Gating neuronal activity in the brain
Willems, J.G.P.

Citation for published version (APA):

General rights
It is not permitted to download or to forward/distribute the text or part of it without the consent of the author(s) and/or copyright holder(s), other than for strictly personal, individual use, unless the work is under an open content license (like Creative Commons).

Disclaimer/Complaints regulations
If you believe that digital publication of certain material infringes any of your rights or (privacy) interests, please let the Library know, stating your reasons. In case of a legitimate complaint, the Library will make the material inaccessible and/or remove it from the website. Please Ask the Library: http://uba.uva.nl/en/contact, or a letter to: Library of the University of Amsterdam, Secretariat, Singel 425, 1012 WP Amsterdam, The Netherlands. You will be contacted as soon as possible.
Parvalbumin interneuron mediated feedforward inhibition controls signal output in the deep layers of the perirhinal-entorhinal cortex

Janske G.P. Willems, Wytse J. Wadman, Natalie L. M. Cappaert

Hippocampus (2018); 28 (4): 281-296
DOI: 10.1002/hipo.22830
Abstract

The perirhinal (PER) and lateral entorhinal (LEC) cortex form an anatomical link between the neocortex and the hippocampus. However, neocortical activity is transmitted through the PER and LEC to the hippocampus with a low probability, suggesting the involvement of the inhibitory network. This study explored the role of interneuron mediated inhibition, activated by electrical stimulation in the agranular insular cortex (AiP), in the deep layers of the PER and LEC. Activated synaptic input by AiP stimulation rarely evoked action potentials in the PER-LEC deep layer excitatory principal neurons, most probably because the evoked synaptic response consisted of a small excitatory and large inhibitory conductance. Furthermore, parvalbumin positive (PV) interneurons - a subset of interneurons projecting onto the axo-somatic region of principal neurons - received synaptic input earlier than principal neurons, suggesting recruitment of feedforward inhibition. This synaptic input in PV interneurons evoked varying trains of action potentials, explaining the fast rising, long lasting synaptic inhibition received by deep layer principal neurons. Altogether, the excitatory input from the AiP onto deep layer principal neurons is overruled by strong feedforward inhibition. PV interneurons, with their fast, extensive stimulus-evoked firing, are able to deliver this fast evoked inhibition in principal neurons. This indicates an essential role for PV interneurons in the gating mechanism of the PER-LEC network.

Key words: excitation/inhibition balance, mouse, patch clamp, pyramidal neurons, parahippocampal region
Introduction

The perirhinal (PER) and the lateral entorhinal (LEC) cortex are both involved in processing object information in the so called ‘what’ pathway (Witter et al., 1989; van Strien et al., 2009; Eichenbaum et al., 2012). The PER and LEC receive afferent projections from the agranular insular cortex (AiP) (Burwell, 2000; Mathiasen et al., 2015), a neocortical area involved in emotional, interoceptive and exteroceptive signal processing (Nieuwenhuys, 2012). In turn, PER and LEC axons project to the hippocampal formation (for review see Witter, 1993). Although there are anatomical projections present to convey information from the neocortex, through the PER and LEC, to the hippocampal formation, neuronal activity is not reliably transmitted through this network (Biella et al., 2002; Pelletier et al., 2004; Willems et al., 2016). This suggests that the PER-LEC network, instead of simply acting as a relay station, actively selects and processes information (de Curtis and Paré, 2004).

Still, the neuronal mechanism behind these selecting and processing capabilities is not fully understood. It is shown though, that principal neurons in both the PER and LEC network stop firing up to 300 ms when a cortical input is received (Pelletier et al., 2004). This suppression presumably originates from inhibitory interneurons as reducing the inhibition by partly antagonizing the GABA$_A$ receptor activity resulted in reliable transmission of neocortical synaptic input, implying a role for GABAergic interneurons in controlling relay of activity in the PER-LEC network (Koganezawa et al., 2008; Willems et al., 2016).

Previous studies also showed that a stimulus in the local PER-LEC network evoked inhibitory as well as excitatory responses, whereas a distal stimulus resulted mainly in excitation (Biella et al., 2001; Martina et al., 2001). This suggests that inhibition is mainly recruited in the local circuitry (Unal et al., 2013). An ultrastructural study revealed that the GABAergic neurons are presumably organized in a feed-forward manner (Pinto et al., 2006). The origin of the functional inhibition in the PER-LEC network is still needs to be determined.

Potential candidates for efficient inhibitory control of principal neurons are parvalbumin positive (PV) interneurons (Pfeffer et al., 2013). This interneuron type is present in all layers of the PER and even more abundantly in the LEC (Wouterlood et al., 1995). PV interneurons are known for their high-frequency firing capabilities and they project onto the axo-somatic region of principal neurons. Hence, PV interneurons are capable of strongly regulating
principal neuron output by shaping oscillatory activity (Cunningham et al., 2006; Sohal et al., 2009). Loss of inhibition in the PER-LEC is associated with pathologies involving hyperexcitability such as temporal lobe epilepsy and psychiatric illness (Cunningham et al., 2006; Kumar and Buckmaster, 2006). Furthermore, PV interneuron numbers decrease tremendously in the PER of epileptic rats (Biagini et al., 2013) and PV interneuron activation can terminate epileptic activity in the mouse model for epilepsy (Assaf and Schiller, 2016).

This study investigated whether the interplay between principal neurons and PV interneurons performs a role in processing of synaptic input to the deep layers of the PER-LEC network. We examined the stimulus evoked synaptic input and action potential firing patterns in principal neurons and PV interneurons to address the functional output of the PER-LEC network once synaptic input is processed in the local circuitry.

Materials and Methods

Animals. Experiments were performed on 22 male and female C57Bl/6 mice (Harlan Netherlands BV, Horst) and 18 male and female Pvalb<sup>tm1(cre)Arbr</sup> (Hippenmeyer et al., 2005)/Gt(ROSA)26Sor<sup>tm1(EYFP)Cos</sup> (Srinivas et al., 2001) (PV/YFP) transgenic mice. Experiments to confirm the reversal potential for fast, chloride mediated inhibition were performed on 4 Pvalb<sup>tm1(cre)Arbr</sup>/Gt(ROSA)26Sor<sup>tm32(CAG-COP4*H134R/EYFP)Hze</sup> (Madisen et al., 2012) transgenic mice. All animals were between the ages of P28 and P42. Animal care and experiments were approved by the Animal Care and Use committee of the University of Amsterdam and were in accordance with European guidelines.

