscillations. There were no significant differences in the frequency between SP stimulation and stimulation of any of the other layers when using a paired \(t\) test with Bonferroni’s corrections (\(p\) value threshold, 0.0167). In C, D, G, and H, error bars represent SEM.
suggesting that theta–gamma phase–phase coupling increases with theta input to the CA1.

To summarize, while the power of CA1 gamma is highly dependent on theta input, the oscillations occupy a specific narrow frequency range that did not shift by more than a few hertz when either the frequency or power of the theta input was varied.

**Pharmacological analysis of CA1 gamma**

It was therefore apparent that optogenetic activation of CA1 consistently activated a single internal gamma generator. We next embarked on a pharmacological characterization of this CA1 gamma generator.

(27 ± 5% increase between 40 and 80% light intensity; Fig. 4I), suggesting that theta–gamma phase–phase coupling increases with theta input to the CA1.

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When slices were left for an incubation period of 10 min, there was a significant decrease in the power of the oscillations over time by 8 ± 2% \( (p = 0.0078, n = 11) \). Therefore, the difference in oscillatory power before and after each drug application was compared with this change. Furthermore, the frequency of CA1 gamma before drug application was found to be different when compared with slices in the control condition, so the change in frequency, rather than the actual values, was used for comparison. Ethanol, DMSO, and NaOH were used as solvents for some
of the reagents, and none of these had a significant effect on the power or the frequency of CA1 gamma (data not shown).

We first sought to whether the CA1 gamma model reported here shares mechanisms with pharmacological models of in vitro gamma oscillations (Fig. 5). Activation of mGlu receptors (mGluRs) causes CA1 gamma oscillations in vitro (Whittington et al., 1995), but mGluRs were not important for optogenetic CA1 gamma, as 500 μM of the mGluR antagonist MCPG had no effect on the power (87 ± 5%, p = 0.3618, n = 11; Fig. 5B) or frequency of the gamma oscillations (98 ± 2%, p = 0.3115; Fig. 5C). Activation of kainate receptors can also induce in vitro CA1 gamma (Fisahn, 1999), but when these receptors were blocked with 20 μM UBP-302, there was no change in the power (97 ± 2%, p = 0.1589, n = 8; Fig. 5B) or frequency of CA1 gamma (100 ± 5%, p = 0.9923; Fig. 5C). It was recently shown that muscarinic receptor activity can induce CA1 gamma oscillations (Pietersen et al., 2014), but these receptors were not involved here, as 10 μM of the muscarinic receptor antagonist atropine had no effect on the power (97 ± 4%, p = 0.2340, n = 7; Fig. 5B) or the frequency (94 ± 2%, p = 0.0222; Fig. 5C) of the CA1 gamma.

Two different models have been proposed for the mechanism of gamma oscillations. The interneuron network gamma (ING) model predicts that recurrently connected interneurons alone are sufficient to generate gamma oscillations, by spiking at their preferred firing rate, which will entrain the entire network (Whittington et al., 1995; Bartos et al., 2007). The second model is the pyramidal–interneuron network gamma (PING) model, whereby pyramidal cell spiking excites interneurons, which then silence pyramidal neurons (as well as other interneurons) via feedback inhibition until the inhibition fades and the pyramidal neurons can fire again, resulting in synchronized rhythmic activity (Fisahn et al., 1998; Whittington et al., 2000). As interneurons are thought to be important for both models, we first tested their involvement in CA1 gamma by blocking GABA_\_A receptor-mediated NMDA receptors was important. At this concentration, GYKI-53655 was then used to identify which of these two excitatory blockers altered the frequency of the gamma oscillations produced (Fig. 7C). Even CTZ, which prolongs EPSCs, had no effect on the power (85 ± 4%, p = 0.0676; Fig. 7B) or the frequency of the oscillations (108 ± 5%, p = 0.0725; Fig. 7C).

We conclude that for CA1 gamma to be generated, both GABA_\_A receptor-mediated inhibition and AMPA receptor-mediated excitation are required. These results suggest that the PING model underpins CA1 gamma. We then sought to confirm this conclusion by conducting single-cell recordings during CA1 gamma.

