Inhibitory connectivity dominates the fan cell network in layer II of lateral entorhinal cortex

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Abbreviated title: Local connectivity of fan cells in LII of LEC

Number of pages: 37

Word count abstract: 222

Word count introduction: 657

Word count discussion: 842

Number of display items: 10 figures + 1 table

Acknowledgements

This study has been supported by grant number 227769 of the Research Council of Norway, The Kavli Foundation, the Centre of Excellence scheme of the Research Council of Norway – Centre for Neural Computation, grant number 223262, and the National Infrastructure scheme of the Research Council of Norway – NORBRAIN, grant number 197467. We thank Pål Kvello for contributing to the initial part of the project, Paulo Girao for Matlab programming, Maximiliano J. Nigro for stimulating discussions, Edvard I. Moser for helpful comments, Jørgen Sugar and Øyvind W. Simonsen for commenting and proofreading the semi-final draft of the manuscript. The authors declare no competing financial interests.
ABSTRACT

Fan cells in layer II of the lateral entorhinal cortex (LEC) form a main component of the projection to the dentate gyrus, CA3 and CA2 of the hippocampal formation. This projection has a counterpart originating from stellate cells in layer II of the medial entorhinal cortex (MEC). Available evidence suggests that the two pathways carry different information, exemplified by a difference in spatial tuning of cells in LEC and MEC. The grid cell, a prominent position-modulated cell type present in MEC, has been postulated to derive its characteristic hexagonal firing pattern from dominant disynaptic inhibitory connections between hippocampal-projecting stellate cells. Given that grid cells have not been described in LEC, we aim to describe the local synaptic connectivity of fan cells, to explore whether the network architecture is similar to that of the MEC stellate cell. Using a combination of in vitro multi-cell electrophysiological and optogenetic approaches in acute slices from rodents of either sex, we show that excitatory connectivity between fan cells is very sparse. Fan cells connect preferentially with two distinct types of inhibitory interneurons, suggesting disynaptic inhibitory coupling as the main form of communication among fan cells. These principles are similar to those reported for stellate cells in MEC, indicating an overall comparable local circuit architecture of the main hippocampal-projecting cell types in the lateral and medial entorhinal cortex.

SIGNIFICANCE STATEMENT

Our data provide the first description of the synaptic microcircuit of hippocampal-projecting layer II cells in the lateral entorhinal cortex (LEC). We show that these cells make infrequent monosynaptic connections with each other, and that they preferentially communicate through a disynaptic inhibitory network. This is similar to the microcircuit of hippocampal-projecting stellate cells in layer II of the medial entorhinal cortex (MEC), but dissimilar to the connectivity observed in layer 2 of neocortex. In MEC, the observed network structure has been proposed to underlie the firing pattern
of grid cells. This opens the possibility that layer II cells in LEC exhibit regular firing patterns in an unexplored domain.

INTRODUCTION

The entorhinal cortex (EC) has a pivotal position in the hippocampal-parahippocampal episodic memory system, serving as the main gateway for information entering the hippocampal formation (HF). EC connects to HF through parallel pathways mediated by the lateral (LEC) and medial (MEC) entorhinal subdivisions (Knierim et al., 2014; Cappaert et al., 2015; Knierim, 2015). These input streams point to functionally distinct and potentially complementary roles for LEC and MEC in hippocampal function (Keene et al., 2016), a notion supported by numerous in vivo studies (Fyhn et al., 2004; Hafting et al., 2005; Hargreaves et al., 2005; Van Cauter et al., 2008; Lu et al., 2013; Tsao et al., 2013; Van Cauter et al., 2013). LEC is involved in encoding contextual information (Wilson et al., 2013; Pilkiw et al., 2017), demonstrated by the ability of cell ensembles to represent time across different contexts (Tsao et al., 2018), or by single cells to represent the past or present position of objects (Deshmukh and Knierim, 2011; Tsao et al., 2013). In contrast, MEC contains elements of the brain’s navigational circuit, including grid (Hafting et al., 2005), object-vector (Høydal et al., 2018), head-direction (Sargolini et al., 2006), border (Solstad et al., 2008) and speed cells (Kropff et al., 2015), collectively supplying HF with context-invariant information signaling the animal’s trajectory.