Slice preparation. Animals were killed by decapitation, whereafter the brain was rapidly removed and stored in ice-cold modified artificial cerebrospinal fluid (mACSF) containing (in mM): 120 choline chloride, 3.5 KCl, 5 MgSO<sub>4</sub>, 1.25 NaH<sub>2</sub>PO<sub>4</sub>, 0.5 CaCl<sub>2</sub>, 25 NaHCO<sub>3</sub>, 10 D-glucose (pH 7.4, 300 - 315 mOsmol), oxygenated with 95% O<sub>2</sub>/5% CO<sub>2</sub> for at least 30 minutes. Horizontal slices (400 µm thick) containing the neocortical AiP, PER and LEC (Figure 1 e, Willems et al., 2016) were cut in ice-cold mACSF using a VT1200S vibratome (Leica Biosystems, Nussloch, Germany). Functional projections from the AiP to the PER and EC are present in this slice preparation (von Bohlen und Halbach and Albrecht, 2002; Mathiasen et al., 2015; Willems et al., 2016). After sectioning, slices were incubated in ACSF containing (in
mM): 120 NaCl, 3.5 KCl, 1.3 MgSO₄, 1.25 NaH₂PO₄, 2.5 CaCl₂, 25 NaHCO₃, 10 D-glucose, oxygenated with 95% O₂/5% CO₂ (pH 7.4, 300 - 315 mOsmol) at 32°C for 15 minutes, thereafter slices were kept at room temperature until the recording started.

Whole cell recordings in principal neurons. In total 81 principal neurons were recorded in the PER and LEC deep layers. The localization of the PER and LEC in our slice preparation was based on the mouse brain atlas (Paxinos and Franklin, 2001). Patch pipettes were pulled using micropipette puller model P-87 (Sutter Instrument, Novato, CA) and had a resistance of 3-5 MΩ. Whole-cell recordings were performed using an intracellular solution containing (in mM): 131.25 K-gluconate, 8.75 KCl, 10 HEPES, 0.5 EGTA, 4 MgATP, 0.4 Na₂GTP, pH adjusted to 7.4, 295 - 300 mOsmol. 1% biocytin (Sigma-Aldrich, Saint Louis, MO) was added to the intracellular solution for post hoc visualization and morphological identification of the recorded neuron. During the recordings, slices were perfused with ACSF of 30°C at a rate of 2 mL/min. Deep layer PER and LEC principal neurons were selected based on large soma size using a Scientifica SliceScope Pro 6000 (Scientifica, Uckfield, UK). Whole-cell recordings were made using an Axopatch 200B amplifier (Molecular Devices, Sunnyvale, CA), filtered at 10 kHz, sampled at 100 kHz and digitized using a NI DAQ usb-6259 (National Instruments, Austin, TX). Software for data-acquisition was custom made in MatLab (MathWorks, Natick, MA). All voltage signals were corrected online for a -14 mV junction potential. Principal neurons were approached with slight pressure on the pipette and when pressure was released the pipet-cell contact had to reach a seal of 1 GΩ before break in. Immediately after break in, the resting membrane potential was recorded in current clamp at a 0 pA holding current. Access resistance was compensated for at least 50 – 60% and recordings with an access resistance higher than 20 MΩ or with more than 25% change during the recording were discarded.

In some experiments, the glutamatergic transmission was blocked by bath application of 20 µM AMPA receptor antagonist CNQX (Abcam, Cambridge, UK) and 10 µM NMDA receptor antagonist APV (Tocris, Bristol, UK). All other chemicals were obtained from Sigma-Aldrich (Saint-Louis, MO).

Electrical stimulation. For electrical stimulation, a bipolar tungsten stimulus electrode (World Precision Instruments, Sarasota, FL) with a tip separation of 125 µm was placed under visual guidance in the superficial layers of the AiP. A single bi-phasic stimulus pulse (160
µs/phase) was applied using a DS4 bi-phasic current stimulator (Digitimer Ltd, Hertfordshire, UK).

Data analysis. The response latency, defined as the onset of an EPSC, EPSP or an excitatory or inhibitory conductance, was determined as the time difference between the stimulus and the point where the signal exceeded 8 times the baseline standard deviation, within 75 ms after the stimulus was applied. If a response latency was detected the peak and the half width of the response were determined. The peak of the response was characterized as the maximum amplitude after the onset latency and the half width was defined as the time between the point where the response crosses the 50% of the maximum response before and after the peak. The peak and peak time of the action potentials was determined using Matlab (peakdet function), to address the presence and rate of action potential firing.

Decomposition of stimulus evoked synaptic currents. The evoked synaptic response in a neuron contains components that originate from excitatory and inhibitory synapses. As blocking some of these components with pharmaceuticals will affect all responses in the network, we linearly decomposed the current into two underlying components that have a different reversal potential. The post-synaptic cell was clamped at potentials between -90 mV and -50 mV, while evoking the same, voltage-independent, synaptic conductance (see inset in Figure 3 b-c). After subtraction of the stimulus independent background current, this results in a membrane current that contains the excitatory synaptic current and the inhibitory synaptic current:

$$I_m(t) = I_{exc}(t) + I_{inh}(t)$$

These currents are the result of the excitatory and the inhibitory synaptic conductances ($G_{exc}(t)$ and $G_{inh}(t)$) and their respective driving forces, being the differences between membrane voltage $V_m$ and the excitation and inhibition reversal potentials ($E_{exc}$ and $E_{inh}$):

$$I_m(t) = G_{exc}(t)(V_m(t) - E_{exc}) + G_{inh}(t)(V_m(t) - E_{inh})$$

The instantaneous relation between membrane current and membrane can, at each moment in time, be summarized by:

$$I_m = (G_{exc} + G_{inh}) * V_m - (G_{exc} * E_{exc} + G_{inh} * E_{inh})$$

The last equation is the linear I/V relation $I_m = a * V_m + b$, which can be calculated at each moment in time and from which the time varying conductances can now be constructed:
\[ G_{\text{inh}}(t) = (b(t) + a(t) \cdot E_{\text{exc}})/(E_{\text{exc}} - E_{\text{inh}}) \]
\[ G_{\text{exc}}(t) = (a(t) - G_{\text{inh}}(t)) \]

We performed this calculation for 100 ms after the stimulus and with 0.1 ms time resolution. If there are only glutamatergic and GABAergic synapses activated and we have exact knowledge of their (time-invariant) reversal potentials (0 mV respectively -70 mV, Purves et al., 2001; Melzer et al., 2012), \(G_{\text{exc}}\) and \(G_{\text{inh}}\) describe the time course of the synaptic conductances in the cell. The reversal potential of the fast, GABAergic mediated inhibition was verified in slices from transgenic mice expressing the light activated channelrhodopsin (ChR2) specifically in PV interneurons. Optical activation of ChR2 in PV interneurons depolarized the PV interneurons and evoked action potential firing. In this way, we specifically induced PV related IPSCs in the post-synaptic principal cells and calculated the reversal potential of that component, revealing the \(E_{\text{inh}}\) (-70.2 ± 0.4 mV, \(n=30\) IPSCs in 2 principal neurons; data not shown). This value, together with the well-established value of 0 mV for the \(E_{\text{exc}}\) was used for the decomposition. The conductances induced by AiP stimulation were averaged over three repetitions. 