**Single-cell recordings**

To understand the underlying mechanisms at the cellular level during CA1 gamma, patch-clamp recordings were undertaken from individual CA1 neurons during optical theta stimulation, while theta and gamma oscillations were monitored with an ex-
A tracellular electrode placed in the SP layer of the slice. Cells were filled with neurobiotin and reconstructed post hoc to confirm the cell type (n = 50 of 62), which allowed for identification of pyramidal neurons, with their distinctive main apical dendrite, and the smaller, more compact, interneurons (Fig. 8A). All reconstructed interneurons resided in deep SP and SO and possessed a range of morphologies (Fig. 8A). The firing properties of all cells from which recordings were made were also analyzed. While some interneurons display an irregular spiking pattern (n = 2 of 18), most displayed a fast-spiking pattern (stuttering, n = 2 of 18; continuous, n = 14 of 18). The latter, exhibiting consistently narrow action potentials with strong afterhyperpolarization, is widely regarded to be important for gamma generation (Buzsáki and Wang, 2012). In contrast, all pyramidal neurons showed action potential broadening and spike frequency adaptation (Fig. 8B).

Cell-attached recordings showed that both pyramidal neurons (Fig. 8Ci) and interneurons (Fig. 8Ciii) increased their spiking activity in response to blue light, demonstrating the effect of ChR2 on the cells. To distinguish between direct activation of cells expressing ChR2 and indirect activation via synaptic excitation from upstream ChR2-expressing neurons, we recorded from cells in the presence of 50 µM AP5 and 20 µM NBQX to block all glutamatergic excitation in CA1. This would stop indirect activation occurring but still allow for direct activation. This block had no effect on pyramidal neurons, with all 11 cells recorded in the presence of the blockers spiking in response to light activation in the same manner as cells without blockers (Fig. 8Dii). In contrast, of the 11 interneurons recorded from in the presence of the blockers, 10 did not spike in response to light in the presence of the excitatory blockers (Fig. 8Diii) and only one continued to show a strong response to the light stimulation. It was still possi-
Figure 8. Characterization and ChR2 expression of CA1 neurons. A, 3D reconstruction of the pyramidal neurons (gray, 32 reconstructed of 36 included) and the interneurons used for the pharmacology (blue, 4 of 11), cell-attached (orange, 7 of 7), and voltage-clamp (pink, 7 of 8) single-cell experiments. The highly distinct morphologies and positioning of the two cell types is obvious. In this schematic CA1 illustration, the relative position and size of the cells were based on the images acquired in the confocal microscope. B, The typical membrane potential response observed in response to current steps of increasing amplitude in pyramidal cells (Bi) and fast-spiking interneurons (Bii). Ci, Typical response of pyramidal neurons (Ci) and fast-spiking interneurons (Cii) to optical stimulation with (right) and without (left) the presence of the glutamatergic blockers NBQX (20 μM) and AP5 (50 μM). Di, Average number of spikes that occurred per theta stimulation cycle for pyramidal neurons in control and in pyramidal neurons in the presence of NBQX and AP5. Dii, Average number of spikes that occurred per theta stimulation cycle for interneurons in control and interneurons in the presence of NBQX and AP5. One interneuron retained its response to light in the presence of excitatory blockers, indicating that it expressed ChR2. This interneuron was not included in the group average in Dii. ***p < 0.0005, independent samples t test with Bonferroni’s corrections. n.s., Not significant. Error bars represent SEM. E, An example of the typical response of interneurons in response to blue-light stimulation in the presence of excitatory blockers. Light stimulation still caused a slight membrane depolarization, indicating there was some low-level ChR2 expression.
ble, however, that there was low ChR2 expression in interneurons that could depolarize the cells, just not enough to initiate action potential firing. So the interneurons were also repatched and recordings from them were made in whole-cell current-clamp mode during light stimulation (Fig. 8E). The blue light caused depolarization of the interneurons by an average of 10.0 ± 1.2 mV (n = 10). Thus, ChR2 was expressed at high levels in all pyramidal neurons and to a small degree in interneurons, directly triggering spiking in only a small percentage of interneurons (<10%).