Consistent with distinct functional specializations is the discovery of different embryological origins of LEC and MEC. LEC originates from caudal parts of the dorsolateral pallium, whereas MEC derives from the medial pallium (Medina et al., 2017). Hence, these areas might conceivably have developed cells and local circuitries that perform different computations to shape information represented and relayed to HF. The layer II (LII) projection to HF arises mainly from two reelin positive (RE+) cell types, the MEC stellate cell and the LEC fan cell (Germroth et al., 1989; Varga et al., 2010; Kitamura et al., 2014; Leitner et al., 2016). These cells have different dendritic morphologies and biophysical
membrane properties (Canto and Witter, 2012a, b), indicating distinctive integrative capacities of the two cell types.

The connectivity between MEC stellate cells has been emphasized as a plausible mechanism supporting the grid pattern, characterized by multiple firing fields arranged in a hexagonal lattice spanning the entire environment explored by the animal (Hafting et al., 2005). One influential class of models predicts this salient activity pattern to arise through attractor dynamics (Fuhs and Touretzky, 2006; McNaughton et al., 2006; Guanella et al., 2007; Burak and Fiete, 2009), achieved by the prevailing disynaptic inhibitory connectivity of stellate cells (Bonnevie et al., 2013; Couey et al., 2013; Pastoll et al., 2013). A detailed description of the local synaptic circuitry of LEC fan cells is lacking.

If grid activity in MEC LII arises due to inhibitory functional connections between stellate cells, the lack of grid cells in LEC (Hargreaves et al., 2005; Yoganarasimha et al., 2011) implies the presence of a principal cell local network architecture governed by different principles. Hence, fan cells, the likely LEC analogue of stellate cells, may depend more on excitatory connectivity. This interpretation is strengthened by their different developmental origins, where LEC appears more different from MEC than the neocortex (Medina et al., 2017), an area with higher probability of local principal cell connectivity in layer 2 than MEC (Lefort et al., 2009; Jouhanneau et al., 2015; Seeman et al., 2018). To test our hypothesis, we did in vitro multi-cell patch clamp recordings to probe connectivity between fan cells. Disproving our hypothesis, we report that monosynaptic connectivity between fan cells is sparse, and that they preferentially interact with two out of three distinct types of inhibitory interneurons. Using optogenetic methods, we substantiate these findings by demonstrating that fan cells receive predominant disynaptic inhibitory inputs, and limited excitatory inputs, arising from the activation of RE+ LII principal cells. Our data indicate that the microcircuitry of fan cells in LII of LEC shows a network motif similar to that described previously for stellate cells in MEC.

**MATERIALS AND METHODS**
Animals

All animals were group housed with up to five animals in one cage, kept at a 12:12 hour reversed day/night cycle, and had *ad libitum* access to food and water. Long Evans rats (postnatal age 23 - 25 days, n=2; postnatal age 28 - 80 days, n=47) and C57 mice (postnatal age 42 - 47 days, n=4) were used for whole-cell current clamp recordings. Mice of the transgenic MEC13-53A enhancer strain (postnatal age 84 – 112 days) were used for whole-cell voltage clamp recordings (n=15 mice) and histological assessment of LEC transgene expression (n=8 mice). All experiments were approved by the local ethics committee and were in accordance with the European Communities Council Directive and the Norwegian Experiments on Animals Act.