**Paired whole cell recordings of principal neurons and PV interneurons.** PV expressing interneurons in the PER and LEC network were identified using transgenic mice conditionally expressing YFP driven by the PV promotor dependent cre-recombinase expression (Supplementary figure 1 d). YFP was excited at 470 nm using LED illumination light source (PE-100,CoolLed Ltd., Andover, UK) and a 479 ± 40 nm emission filter (Thorlabs Inc., Newton NJ). Paired whole-cell recordings of one PV interneuron and one principal neuron were performed with a maximal inter-soma distance of 200 µm. The firing properties of the cells were recorded by injecting a membrane current that set the membrane voltage from -100 to -30 mV in steps of 5-10 mV. Connectivity between the principal neuron and PV interneuron was tested by evoking action potentials in the principal neuron at reproducible random moments using a frozen noise current injection (Zeldenrust et al., 2013) and recording unitary excitatory post-synaptic currents (uEPSCs) in the PV interneurons clamped at -70 mV (Figure 6 a). We strived to induce a firing rate of 1-2 Hz in the principal neuron. The reversed configuration was used to establish PV to principal neuron connectivity, holding the principal cell at -50 mV in order to record unitary inhibitory post-synaptic currents (uIPSCs) (Figure 6 b).
Next, we addressed the stimulus-evoked synaptic current in voltage clamp (-70 mV) and action potential firing in current clamp in response to AiP stimulation in both principal neuron and PV interneuron. The maximum stimulus intensity was $836 \pm 43 \mu$A, we adjusted the stimulation strength on the response of the principal neuron. Subsequently, evoked synaptic currents were recorded at five holding potentials (-90 to -50 mV) in the principal neuron and PV firing was recorded in current clamp at the same time to compare the estimated evoked inhibitory conductance in the principal neuron to the spiking of PV neurons.

**Histology.** For visualization of the recorded neurons, slices were fixed in 4% PFA in PBS overnight at 4°C after the recording. After 5 washes in PBS (10 minutes each), sections were permeabilized with 0.25% Triton in PBS and biocytin was labeled using Streptavidin-Alexa 488 conjugate (Sigma-Aldrich, Saint Louis, MO) 1:200 diluted in 0.25% Triton in PBS, incubated overnight at 4 °C. After staining, slices were washed in PBS and mounted in Vectashield (Vector Laboratories Inc., Burlingame, Ca). Biocytin filled neurons were visualized using an A1 confocal microscope (Nikon Instruments Europe, Amsterdam, The Netherlands) and their morphology was further examined in ImageJ (Schindelin et al., 2012, 2015).

For the verification of YFP expression specifically in PV interneurons in PV/YFP mice, brains were removed like the above procedure and fixed in 4% PFA in PBS for 8 hours at 4 °C. After fixation, brains were placed in 15% sucrose in PBS for 2 hours and in 30% sucrose in PBS until submerged for cryoprotection. Brains were snap frozen in dry ice and 40 µm cryosections were made and slices were kept free floating in antifreeze at -20 °C. For immunostaining, slices were rinsed in PBS and blocked with 10% normal donkey serum in 0.4% Triton in PBS for 1 hour. Subsequently, slices were incubated with the primary antibodies Rabbit-anti-PV (ab11427, Abcam, Cambridge, UK) and anti-GFP-488 conjugated (A21311, Invitrogen, Carlsbad, CA) to visualize YFP expression. The primary Rabbit-anti-PV antibody was visualized by a Donkey-anti-Rabbit-Cy3 secondary antibody (Jackson ImmunoResearch Laboratories, West Grove, PA). Cell nuclei were counterstained with Hoechst 33258 (H3569, ThermoFisher Scientific, Waltham,MA).

**Statistics.** All values are reported as mean and standard error of the mean (SEM). Statistical analysis was performed using Matlab or Prism 6 (Graphpad Software Inc., La Jolla, CA). Unless otherwise mentioned, pairwise comparisons were made using Student’s t-test; multiple comparisons were performed using ANOVA with the appropriate post-hoc tests and
correlations were analyzed using linear regression. P<0.05 was assumed to reject the null hypothesis.

Results

Three classes of principal neurons in the PER-LEC deep layers

Whole cell recordings of 81 deep layer principal neurons in horizontal mouse brain slices were performed. Deep layer principal neurons were identified by their physiological properties and post-recording visualization of their localization, i.e. pyramidal like cell body and basal dendrites in the deep layers (Hamam et al., 2002; Canto and Witter, 2012). Intrinsic membrane properties and action potential firing were examined by an 800 – 1000 ms current injection with an amplitude evoking hyper- or depolarizations from -100 to -30 mV, in steps of 5 - 10 mV. We characterized three principal neuron types in the deep layers throughout the rostro-caudal extent of the PER and LEC network (Figure 1, Table 1, Supplementary figure 1 a-c), analyzing the following characteristics: (1) the presence of a voltage sag upon

<table>
<thead>
<tr>
<th>Table 1. Intrinsic properties of principal neurons and PV interneurons</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Property</strong></td>
</tr>
<tr>
<td>RMP (mV)</td>
</tr>
<tr>
<td>Input resistance (MΩ)</td>
</tr>
<tr>
<td>Membrane τ (ms)</td>
</tr>
<tr>
<td>Sag (mV)</td>
</tr>
<tr>
<td>Time to first AP (ms)</td>
</tr>
<tr>
<td>AP threshold (mV)</td>
</tr>
<tr>
<td>AP amplitude (mV)</td>
</tr>
<tr>
<td>AHP amplitude (mV)</td>
</tr>
<tr>
<td>Spike half width (ms)</td>
</tr>
<tr>
<td>Frequency 1st AP (Hz)</td>
</tr>
<tr>
<td>Frequency last AP (Hz)</td>
</tr>
</tbody>
</table>

All values are mean ± SEM. All values are measured at the current step above threshold.

1 AHP amplitude is measured from threshold to maximal afterhyperpolarization
2 frequency is determined as the inverse of the first inter-spike interval or last inter-spike interval

Abbreviations: RS, regular spiking; LS, late spiking; BF, burst firing; PV, parvalbumin; RMP, resting membrane potential; AP, action potential; AHP, afterhyperpolarization.
hyperpolarization, (2) latency to the first spike, (3) burst firing (consistent frequency of spike 1 and 2), and (4) spike frequency adaptation (Faulkner and Brown, 1999; Beggs et al., 2000; Canto and Witter, 2012; Fuchs et al., 2016). Regular spiking (RS) neurons (58/81) showed a