We then examined the relationship between single-cell events and CA1 gamma. We first used cell-attached patch clamp as a minimally invasive method of recording action potentials from the CA1 neurons. To define the phases of the concurrent gamma oscillation, a CWT was constructed for the theta cycle and the flanking 50 ms of activity either side. This was then used to identify the time points of the trough of the cycles before and after each event by using the negative peak in the gamma range (30°–120° Hz) and were used to define 0° and 360° in the calculation of the phase of the selected action potential relative to CA1 gamma. B. The average gamma cycle recorded from each individual slice (gray lines) and the average across all data used in the cell-attached experiments (black line, n = 6443 from 19 slices). C. The mean angle and r value for all pyramidal neurons (black circles) and interneurons (gray squares) from which recordings were made. D. Polar histogram plots (rose plots) to display the phase at which all spikes occurred relative to CA1 gamma for pyramidal neurons (Di; n = 3766 from 12 slices) and interneurons (Dii; n = 2680 from 7 slices). The black arrows are the circular mean and accompanying r value for each group (the axis for these arrows is between 0 and 1 at the bottom and top of the histogram respectively).

Once we had elucidated the cells’ output during CA1 gamma, we next turned to their inputs. Recording in voltage-clamp mode and clamping the cell at −10 mV (the reversal potential of EPSCs) and −70 mV (the reversal potential of IPSCs) enabled us to measure respectively inhibitory and excitatory inputs into the cells during CA1 gamma (Fig. 10A). The IPSC input into all pyramidal neurons tested was significantly nonuniform (p < 0.001 in all cases). Of the eight interneurons tested, all demonstrated IPSCs and EPSCs during CA1 gamma apart from one interneuron that received no detectable EPSCs, and one interneuron that received no detectable IPSCs, and one interneuron that received no detectable EPSCs, and were therefore excluded from further analysis. The phase of IPSC input into the seven interneurons was all strongly nonuniform with respect to CA1 gamma (all p < 0.001 apart from one cell that had p = 0.001). Likewise, the EPSCs into each of the seven interneurons were found to be significantly nonuniform (p < 0.001 in all cases).

IPSCs occurred at the peak of the gamma cycle, and shared the same phase preference regardless of whether they were recorded in interneurons (mean angle, 201°; r = 0.58; n = 2178) or pyramidal neurons (mean angle, 181°; r = 0.56; n = 2407; Fig. 10B). EPSCs in interneurons were strongly phase-locked to the ascending phase of CA1 gamma (mean angle, 82°; r = 0.5; n = 1483). This was consistent across all cells, apart from the EPSCs in one
interneuron, which had a mean angle of 163° ($r = 0.78$, $n = 76$; Fig. 10C). There was therefore a delay between excitatory and inhibitory events that matched the delay between spiking of pyramidal neurons and interneurons.

To assess the contribution of excitatory and inhibitory events to the signal recorded in the LFP, we compared the fluctuations in PSC amplitude to fluctuations in the magnitude of the simultaneously recorded gamma cycle. While there was no correlation between EPSC magnitude and the concurrent gamma cycle magnitude (Fig. 10D), there was a strong correlation between IPSC magnitude and gamma cycle magnitude, which was present for the IPSCs recorded from both pyramidal neurons and interneurons (Fig. 10E). This therefore reinforces the theory that LFP signals comprise mainly inhibitory events (Oren et al., 2010).

These single-cell results suggest the following mechanism by which CA1 is able to intrinsically generate gamma oscillations (Fig. 11). During each gamma cycle (Fig. 11A), pyramidal neurons first fire action potentials at the start of the cycle (Fig. 11B). The resulting EPSCs then arrive in interneurons (Fig. 11C), causing them to produce action potentials (Fig. 11D). This then results in IPSCs reaching both pyramidal neurons and interneurons, resulting in their inhibition (Fig. 11E) and a silent CA1 until the start of the next gamma cycle. This activity will then persist for as long as there is sufficient theta input into the CA1 to maintain

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**Figure 10.** Voltage-clamp recordings during optogenetic CA1 gamma oscillations. A. Representative traces of intracellular currents (top) recorded from pyramidal neurons (left) and interneurons (center, right) while the cells were held at $-10$ mV (left, center) and $-70$ mV (right) to record IPSCs and EPSCs respectively. The concurrent LFP is displayed below each recording. B. Polar histogram plot (rose plot) to display the phase at which all recorded IPSCs in pyramidal neurons (B1; $n = 2111$ from 13 cells) and interneurons (BII; $n = 2179$ from 7 cells), and EPSCs in interneurons (BIII; $n = 1440$ from 7 cells) occurred relative to CA1 gamma. The black arrows are the circular mean and accompanying $r$ value for the group (the axis for these arrows is between 0 and 1 at the bottom and top of the histogram respectively). C. Mean angle and $r$ value for each cell for all the different conditions. D. EPSCs and the size of the concurrent gamma oscillation were normalized to their maximum and linear regression used to calculate a line of best fit for the data from each cell (gray traces) and the average across all cells (black trace; $n = 7$). E. The same as in D but for IPSCs recorded from both pyramidal neurons (gray solid lines; $n = 13$) and interneurons (gray dashed line; $n = 7$) and the average regression across all cells from which recordings were made (black line; $n = 20$).
pyramidal neuron spiking (as evidenced by our experiment modulating the frequency of the theta stimulation).