Preparation of MEC13-53A mice

Cloning of transgenic constructs and pronuclear injection

The enhancer sequence (coordinate mouse genome: chr8:49,906,388-49,908,569 in mm9) was cloned from BACs (chori.org) and transferred to pENTR™/D-TOPO® vectors by TOPO® cloning (Invitrogen, K2400-20). The enhancer was transferred to an injection plasmid by gateway cloning® (Invitrogen, 11791-019). The resulting injection plasmid consisted of a putative enhancer followed by a mutated heatshock promoter 68 (HSP68), a tTA gene, a synthetic intron and a WPRE element. The injection plasmid was linearized by enzyme digestion with XmnI and EcoRV to keep the relevant elements but remove the bacterial elements of the plasmids. Linearized vector of 7.5 Kbp was run on a 1% agarose gel and isolated using a Zymoclean Gel DNA Recovery Kit (Zymo research, D4001). Fertilized egg cells were injected with 1 µL of DNA at concentrations of 1 ng/µL, leading to pups of which 18 were genotypically positive and 6 expressed the transgene in the EC (Blankvoort et al., 2018). Pronuclear injections were done in the transgenic mouse facility of the University of Oregon.
Genotyping was done on ear tissue using a Kapa mouse genotyping kit (Kapa Biosystems, Cat# KK7302). Primer pairs for tTA (5’-GGACAAGTCCAAGGTGATCAAC-3’ and 5’-CTGGTGCTGAACAGCTCG-3’, 591bp product) and internal controls (5’-CTAGGCCACAGAATTGAAAGATCT-3’ and 5’-TAGGTGAAATTCTAGCATCATCC-3’, 324bp product) were added to the PCR mixture at a final concentration of 10 µM. The PCR reaction was done by an initial step of 4 minutes at 95 °C, then 20 cycles of 1 minute at 95 °C, 30 seconds at 70 °C reduced by 0.5 °C each cycle, and 30 seconds at 72 °C. This was followed by 20 cycles of 30 seconds at 95 °C, 30 seconds at 60 °C, and 30 seconds at 72 °C. Then a final 7 minute step at 72 °C. The products were run on a 1% agarose gel along with positive and negative controls.

AAV-TRE-tight-oChIEF-Citrine purification

Creation of pAAV-TRE-tight-WPRE was achieved by replacing the chimeric CMV-promoter/beta-globin intron sequence of pAAV-CMV-β-globin-intron-MCS-WPRE (Agilent) by the TRE-tight promoter from pTRE-tight vector (Clontech). Transgene oChIEF-Citrine was PCR amplified from plasmid #50974 (Addgene) and cloned into the plasmid pAAV-TRE-tight-MCS-WPRE which resulted in pAAV-TRE-tight-oChIEF-Citrine-WPRE. The positive clones were confirmed by restriction digestion analyses and subsequently by DNA sequencing. Endotoxin free plasmid maxipreps (Qiagen) were made for AAV preparations. The day before transfection, 7x 10^6 AAV 293 cells were seeded in DMEM containing 10% fetal bovine serum (ThermoFisher) and pencillin/streptomycin antibiotics into a 150 mm cell culture plate. Calcium chloride mediated cotransfection was done with 22.5 µg pAAV-containing the transgene, 22.5 µg pHelper, 11.3 µg pRC, 11.3 µg of either pXR1 or pAAV9 capsid plasmid. After 7 hours, the medium was replaced by fresh 10% FBS containing DMEM. The cells were scraped out after 72 hours, then centrifuged at 200xg and the cell pellet was subjected to lysis using 150 mM NaCl-20 mM Tris pH 8:0 buffer containing 10% sodium deoxycholate. The lysate was then treated with...
Benzonase nuclease HC (Millipore) for 45 minutes at 37 °C. Benzonase treated lysate was centrifuged at 3000xg for 15 mins and the clear supernatant was then subjected to HiTrap® Heparin High Performance (GE) affinity column chromatography using a peristaltic pump (McClure C JOVE 2011). The elute from the Heparin column was then concentrated using Amicon Ultra centrifugal filters (Millipore). The titer of the viral stock was determined as approximately $10^7$ infectious particles/mL.

**Acute slice preparation**

Rats or mice of either sex were deeply anesthetized with isoflurane and decapitated. The brain was quickly removed and immersed in cold (0 °C) oxygenated (95% O$_2$/5% CO$_2$) artificial cerebrospinal fluid (ACSF) containing 110 mM choline chloride, 2.5 mM KCl, 25 mM D-Glucose, 25 mM NaHCO$_3$, 11.5 mM sodium ascorbate, 3 mM sodium pyruvate, 1.25 mM NaH$_2$PO$_4$, 100 mM D-Mannitol, 7 mM MgCl$_2$ and 0.5 mM CaCl$_2$, pH 7.4, 430 mOsm. The brain hemispheres were subsequently separated and 400 µm thick semicoronal slices containing the lateral entorhinal cortex were cut with a vibrating slicer (Leica VT1000S, Leica Biosystems, Nussloch, Germany). The slices were cut with an angle of 20° with respect to the coronal plane to optimally preserve neurons and local connections (Tahvildari and Alonso, 2005; Canto and Witter, 2012a). Slices were moved to a holding chamber and incubated at 35 °C in oxygenated ASCF containing 126 mM NaCl, 3 mM KCl, 1.2 mM Na$_2$HPO$_4$, 10 mM D-glucose, 26 mM NaHCO$_3$, 3 mM MgCl$_2$ and 0.5 mM CaCl$_2$ for 30 minutes and then kept at room temperature for at least 30 minutes before use.