Figure 1. Intrinsic properties of 3 types of principal neurons and PV positive interneurons in the PER-LEC deep layer network. Typical example of the spike pattern (left), and spike-frequency plot (right) of regular spiking (a), late spiking (b) and burst firing (c) principal neurons and PV positive interneurons (d, gray). a-d left. Top: spike raster plot of four current injections of increasing amplitudes. Bottom: membrane voltage in response to a hyperpolarizing and threshold depolarizing current injection. a-d right. Spike-frequency of every spike at four current injections of increasing amplitudes. e. Left top: schematic representation of the lateral view of the mouse brain, the dotted line indicates the slice location. Left bottom: Representation of the distribution of the three principal neuron types (black) and PV interneurons (gray) recorded in a schematic horizontal mouse brain slice containing the AiP as the stimulated neocortical area, the PER and LEC. Right: Evoked synaptic currents in response to AiP stimulation in three classes of principal neurons (black) i.e. RS, LS, BF neurons, and
regular, adapting firing pattern and a hyperpolarization-induced voltage sag (2.4 ± 0.2 mV, Figure 1 a, Table 1). Late spiking (LS) neurons (11/81) showed delayed firing (142.9 ± 13.3 ms) and lacked a voltage sag upon hyperpolarization (Figure 1 b, Table 1). Burst firing (BF) neurons (12/81) showed a voltage sag (2.5 ± 0.4 mV; Figure 1 c, Table 1) upon hyperpolarization and typically fired the first two action potentials at a high frequency (68.4 ± 19.0 Hz) after which action potential firing showed adaptation (Figure 1 c).

**Stimulus-evoked synaptic input rarely evokes action potential firing in PER and LEC deep layer principal neurons**

The recruitment of the PER and LEC deep layers in response to an AiP synaptic stimulation was investigated by applying an electrical stimulus in the AiP superficial layers at the intensity evoking the maximal synaptic response in 74 principal neurons (average stimulus intensity was 754 ± 30 µA) while the voltage was clamped at -70 mV (Figure 2 a, b). The stimulus was applied three times, with an inter-stimulus interval of 4 seconds; consecutively recorded responses were averaged. The majority of principal neurons (66/74, 89%) received synaptic input from the AiP superficial layers after electrical stimulation. The onset latency and peak amplitude of the synaptic response were compared between the three types of principal neurons. Although BF neurons were more prominently localized in the PER, RS, LS and BF neurons did not differ in latency, in case the latency was corrected for the distance from the stimulus electrode (latency: F(2,63) = 3.10, p = 0.052). The peak synaptic response was also comparable between the three principal neuron subtypes (F(2, 63) = 0.92, p = 0.4; Figure 1 e, Figure 2 b’, b”). The principal neurons are therefore combined into one experimental group and analyzed together (n = 66). Analysis of the evoked synaptic current revealed that the latency increased with distance along the rostro-caudal PER-LEC axis. If we assume a synaptic delay of 0.5 ms in all PER-LEC neurons, the propagation velocity of activity is described by the slope of the robust linear fit of the relationship between the latency of responses and distance to the stimulus electrode. AiP stimulation evoked activity which propagated with 0.199 m/s (Figure 2 b’). The peak of responses was not related to the distance from the electrode (regression analysis n.s.; Figure 2 b”) and had an average amplitude of 327 ± 38 pA.

To address the output of deep layer principal neurons in response to their synaptic input, the evoked postsynaptic potentials were recorded in current clamp (Figure 2 c). Only 3 out
of 51 principal neurons (1 BF and 2 RS neurons) fired a single action potential in response to synaptic input. The peak amplitude of the evoked postsynaptic potentials in non-firing principal neurons did not relate to distance from the stimulus electrode (regression analysis n.s.; Figure 2 c’). Since stimulus application was repeated three times, we were able to address the success rate for the AiP evoked action potentials. The AiP stimulus evoked 3/3 times an action potential in 2 principal neurons and 1/3 times an action potential in 1 principal neuron (Figure 5 h, black dots). This led to the probability of the AiP stimulus evoking an action potential in spiking neurons of 0.78 ± 0.07.

**Figure 2. Evoked synaptic input and postsynaptic potentials in PER-LEC deep layer principal neurons.**

- **a.** Schematic overview of all principal neurons recorded in a horizontal mouse brain slice of one hemisphere containing the AiP (neocortical area), the PER and LEC. The stimulus electrode was placed in the AiP superficial layers. PER recorded cells are indicated with an open triangle (△) and LEC recorded cells are indicated with a closed circle (●). Colors represent three subtypes of principal neurons, i.e. regular spiking (black), late spiking (cyan) and burst firing (orange) neurons.
- **b.** AiP stimulus evoked postsynaptic currents in principal neurons sorted based on their distance to the stimulus electrode (colors represent three subtypes of principal neurons, see a). Principal neurons were voltage clamped at -70 mV, the arrowhead (▼) indicates the moment the stimulus was applied. **b’**. The EPSC onset latency increased when recordings were performed at increasing distance from the AiP stimulus electrode (n = 66). Colors represent the three principal neuron subtypes. **b’’**. The stimulus electrode distance did not correlate with the EPSC peak amplitude.
- **c.** Current clamp recordings of stimulus evoked postsynaptic potentials in 51 principal neurons. EPSPs were consecutively recorded three times at resting membrane potential. Only three out of 51 principal neurons responded with a single action potential after the stimulus, the other 48 neurons only showed an EPSP. Colors represent the three principal neuron subtypes. **c’**. EPSP amplitude of the neurons which only showed an EPSP, so no firing, after stimulus application revealed no relationship between EPSP amplitude and the distance of the recorded neuron to the NC stimulus electrode. Colors represent the three principal neuron subtypes.

**Abbreviations:** AiP, agranular insular cortex; PER, perirhinal cortex; LEC, lateral entorhinal cortex; R, rostral; C, caudal; L, lateral; M, medial; EPSC, excitatory post-synaptic current; RS, regular spiking; LS, late spiking; BF, burst firing; EPSP, excitatory post-synaptic potential; PrN, principal neuron.
Synaptic input onto deep layer principal neurons is composed of a small excitatory and a larger inhibitory conductance

Since the deep layer principal neurons rarely emitted action potentials in response to a synaptic input, we hypothesized that the inhibition-excitation balance was in favor of inhibition and aimed to address the relation between the stimulus-evoked inhibition and excitation. The synaptic currents recorded at 4-5 holding potentials ranging from -90 to -50 mV in response to 50% and 100% of the maximum stimulus intensity (Figure 3 b-c, insets) were used to estimate the synaptic conductance changes (Figure 3 b-c) evoked by the afferent input in 53 principal neurons along the PER-LEC axis (Figure 3 a). The evoked synaptic conductance at two stimulus intensities was linearly decomposed into an excitatory and inhibitory conductance during 100 ms, under the assumption of a reversal potential of 0 mV for excitation and -70 mV for the fast, chloride mediated GABA A dependent inhibition.