Discussion

Our three main findings are as follows: (1) optogenetic stimulation of the CA1 activates a single intrinsic gamma generator; (2) the induced oscillation is robust and largely insensitive to changes in the frequency and power of theta input; and (3) CA1 gamma is generated by a PING mechanism.

Optogenetic theta stimulation of the CA1 induces a single intrinsic gamma oscillation

Expression of ChR2 in the CA1 allowed for robust induction of gamma oscillations in response to theta stimulation. There are ≥3 distinct gamma oscillations in CA1 in vivo, distinguished by their anatomical site of generation and classified by their frequency ranges as slow (30–80 Hz), medium (60–120 Hz), and fast gamma (>100 Hz; Schomburg et al., 2014). Recent in vivo recordings found the CA1 not to be entrained to the slow or medium gamma from its afferent regions, but rather CA1 pyramidal neurons were phase-locked to fast gamma, suggesting an intrinsic CA1 gamma oscillation generator (Schomburg et al., 2014). We provide further evidence for an intrinsic CA1 gamma generator here by demonstrating that optogenetic theta stimulation of the CA1 causes a single gamma oscillation of a narrow frequency range that is independent of activity in the afferent regions. This in vitro CA1 gamma has several similarities with CA1 gamma in vivo, including a relatively fast frequency (Belluscio et al., 2012), similar phase-locking of CA1 gamma to the concurrent theta oscillation (Schomburg et al., 2014) and a similar phase delay between pyramidal neuron firing and interneuron firing within each gamma cycle (Senior et al., 2008; Lasztoczi and Klausberger, 2014).

The presence of the alternating sink–source pair within the CA1 shows that the current generator is within the CA1 itself, as reflected by the reversal of the LFP between the SP and SLM. The gamma oscillations could also be recorded from the neighboring dentate gyrus and subiculum, but as these oscillations were of much smaller amplitude and as no phase reversal was observed in these regions, it is likely that this was due to volume conduction from the CA1, rather than due to the activity of any local generator. Indeed, when the light was directed over these neighboring areas, local gamma oscillations with characteristics distinct from those of CA1 gamma were observed, and in this case phase reversal was detected between the different local layers (data not shown). This highlights the potential of the model presented here for comparing gamma oscillations from different regions of the mouse brain in vitro. The ability of the CA1 to generate its own gamma oscillation independent of afferent regions makes it unlikely that gamma oscillations are an interregionally synchronized process, with each packet of information bound to a specific gamma cycle, which carries it along the different steps of the hippocampal circuit. Instead it seems more likely that gamma oscillations are for unidirectional signal propagation, producing local synchrony and delivering each packet to the next downstream region, whereupon a new gamma oscillation from the local generator takes over transmission of the signal. Indeed, the differences in frequencies between the hippocampal gamma oscillations demonstrate that the gamma generators do not need to align their frequency ranges for successful cognitive processing. Having different frequencies would result in asynchrony between the different signals. Perhaps this even enhances processing capability by allowing each area to select input signals by the gamma frequency that carries those signals.

Theta input has only a small effect on CA1 gamma

CA1 gamma was remarkably robust to changes in the frequency and power of theta stimulation (Fig. 4). When the frequency of the theta stimulation was altered, the gamma power was highest for the middle of the theta range. Such theta resonance also exists in vivo (Stark et al., 2013), further supporting the physiological relevance of this model of intrinsic CA1 gamma. When we manipulated the strength of the theta input, the power of the gamma oscillation increased as theta stimulation intensity was increased. Interestingly, both theta power (Chen et al., 2011) and gamma power >60 Hz (Ahmed and Mehta, 2012) was found to increase with running speed in vivo. Therefore, perhaps the increase in gamma power is a simply a consequence of the increase in theta power, rather than being directly caused by the increase in running speed.