**Multi-cell current clamp recordings**

Cells were visualized using infrared differential interference contrast (IR-DIC) optics on two different patch clamp setups, either with a 40x/0.8 NA water immersion (WI) objective (Olympus BX51WI, Tokyo, Japan) or 20x/1.0 NA WI objective (Zeiss Axio Examiner.D1, Carl Zeiss, Jena, Germany). Groups of LII cells located deeper than 50 µm in the slice were selected for recording. Patch pipettes (3-8 MQ)
were pulled from standard wall borosilicate glass capillaries and filled with intracellular solution
containing 120 mM K-gluronate, 10 mM KCl, 10 mM Na₂-phosphocreatine, 10 mM HEPES, 4 mM Mg-
ATP, 0.3 mM Na-GTP, with pH adjusted to 7.3 and osmolality to 300-305 mOsm. Biocytin-HCl (0.5%,
Sigma-Aldrich, St Louis, MO, USA) was added to the pipette solution for recovery of neuronal
morphology. In a subset of experiments, different spectral variants of Alexa Fluor (AF) dyes (405, 488,
568, 633; Molecular Probes, Invitrogen, Thermo Fisher Scientific, Waltham, MA, USA) was included in
the individual pipettes instead of biocytin. Chloride reversal potentials were -47.0 mV for the internal
solution with biocytin-HCl, and -69.10 mV for the internal solution with Alexa Fluor dyes, calculated
using the Nernst equation. All recordings were carried out at 35 °C and the slices were continuously
superfused with oxygenated ACSF containing 126 mM NaCl, 3 mM KCl, 1.2mM Na₂HPO₄, 10 mM D-
Glucose, 26 mM NaHCO₃ 1.5 mM MgCl₂ and 1.6 mM CaCl₂.