Figure 3. Evoked excitatory and inhibitory conductance in response to AiP stimulation at two intensities. a. A schematic overview of the recorded PER (Δ) and LEC (●) principal neurons (n = 48). b. Typical example of the G exc (black trace) and the GABA A mediated G inh (red trace) calculated from the evoked post-synaptic currents shown in the inset, evoked at 50% of the stimulus intensity evoking the maximum response. The arrowhead (▼) indicates the moment the stimulus is applied. c. The G exc (black trace) and the GABA A mediated G inh (red trace) calculated from the evoked post-synaptic currents shown in the inset, of the same cell as in b but with 100% stimulus intensity. d-f. Quantification of the onset latency (d), peak amplitude (e) and half width (f) of the G exc (black) and G inh (red) recorded in PER and LEC principal neurons at 50% and 100% of the stimulus intensity evoking the maximum response. g. Plot of the difference between the G exc and G inh latency in relation to the distance from the stimulus electrode in the recorded PER and LEC principal neurons after stimulation at 100% intensity.

Abbreviations: G exc, excitatory conductance; G inh, inhibitory conductance; PER, perirhinal cortex; LEC, entorhinal cortex; R, rostral; C, caudal; L, lateral; M, medial.
The latency of the evoked conductances was defined (Figure 3 d). If an evoked conductance was detected (n = 49 principal neurons), the peak amplitude and half width were determined (Figure 3 e, f). Combined evaluation of the latency, peak amplitude, and half width of the inhibitory and excitatory conductances after stimulation at the intensity evoking the maximum response (100% intensity) (F(2.289, 110.3) = 22.68, p<0.0001). Sidak’s multiple comparison post hoc analysis showed that the evoked excitatory conductance had a shorter latency than the evoked inhibitory conductance (latency$_{exc} = 7.2 ± 0.4$ ms, latency$_{inh} = 10.5 ± 0.6$ ms, p<0.0001; Figure 3 b-d), the peak of the inhibitory conductance was larger (peak$_{exc} = 3.8 ± 0.5$ nS, peak$_{inh} = 8.4 ± 1.1$ nS, p<0.0001; Figure 3 b, c, e), and the inhibitory response lasted longer (half width$_{exc} = 14.0 ± 1.2$ ms, half width$_{inh} = 17.6 ± 2.0$ ms, p=0.013; Figure 3 b, c, f) compared to the excitatory response. To examine whether the composition of synaptic responses changed when the input is weaker, we also stimulated the AiP at the intensity evoking the half maximum response (50% intensity, Figure 3 b – f). We found that both excitation and inhibition had a slightly longer latency (latency$_{exc} = 8.2 ± 0.4$ ms p<0.0001, latency$_{inh} = 12.5 ± 0.9$ ms, p<0.001), lower peak amplitude (peak$_{exc} = 2.1 ± 0.3$ nS p<0.001, peak$_{inh} = 5.2 ± 0.9$ nS p<0.0001) and a comparable half width (half width$_{exc} = 16.5 ± 2.0$ ms, half width$_{inh} = 23.5 ± 3.4$ ms) compared to responses evoked at maximum stimulus intensity.

To address whether the delay between the recruitment of excitation and inhibition was different along the rostro-caudal axis of the PER and LEC, we tested whether the difference between the latency of the $G_{exc}$ and $G_{inh}$ changed with the distance from the stimulus electrode (Figure 3 g). We found that there was no relation between the latency difference of the $G_{exc}$ and $G_{inh}$ and the distance (average delay was 3.3 ± 0.4 ms; regression analysis n.s., Figure 3 g), indicating that the delay between excitation and inhibition does not depend on conduction velocity from the stimulus to the site of recording, which led to the hypothesis that $G_{inh}$ is likely recruited in the local PER and LEC network.

**AiP evoked fast inhibition is recruited in the local PER-LEC network**

The latency of the inhibitory conductance we recorded in principal neurons was relatively short, which could imply the presence of direct, monosynaptic inhibitory input from the stimulated neocortical AiP. This hypothesis is in line with the long-range inhibitory projections...
from neocortical areas towards the PER and EC described by Pinto et al. (2006). To address the monosynaptic inhibition hypothesis, we bath applied ACSF containing 20 µM CNQX and 10 µM APV to block the AMPA and NMDA receptor mediated excitatory input. Besides mono and polysynaptic excitation, this prevents polysynaptic recruitment of interneurons in the local circuitry, only allowing possible monosynaptic, long-range GABAergic projections from the AiP to evoke an inhibitory response in principal neurons. After obtaining the AiP evoked conductances in control ACSF (Figure 4 a), we obtained the conductances while excitatory transmission was blocked (Figure 4 b). This abolished both excitatory and inhibitory conductances (Figure 4 b, c, p=0.0018, n = 8, Friedman test), suggesting the absence of a direct inhibitory connection from the AiP onto deep layer principal neurons in this mouse brain slice preparation. This implies that the inhibitory conductance evoked in PER-LEC deep layers must originate from local inhibitory neurons.
**PV interneurons are strongly recruited by synaptic input**

AiP synaptic input to principal neurons in the PER-LEC network evoked a large inhibitory and a smaller excitatory synaptic conductance (Figure 3). PV expressing fast spiking interneurons are, amongst other interneuron types, present in the local PER and LEC network (Wouterlood et al., 1995; Barinka et al., 2012) and make synaptic contacts onto the axosomatic region of principal neurons (Markram et al., 2004; Klausberger and Somogyi, 2008; Kubota et al., 2016). PV interneurons are capable of high frequency firing upon depolarization (Figure 1 d), and can therefore exhibit strong inhibitory action onto principal neurons (Pfeffer et al., 2013). This led to the hypothesis that PV interneurons could well be strongly recruited by the AiP input, to account for the large, locally activated, inhibitory conductance (Figure 3, Figure 4). To address this hypothesis, the synaptic input (Figure 5 a - d) and action potential firing (Figure 5 e - h) in response to neocortical AiP stimulation was examined in horizontal slices of transgenic mice conditionally expressing YFP in PV interneurons (Supplementary figure 1d). PV interneurons were recorded along the rostro-caudal axis of the PER and LEC deep layer network (Figure 5 a, left) and were characterized by a small membrane time constant, a lack of a hyperpolarization induced voltage sag, a short onset latency to the first induced action potential, and a smaller AP amplitude than principal neurons (Figure 1 d-e, Table 1).

AiP stimulation evoked complex synaptic responses in 53/56 the PV interneurons (95%). Assuming a synaptic delay of 0.5 ms, the latency increased with distance from the stimulus electrode (slope = 0.192 mm/ms; Figure 5 a right, b) and the peak amplitude of the response decreased with distance along the rostro-caudal extent of the PER and LEC (slope = -0.0002 mm/pA, R² = 0.32, F(1,51) = 24.01, p<0.0001; Figure 5 a right, c). Long latencies were accompanied with a small peak amplitude (Figure 5 d), at larger peak amplitudes, the latency did not show large values.