Despite dramatically varying both the intensity and the frequency of the theta stimulation, the frequency of the CA1 gamma never strayed by more than a few hertz from the original fre-
quency, indicating that it is a rigid phenomenon. As the frequency for intrinsic CA1 gamma was far higher than for slow gamma as seen in vitro (as well as the cholinergically induced slow gamma in vitro; Fisahn et al., 1998), CA3 activity might be required for slow gamma in CA1 (Csicsvari et al., 2003; Colgin et al., 2009). This finding supports the theory that CA3 interneurons and CA3–CA1 feedforward inhibition are important for CA1 slow gamma (Zemankovicvs et al., 2013). It is likely that, in terms of gamma generation in the CA1, there are ≥2 different circuits consisting of interneuron populations with different inhibitory time constants (which would set the different oscillation frequencies), and in the model presented here, predominantly one such population is being activated. All interneurons tested here were found to participate in CA1 gamma and all had similar phase preferences. Our sampling of interneurons was limited to fast-spiking cells in the deep SP and SO. Therefore it is possible that targeting other interneuron populations may uncover some possessing a different relationship with CA1 gamma.

It has been assumed that one role of gamma oscillations is to synchronize the timing between networks, thus allowing for successful neural communication by providing a rigid temporal framework (Fries, 2005). When theta stimulation intensity was increased, this caused an increase in the phase locking of gamma cycles to their respective phases of the theta cycle, and therefore an increase in the precision and predictability of activity. This increase in the rigidity of the timing of gamma cycles during levels of high theta activity would enable the CA1 to increase its precision during times of high cognitive load, ensuring accurate communication between different cell assembles.

**CA1 gamma is generated by a PING mechanism**

Through the use of both pharmacology and single-cell recordings, we have shown that intrinsic CA1 gamma is generated by a PING mechanism. Although ChR2 expression was not entirely limited to pyramidal neurons, it is unlikely that the small amount of ChR2 expressed in interneurons was important for the generation of CA1 gamma. Furthermore, changes in extracellular potassium levels can induce CA1 gamma oscillations (LeBeau et al., 2002), and we cannot rule out the possibility that extracellular potassium concentration is also important for the gamma oscillations recorded here. The pre-existing models of CA1 gamma are generated by a mixture of PING (Traub et al., 2003; Pietersen et al., 2014) and ING mechanisms (Whittington et al., 1995; Craig and McBain, 2015). Whether CA1 gamma in vitro is generated by a PING or ING mechanism is still not clear, although many studies have reported a delay in firing in CA1 interneurons compared with pyramidal cells consistent with the monosynaptic delay that is characteristic of PING (Bragin et al., 1995; Csicsvari et al., 2003; Senior et al., 2008; Korotkova et al., 2010).

Optogenetic technology has also been used in other in vitro models of gamma oscillations in the CA1’s afferent regions, mEC and CA3. Indeed, Pastoll et al. (2013) demonstrated that optogenetic theta stimulation generated gamma oscillations in the mEC with similar properties to those observed here. This would suggest that the gamma generators present in the mEC and CA1 are analogous. Interestingly, the optogenetic mEC gamma was also phase-locked to the trough of the theta oscillation, and it remains to be seen whether conduction delays would result in propagated mEC gamma occurring at a different theta phase in CA1, as is the case in vitro (Schomburg et al., 2014).

Similarly, optogenetic stimulation of CA3 generated gamma oscillations of a similar frequency to those seen here (Akam et al., 2012), suggesting that another analogous generator is also present in CA3. This is also the case for pharmacological models, with in vitro CA3 (Fisahn et al., 1998) and CA1 gamma (Traub et al., 2003) having a similar frequency and slower than that of optogenetic gamma (~40 Hz). These in vitro similarities make it likely that the different hippocampal gamma generators all share the same basic architecture and each have slight differences, such as the presence of recurrent excitation in CA3, that result in the differences in frequency of the different hippocampal gamma oscillations seen in vivo.

In summary, excitation of CA1 pyramidal neurons is sufficient to generate intrinsic CA1 gamma oscillations in vitro with similar properties to in vivo CA1 gamma oscillations. This model of CA1 gamma oscillations should therefore be useful to further our understanding of the complex interactions of distinct gamma oscillations present in CA1.

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