Data acquisition and analysis

Whole-cell current clamp recordings with one rig were acquired with two Multiclamp 700 A/B
amplifiers (Axon Instruments, Molecular Devices, Foster City, CA) with a sampling rate of 10 kHz and
digitized with an ITC-1600 A/D board (HEKA Elektronik, Lambrecht, Germany). Data acquisition was
controlled by custom-written protocols in Igor Pro (WaveMetrics, Inc., Lake Oswego, OR, USA). Whole-
cell current clamp recordings on the second rig were collected with an EPC 10 Quadro USB amplifier,
controlled by the acquisition software Patchmaster (Heka Elektronik). Acquired data were low pass
Bessel filtered at 4 kHz and digitized at 10 kHz. For all recordings, pipette capacitance was
compensated and online bridge adjustments were performed (series resistance: median, 16 MΩ; 25th-
75th percentiles, 10-25 MΩ). Membrane potentials were not corrected for the liquid junction potential
(15.8 mV for the potassium based internal solution, calculated using the junction potential calculator
(JPCalc) plugin in pCLAMP (Molecular Devices). Synaptic connectivity was tested by inducing spike
trains at various frequencies (10, 20, 40 and 70 Hz) sequentially in each individual cell and
simultaneously monitoring the membrane potential of the other cells (Couey et al., 2013). Cells were recorded and connectivity tested at their native resting potentials (median, -70.25 mV; 25th-75th percentiles, -73.38 - -67.0 mV), but in rare cases stabilized with negative current to prevent spontaneous firing of action potentials. Possible disynaptic interactions were tested by activating two or three cells simultaneously and recording the membrane potential of the last unstimulated cell. The connectivity protocol was repeated up to 20 times. Possible connections were scrutinized during recording and later during analysis. Connections were accepted if their amplitudes evoked by 70 Hz stimulation were larger than 2.5 times the median of absolute deviations from the median (2.5*MAD). MAD is defined as $MAD = \text{median}(|X_i - \text{median}(X)|)$. In order to be considered a synaptic connection evoked specifically by stimulation of one of the other cells, the membrane potential deflection needed to occur within the time window where the presynaptic cell was active. All cells with resting membrane potential lower than -55 mV were screened for synaptic connections. Possible postsynaptic responses were analyzed by computing average membrane potential traces from 5-20 individual sweeps for each presynaptic stimulation protocol (10, 20, 40, 70 Hz). Due to a low signal-to-noise ratio of the small amplitude synaptic potentials in single sweeps, the averages were used to calculate synaptic latency, amplitude, latency to peak and half-width. Postsynaptic potential (PSP) amplitude was extracted from the deflection in response to the first action potential in the 10 Hz presynaptic train (single stimulus), as well as the peak of the postsynaptic membrane deflection following a presynaptic 70 Hz action potential train. The amplitude was defined as the potential difference between PSP peak and baseline potential (baseline measured as the average potential of a 50 ms interval immediately prior to the onset of presynaptic stimulation). PSP latency was calculated from the first PSP in response to 10 Hz presynaptic stimulation, and defined as the time interval between the peak of the averaged first presynaptic action potential to the onset of PSP. The onset of PSP was defined as the point where the PSP had reached 15% amplitude. Conducting the PSP analyses on the average trace precluded an investigation of the variability in onset times between trials, a
feature important to assess monosynapticity (Feldmeyer et al., 1999; Koelbl et al., 2015). Therefore, we defined PSPs with latencies < 3 ms evoked in pairs of excitatory cells as presumed monosynaptic (Peng et al., 2017). All connections in pairs of excitatory cells with longer latencies were classified as putative polysynaptic connections. In synaptically coupled cell pairs in which the shape of the presynaptic action potential was distorted due to high (>40 MΩ) access resistance, the connection was defined as unclassifiable. PSP half width was defined as the time difference between rising phase and falling phase of the PSP measured at 50% PSP amplitude. PSP latency to peak represented the time difference between peak of the presynaptic action potential and the peak of the PSP. Intrinsic membrane properties were measured from membrane voltage responses to step injections of hyperpolarizing and depolarizing current (500 ms duration, -400 pA to +500 pA, 100 pA increments, 3 seconds interpulse interval). Relevant intrinsic membrane properties were calculated for cells with series resistance less than 25 MΩ, and included action potential (AP) half-width, maximum AP firing rate, sag ratio, rebound potential and input resistance. AP half-width was defined as the time difference between the rise and decay phase measured at 50% amplitude of the first elicited AP. AP amplitude was calculated as the difference in voltage between the AP peak and threshold, defined as the point where the voltage strongly accelerate from rest (Sekerli et al., 2004). The maximum AP firing rate observed during depolarizing current step stimuli (100-500 pA) was defined as the maximum AP rate of the cell. The sag ratio was calculated from the -300 pA current sweep as the ratio between steady state and maximum membrane voltage deflection. Rebound potential was defined as the difference between baseline and the maximal depolarizing voltage deflection immediately following the strongest hyperpolarizing current sweep. Input resistance was estimated by taking the voltage deflection between baseline and steady state during the weakest hyperpolarizing current, divided by the injected current (Ohm’s law: R = V/I). Data analysis was performed in Igor Pro and using custom written scripts in Matlab (MathWorks, Natick, Massachusetts, USA). Recorded interneurons were categorized into three main groups. This classification was performed by doing a k-means cluster
analysis in Matlab, using input resistance, sag ratio, maximum AP rate, AP half-width and rebound potential as input parameters. All input values were standardized using the z-score function in Matlab. The number of predefined clusters were set to three \((k=3)\) after inspection of the elbow plot, showing the relationship between the sum of squared errors and the number of clusters \((k)\).