We next examined whether the PV interneurons fired action potentials in response to AiP stimulation (Figure 5 e-h). We recorded 51 PV interneurons in current clamp to determine the postsynaptic potentials and evoked firing in response to AiP stimulation in three consecutive repeats. We found that in total 31/51 deep layer PV interneurons recorded over the whole extent of the PER and LEC (Figure 5 a, e, f), fired action potentials in response to AiP stimuli with a success rate for the AiP evoked synaptic input of 0.90 ± 0.04 in firing PV
interneurons. Assuming a constant synaptic delay of 0.5 ms, the latency of the first evoked spike increased with distance from the stimulus electrode (Figure 5 h, slope = 0.158 mm/ms), with a velocity which was in the range of axonal conduction velocity (Telfeian and Connors, 2003; Willems et al., 2016). This conduction velocity was slightly lower than the velocity.
calculated based on the response latency, likely because of the combination of mono- and polysynaptic origin of the responses, leading to more variable spike timing.

The standard deviation of the latency of the first evoked spike in all three consecutive repetitions in response to AiP stimulation was used as an indicator for the spike jitter and was $1.0 \pm 0.6$ ms, suggesting a very reproducible recruitment of PV interneurons directly after synaptic input is received. Although the peak of the evoked response decreased with distance along the PER-LEC rostro-caudal axis, the average frequency of the evoked firing in PV interneurons was not related to distance (regression analysis n.s.; Figure 5 f). A subset of PV interneurons (4/31) persistently continued firing after the stimulus (Figure 5 g, h) indicating that PV interneurons are strongly recruited in the PER and LEC network and can therefore exhibit strong inhibitory control onto deep layer principal neuron activity.

**Connectivity between local PV interneurons and principal neurons**

Since local PV interneurons strongly respond to AiP synaptic input (Figure 5), we hypothesized that these PV interneurons project onto principal neurons locally in the PER-LEC deep layer network. To address the connectivity of deep layer PV interneurons and principal neurons in the PER and LEC deep layers, we performed 29 paired recordings of principal neurons and PV interneurons. After successfully obtaining whole-cell configuration in both neurons within a $200 \, \mu m$ inter-neuron-distance, random frozen noise was injected in the principal neuron or PV interneuron to evoke reproducible, randomly distributed action potential firing at an average frequency of 1-2 Hz (Figure 6 a, b). Principal neuron-PV pairs were considered connected when the onset of the unitary postsynaptic currents (uEPSC or uIPSC) following the evoked spikes clustered within 4 ms after the spikes (Figure 6 a”, b” and the probability of spike transmission was at least 0.6 (Csicsvari et al., 1998; Miles, 1990).

First, principal neuron firing was induced and the PV interneuron was clamped at -70 mV to simultaneously record uEPSCs in response to a principal neuron action potential (Figure 6 a, a”). In 11/29 recordings (38%), the principal neuron projected onto the simultaneously recorded PV interneuron (Figure 6 a - a”). The latency of the uEPSC in the PV neuron was $2.5 \pm 0.2$ ms after the peak of the principal neuron action potential (Figure 6 c, n = 312 action potentials from 11 principal neurons). The uEPSC rise time in PV interneurons was $0.61 \pm 0.15$
ms (Figure 6 d). The probability of the principal neuron spike evoking a uEPSC was 0.83 ± 0.04 in the 11 connected principal neuron to PV interneuron pairs (Figure 6 e).
Second, the projection from a PV interneuron onto a principal neuron was tested by recording PV interneuron firing-induced uIPSCs in the principal neuron, clamped at -50 mV to reveal the outward inhibitory currents (Figure 6 b, b'). In 14% of the pairs (4/29), a PV to principal neuron projection was detected. The latency of the uIPSCs was 1.33 ± 0.05 ms after peak time of the PV spike (Figure 6 b’’- c, n = 99 action potentials from 4 PV interneurons). The uIPSC rise time in principal neurons was 2.68 ± 0.15 ms (Figure 6 d). The probability that a PV spike evoked a uIPSC was high, 0.98 ± 0.025 (Figure 6 e, n = 4 pairs). Although the latencies of both uEPSCs and uIPSCs showed a comparable distribution (Figure 6 c), the distribution of the rise time of uEPSCs was skewed compared to the distribution of the rise time of uIPSCs (Figure 6 d), indicating that uEPSCs had faster kinetics than uIPSCs.

Only a subset (45%) of the recorded pairs was connected, 52% of the recorded pairs were not connected in this study and 2/29 recorded pairs (7%) were reciprocally connected.

**Relation between synaptically evoked PV firing and inhibitory conductance in principal neurons**

We next examined if PV interneurons could induce the fast, large inhibitory conductance observed in principal neurons in the local deep layer PER and LEC network. The temporal dynamics of PV interneuron recruitment should be fast enough to explain the fast inhibitory conductance which we recorded in the principal neurons (Figure 3). To compare the recruitment of the PV and principal neurons in response to AiP stimulation, we simultaneously recorded evoked synaptic currents in principal neurons and PV interneurons (n = 18 pairs, Figure 7 a). The latency of the evoked synaptic responses in the PV interneurons (8.5 ± 0.9 ms) was shorter than in principal neurons (10.7 ± 1.3 ms; t(17) = 2.5, p = 0.02; Figure 7 b, c). The peak amplitude of the evoked synaptic response in PV interneurons (714 ± 165 pA) was larger than the evoked synaptic response in principal neurons (343 ± 104 pA; t(17) = 3.1, p = 0.007; Figure 7 b, d). These results suggest, together with the absence of firing in principal neurons (Figure 2, Figure 6), that AiP stimulation recruits the inhibitory network predominantly in a feedforward manner. Furthermore, the conduction velocity of the synaptic input in both principal neurons and PV interneurons was comparable, suggesting that both types receive input from the same axon fibers.
Finally, we examined the relation between PV firing and the evoked inhibitory conductance in principal neurons in simultaneously recorded PV-principle neuron pairs. We compared the peak latency of the first PV spike to the $G_{inh}$ onset latency in the simultaneously recorded principal neuron in response to an AiP stimulus (Figure 7 e). We found that the latency of the $G_{inh}$ and the latency of the first evoked PV spike were not different (Figure 7 f;
Wilcoxon signed rank test n.s., n = 9 pairs). Additionally, the number of emitted spikes in the PV neuron strongly correlated with the peak G$_{inh}$ in the simultaneously recorded principal neuron (Figure 7 g; $R^2 = 0.84$, p<0.0001, n = 14 pairs), indicating that the number of spikes in the PV interneuron predicts the amount of inhibition in the simultaneously recorded principal neuron.

To get an indication of the necessary inhibitory input onto principal neurons in response to AiP stimulation, we estimated which inhibitory spike pattern was needed to reconstruct the recorded G$_{inh}$ in principal neurons. The mean time locked uIPSC in the principal neuron that was linked to a single action potential from the PV interneuron (Figure 6, Figure 8 a), allowed us to determine the conductance change of such single response, using the uIPSC and the driving force to calculate the unitary G$_{inh}$ (uG$_{inh}$). Subsequently, all recorded uG$_{inh}$ traces (n = 101 uG$_{inh}$ from 4 connected PV-principal neuron pairs recorded in 4 mice) were averaged to obtain one standardized uG$_{inh}$ (Figure 8 a). Using this standard PV interneuron firing-induced uG$_{inh}$, we could reconstruct the hypothetical pattern of inhibitory input received by a principal cell based on the firing patterns we recorded from the PV neurons (Figure 5). As shown above, 31 out of 51 PV interneurons emitted action potentials in response to AiP stimulation (Figure 5 e) and stimulation was repeated three times. Most, but not all, PV interneurons responded three out of three consecutive recorded repeats (Figure 5 h), which resulted in 83 AiP-evoked PV spike patterns. Figure 8 b (top traces) shows examples of six typical PV spike patterns. We assumed that the 83 recorded spike patterns describe a representative set of PV interneurons, which characteristically responded to the synaptic input.