**Surgery**

Mice of the MEC13-53A enhancer strain expressing the tetracycline-controlled transcriptional activation \((tTA)\) system in RE+ cells in LII of the EC were used for the optogenetic experiments (Blankvoort et al., 2018). 23 mice were injected with tTA dependent AAV viruses \((AAV2/9-TRE-tight-oChIEF-Citrine)\) was used for DG/CA3 injections, AAV2/1-TRE-tight-oChIEF-Citrine was used for LEC injections, and control animals for histology were injected with AAV2/1-tetO GFP. The animals were anesthetized with isoflurane in an induction chamber \((4\%, \text{Nycomed Zurich, airflow 1 L/min})\), after which they were moved to a surgical mask on a stereotactic frame \((\text{Kopf Instruments, Tujunga CA})\). The animals were placed on a heating pad \((37 \, ^\circ \text{C})\) to maintain stable body temperature throughout the surgery. The dorsal surface of the head was shaved with an electrical shaver, and eye ointment was applied to the eyes of the animal to protect the corneas from drying out. The animals were injected subcutaneously with buprenorphine hydrochloride \((0.1 \, \text{mg/kg, Temgesic}^\circ \text{ Indivior})\), meloxicam \((1 \, \text{mg/kg, Metacam}^\circ \text{ Boehringer Ingelheim Vetmedica})\) and bupivacaine hydrochloride \((\text{Marcain}^TM \, 1 \, \text{mg/kg, AstraZeneca})\), the latter at the incision site. The head was fixed to the stereotaxic frame with ear bars, and the skin overlying the skull at the incision site was disinfected with ethanol \((70\%)\) and iodide before a rostro-caudal incision was made. Anterior-posterior coordinates were measured from bregma or the posterior transverse sinus, mediolateral coordinates were measured from the mid-sagittal sinus, and dorso-ventral coordinates were measured from the surface of the brain. A craniotomy was made around the approximate coordinate for the injection, and precise measurements were made with the glass capillary used for the virus injection. Viruses were injected...
with a nanoliter injector (Nanoliter2010, World Precision Instruments, Sarasota, FL, USA) controlled by a microsyringe pump controller (Micro4 pump, World Precision Instruments). 400-500 nL of virus was injected with a speed of 40 nL/min. After completion of the injection, we waited 10 minutes to give the virus time to diffuse before retracting the capillary. Finally, the wound was rinsed and the skin was sutured. The animals were left to recover in a heating chamber, before being returned to their home cage, where their health was checked daily.

**Whole-cell voltage clamp recordings**

After 2 weeks, MEC13-53A mice that had undergone surgery were sacrificed and acute 400 µm semicoronal slices were prepared as described above (see [acute slice preparation](#)). Experimental conditions were the same as for whole-cell current clamp recordings, except here LII principal cells were recorded in whole-cell voltage clamp mode with the following intracellular solution: 117 mM cesium gluconate, 13 mM CsCl, 2 mM MgCl₂, 10 mM HEPES, 10 mM Na₂-phosphocreatine, 0.3 mM Na-GTP 4 mM Mg-ATP, 5 mM QX314-Cl. Biocytin (Iris Biotech, Marktredwitz, Germany) was added at a concentration of 5 mg/mL (0.5%). Recordings were performed with an EPC 10 Quadro USB amplifier, controlled by the acquisition software Patchmaster. Acquired data were low pass Bessel filtered at 4 kHz and digitized at 10 kHz. Only recordings with series resistance ≤ 25 MΩ were included (median, 18 MΩ; 25ᵗʰ-75ᵗʰ percentiles, 13-22 MΩ). Series resistance was continually monitored and compensated. Series resistance compensation was carried out by increasing the amount of compensation until oscillations were observed in the current trace, and then reduced and maintained at a level just below this point (65-85% compensation). Remaining uncompensated series resistance was estimated to produce an error in the holding voltage of usually less than a millivolt (median, 0.45 mV; 25ᵗʰ – 75ᵗʰ percentiles, 0.26 – 0.73 mV). In all experiments, excitatory postsynaptic currents (EPSCs) or inhibitory postsynaptic currents (IPSCs) were recorded by clamping the cell potential near the reversal potential for excitatory currents \( E_{\text{exc}} \approx 0 \text{ mV} \) or inhibitory, chloride-mediated currents \( E_{\text{Cl}} = -49 \text{ mV} \).
respectively. Recordings were corrected for the liquid junction potential (12.1 mV for the cesium based internal solution, calculated using the junction potential calculator (JPCalc) plugin in pCLAMP (Molecular devices); the reversal potentials were subsequently confirmed experimentally). In several experiments, bath application of bicuculline (10 µM, Sigma-Aldrich) or combined DNQX (10 µM, Tocris Bioscience, Abingdon, UK) and APV (50 µM, Tocris Bioscience) was used to block GABAergic or glutamatergic synaptic transmission, respectively.