Next, the predicted G$_{inh}$ evoked by these 83 spike patterns were calculated using the standard uG$_{inh}$ (examples are shown in Figure 8 b, bottom traces). The 83 predicted G$_{inh}$ traces were used to perform a nonnegative linear regression fit (Matlab) to find the weight of every PV interneuron spike pattern to reconstruct the recorded G$_{inh}$ in a principal neuron. A constraint was that the weight of the predicted G$_{inh}$ of every PV interneuron included, had to be at least 1. Figure 8 c shows a typical example of a recorded (black trace) G$_{inh}$ in a principal neuron and the accompanying predicted G$_{inh}$ (red trace). In this typical example, 41 PV interneuron spike patterns, all with weights between 1 and 5, were required to reconstruct the recorded G$_{inh}$ in this principal neuron. The number of PV interneuron spike patterns and
their weights were used to reconstruct the inhibitory spike pattern the principal neurons would have received, by multiplying the 41 included PV spike patterns by their weights and in that way creating the predicted spike raster plot of the total inhibitory spike pattern (Figure 8 d). We were able to fit the recorded $G_{\text{inh}}$ of 49 principal neurons. Figure 8 e shows which
PV spike patterns were included for each principal neuron $G_{\text{inh}}$ reconstruction. With the population of 83 PV spike patterns we had enough variation in inhibitory spike patterns to reconstruct the recorded $G_{\text{inh}}$ in all 49 principal neurons.

To address when the majority of inhibitory spikes have to occur to evoke the recorded $G_{\text{inh}}$, we reconstructed peristimulus time histogram (PSTH). The PTSH was normalized to the onset latency of the recorded $G_{\text{inh}}$, binned, averaged over the PSTH of the 44 principal neurons of which the inhibitory spike pattern was reconstructed and normalized to the bin with the maximum spike count (Figure 8 f). The inhibitory activity was most prominent in the first 20 ms of the inhibitory input, but continues during the time course of 100 ms with fewer spikes, probably to create long lasting suppression of activity within the PER-LEC network. We predicted that principal neurons received 4 – 733 inhibitory spikes in response to AiP stimulation, based on the 49 reconstructed spike patterns. In conclusion, with our set of recorded PV interneurons, with differential stimulus evoked spike patterns, we could explain the $G_{\text{inh}}$ recorded in all 49 principal neurons.

**Discussion**

Information from the neocortex travelling towards the hippocampus for memory consolidation is relayed by the PER-LEC network (Buzsáki, 1996; Pennartz et al., 2002), where transmission is most likely regulated by inhibition (de Curtis and Paré, 2004). This study addressed the recruitment of the inhibitory and excitatory neuronal local circuitry in the deep layers of the PER and LEC network of the mouse. Stimulation of the superficial layers of the AiP, a neocortical afferent of the PER and LEC, revealed that the PV interneurons are involved in eliciting strong inhibition of principal neurons in the deep layer network.

**Three PER-LEC principal neuron types and their evoked synaptic input**

We recorded principal neurons in the deep layers of the PER-LEC network because the LEC deep layers are considered to play a significant role in gating activity transmission, likely regulated by the inhibitory circuitry (Koganezawa et al., 2008; Willems et al., 2016). Examination of the intrinsic properties like the hyperpolarization-induced sag and firing properties of these deep layer principal neurons revealed three subtypes, i.e. RS, LS and BF
neurons in mouse brain slices, which are comparable with the assumed excitatory, glutamate containing neurons in rats (Somogyi et al., 1998; Faulkner and Brown, 1999; Moyer et al., 2002) and guinea pigs (Martina et al., 2001).

Superficial AiP stimulation evoked synaptic responses in both PER and LEC deep layer principal neurons. The three subclasses responded similarly to AiP stimulation, as the latency and peak amplitude did not differ and all types refrained from action potential firing. This phenomenon might depend on the stimulated afferent, as seen in the piriform cortex, where BF and RS neurons respond similarly to layer Ib, but differentially to lateral olfactory tract input (Suzuki and Bekkers, 2006). Our data suggest that synaptic input from the AiP to PER-LEC deep layer neurons is not principal neuron subtype specific. It is however still possible that these neuronal subtypes react differently to the same synaptic input when the membrane potential is around firing threshold, as a result of their different intrinsic properties. Since the evoked EPSP almost never induced action potential firing in these experiments, the three subtypes of principal neurons were pooled.

The latency of the synaptic input in principal neurons gradually increased with the distance from the stimulation electrode, comparable with findings of Biella et al. (2001) and Unal et al. (2012). The synaptic input is shown to be both mono and polysynaptic in deep layer PER and LEC neurons (Deacon et al., 1983; Burwell and Amaral, 1998a; de Villers-Sidani et al., 2004; Unal et al., 2013). A combination of conduction velocity, distance and polysynaptically transmitted activity can explain the increasing latency of synaptic responses in neurons situated more caudal in the PER-LEC network. The polysynaptic response in the LEC could originate from PER neurons projecting to the apical dendrites of LEC deep layer neurons (Burwell and Amaral, 1998b; Biella et al., 2002; de Villers-Sidani et al., 2004). Both the peak EPSC and EPSP in principal neurons were not related to distance along the rostro-caudal axis, suggesting no difference between PER and LEC excitation. This finding is in line with Mathiasen et al. (2015) who did a tracing study showing that the neocortical AiP is a presynaptic target of both deep layer PER and deep layer LEC neurons in the rat.

**Output of principal neurons and PV interneurons**

Although 89% of the recorded principal neurons received synaptic input, only 6% (3 out of 51) of them spiked, while 61% (31 out of 51) of the PV neurons fired after AiP stimulation
in this horizontal mouse brain preparation. Pelletier et al. (2004) found percentages of firing neurons in the deep layers of the PER (40%) and the EC (1.4%) in vivo. These results together suggest that superficial layer AiP can evoke synaptic activity in deep layer PER and LEC principal neurons, but this activity is not transmitted from the LEC to the postsynaptic targets (Biella et al., 2002). However, since the brain consists of approximately 80% excitatory neurons and 20% inhibitory neurons (Markram et al., 2004), this small percentage of firing principal neurons might be effective. If we take an example population of 1000 neurons, there will be 800 principal neurons and 200 interneurons. We found 6% firing principal neurons, resulting in 48 (800*0.06) firing principal neurons, and 61% firing PV interneurons, leading to 122 PV interneurons firing action potentials in response to an input in our example population. This finding supports the sparse coding strategy, which assumes that only a small portion of the cortical principal neurons fire in a certain event, responsible for information transfer in the EC-hippocampal circuitry (Mizuseki and Buzsáki, 2013) and this balance might be critical to maintain a self-organized and controlled activity in large scale networks. Furthermore, computer models showed that a small portion of firing excitatory neurons with strong synaptic weights can be sufficient to have ongoing network activity (Ikegaya et al., 2013).

**AiP recruits feedforward inhibition**

The low firing probability of principal neurons in response to synaptic input could be due to a marginal excitatory input or a massive inhibition. We found that AiP stimulation evoked a larger inhibitory than excitatory conductance in the PER-LEC deep layer principal neurons, suggesting more GABA\textsubscript{A} than glutamate receptor activation at principal neuron postsynapses. This phenomenon is not necessarily surprising since a larger inhibitory synaptic conductance than excitatory conductance can lead to balanced inhibitory and excitatory synaptic currents, due to a smaller driving force for inhibition than for excitation (Puzerey and Galán, 2014). However, the short latency difference between the excitatory and inhibitory conductance and especially the larger, longer lasting inhibitory conductance could prevent firing of deep layer principal neurons once they are depolarized.

It has been previously shown that stimulation of the temporal neocortex in brain slices of guinea pigs evoked a pure excitatory response when the recording electrode was more than
1 mm away from the stimulus electrode, while more closely situated neurons showed a sequence of excitatory and inhibitory potentials (Martina et al., 2001). In contrast, in this study we find both excitatory and inhibitory evoked components recruited in the same fashion along the rostro-caudal extent of the PER and LEC network. This indicates that the AiP projections in mouse horizontal brain slices evoke both excitation and inhibition in the whole PER-LEC network.

A broad range of interneurons are defined, based on several characteristics like morphology, physiological and connectional properties. Good candidates for delivering strong inhibition on principal neurons are the PV expressing interneurons. These interneurons target the axo-somatic region of neurons and therefore evoke large inhibitory currents in the post-synaptic neuron upon firing (Pfeffer et al., 2013; Jiang et al., 2015). The PV neurons received a larger evoked synaptic current than principal neurons, which can be due to the presence of larger or more glutamatergic terminals on interneuron dendrites, resulting in more effective activity transmission in interneurons (for review see Buzsáki et al., 2007). This fits neatly with the discovered PV firing patterns, often showing multiple action potentials, at a high frequency, in response to AiP stimulation. Although the role of different interneuron subtypes in evoking the large $G_{\text{inh}}$ in principal neurons is not yet clear, we showed a complete reconstruction of the recorded $G_{\text{inh}}$ in principal neurons solely based on the firing patterns of PV interneurons. This indicates that variation of responses in PV interneurons possibly is enough to explain the inhibitory input in principal neurons. Future studies may reveal the role of other interneuron types in this large $G_{\text{inh}}$ in the PER-LEC deep layer principal neurons.

Moreover, the synaptic input from the AiP is first received by PV and then by principal neurons in the PER and LEC deep layers when simultaneously recorded and the principal did not fire in response to AiP stimulation. Based on the comparable conduction velocities derived from the evoked EPSCs in principal neurons and PV interneurons we surmise that the same fibers innervate both neuron types. We therefore propose that deep layer PV interneurons are recruited in a feedforward manner by the AiP synaptic input and substantially contribute to strong principal neuron inhibition in the PER and LEC network. This is supported by anatomical data showing a high incidence of excitatory projections from the PER to GABAergic neurons in the EC (Pinto et al., 2006). However, since bidirectional
connections between PV and principal neurons are found, it is still likely that PV neurons can also provide feedback inhibition if local principal neuron would fire.

**Functional relevance of feedforward inhibition in the PER-LEC network**

The recruitment of feedforward inhibition preventing deep layer principal neurons from firing can have two plausible functions: First, inhibition of activity in the deep layers in response to superficial neocortical input is in line with the general hypothesis that information travelling towards the hippocampus is mainly transmitted via the superficial layers of the PER-EC network (Ruth et al., 1988; Witter, 1993) and that the deep layers return the information from the hippocampus to the neocortical areas (Buzsáki, 1996; Canto et al., 2008). We therefore hypothesize that this strong inhibitory response in the deep layers blocks the output pathway of the hippocampus, while possibly facilitating the input pathway via the superficial layers. This hypothesis is supported by data showing that PER superficial layer stimulation results in a significant monosynaptic activation of LEC superficial dendrites (de Villers-Sidani et al., 2004). Besides, a current source is found in layer V of the LEC *in vivo*, which could either be the result of apical dendrite activation of layer V principal neurons or deep layer inhibition (de Villers-Sidani et al., 2004). A similar concept of blocking activity propagation was found for regulation of the output from the hippocampus towards the EC (Gnatkovsky and de Curtis, 2006). They showed that the hippocampal output transfers through the EC deep layers, while the superficial MEC neurons were simultaneously inhibited, leaving only MEC deep layer neurons excitable by synaptic input arriving from the hippocampus.

Second, PV interneuron activity can be controlled by inputs from regulatory regions. By inhibiting the interneuron mediated inhibition, a window of opportunity could be created to transmit activity through the PER and EC. For example cholinergic inputs from the basal forebrain regulate inhibitory activity in the EC as well as in the auditory cortex (Apergis-Schoute et al., 2007; Kuchibhotla et al., 2016) and hippocampal long range inhibitory projections to the EC specifically target interneurons (Melzer et al., 2012). Such mechanism would be useful to release the EC from a strong intrinsic inhibitory control, to regulate information transmission to the hippocampus.
Altogether, our study shows a strong recruitment of PV interneuron mediated inhibition in the deep layers of the PER and LEC network by the neocortical AiP. This inhibition is likely to play a key role in regulating selective transmission of information travelling to and coming from the hippocampus.

Conflict of Interest

The authors declare no competing financial interests.

Acknowledgements

We gratefully thank Lieneke Kooijman for the help with the immunohistochemistry, Hendrikus van Heesbeen and Christiaan Levelt for providing the eYFP-floxed and PV-Cre animals, respectively, and the Van Leeuwenhoek Centre for Advanced Microscopy for help with the confocal imaging.
Supplementary material
Supplementary figure 3.1. Morphology of 3 types of principal neurons and PV positive interneurons in the PER-LEC deep layer network. a-c. Confocal images of biocytin filled regular spiking (a), late spiking (b) and burst firing (c, arrowhead) principal neurons and PV positive interneurons (c, red asterisk). Scale bar 100 µm. d. Immunohistochemical staining of PV (red, top), eYFP (green, middle) and the merge (bottom) showing complete overlap of PV and eYFP, scale bar 50 µm.

Abbreviations: PV, parvalbumin; eYFP, enhanced yellow fluorescent protein.