**Laser scanning photostimulation and voltage clamp data analysis**

Photostimulation of oChIEF positive fibers was carried out with a 473 nm laser controlled by a UGA-42 GEO point scanning system (Rapp OptoElectronic, Wedel, Germany), and delivered through a 20x/1.0 NA WI objective (Zeiss Axio Examiner.D1). Laser pulses had a beam diameter of 35 µm and a duration of 1 ms. The tissue was illuminated with individual pulses at a rate of 1 Hz in a 4x5 grid pattern. Laser intensity (1.5-5.0 mW) was adjusted for each recording to evoke small, repeatable inward currents (EPSCs), and the same laser pulses were used to evoke both monosynaptic EPSCs and disynaptic IPSCs. Current traces from individual stimulation spots were averaged over 5-15 individual sweeps to create an average response for each point in the 4x5 grid. Deflections of the average current trace exceeding 10 standard deviations (±10 SD) of the baseline were classified as synaptic responses. Deflections that did not meet our inclusion criterion were invariably less than 10 pA in amplitude. Postsynaptic current amplitudes were calculated as the difference between the peak of the evoked synaptic current and the baseline current measured before stimulus onset. Postsynaptic current latency was defined as the time interval between light onset and the point where the current trace exceeded 10% amplitude. Data analysis was performed using custom written scripts in Matlab.

**Histology**
After electrophysiological recordings, the 400 µm thick brain slices were put in 4% paraformaldehyde (PFA, pH 7.4, Merck Chemicals) for 48 hours at 4 °C. Slices containing cells filled with Alexa Fluor dyes were dehydrated after the fixation (see details below) and prepared for confocal microscopy. Slices containing biocytin filled cells were stained with streptavidin conjugated to a fluorescent tag (see details below). A selection of slices from current clamp experiments were stained with antibodies against reelin and some of the slices containing fast spiking cells were stained with antibodies against parvalbumin (see immunohistochemistry for complete protocols). All slices from optogenetic experiments were stained with antibodies against reelin and GFP.

To confirm the specificity of transgenic labelling and assess the overlap with RE+ positive cells in EC LII, four transgenic MEC13-53A mice were injected with AAV-tetO-GFP and another four with AAV-TRE-tight-oChIEF-Citrine. Two weeks after injection these animals were anesthetized with isoflurane before being killed with a lethal intraperitoneal injection of pentobarbital (100 mg/kg, Apotekerforeningen). Animals were transcardially perfused using a peristaltic pump (World Precision Instruments), first with Ringer solution (0.85% NaCl, 0.025% KCl, 0.02% NaHCO3) and subsequently with freshly prepared 4% PFA. The brains were removed from the skull, post-fixed in PFA overnight and put in a cryo-protective solution containing 20% glycerol, 2% dimethyl sulfoxide (DMSO) diluted in 0.125 M phosphate buffer (PB). A freezing microtome was used to cut the brains into 40 µm thick sections, in four equally spaced series. One series from each brain was immunostained against GFP and reelin (see details below). After staining the sections were mounted in Tris-HCl on Superfrost™Plus microscope slides (Thermo Fisher Scientific), cleared in toluene (VWR) and cover slipped in a mixture of toluene and entellan (Merck Chemicals).

Immunohistochemistry

Reelin and GFP
Thick (400 μm) slices from electrophysiological recordings selected for immunohistochemical staining were first washed 4x15 minutes in 0.125 M PB at room temperature, before going through an antigen retrieval process, in which the slices were incubated for 1 hour at 60 °C in PB. After this the sections were permeabilized 2x10 minutes in PB+0.5% Triton X-100 (PBT; Merck Chemicals) and pre-incubated in a blocking medium containing PBT and 5% Normal Goat Serum (NGS, Abcam Cat# ab7481, RRID: AB_2716553) for 90 minutes at room temperature. After this, the tissue was incubated with primary antibodies, chicken anti-GFP 1:400 and/or mouse anti-reelin 1:800 (anti-GFP: Abcam Cat# ab13970, RRID: AB_300798, recombinant full-length protein corresponding to GFP as the immunogen; anti-reelin: Millipore Cat# MAB5364, RRID: AB_2179313, recombinant reelin amino acids 164-496 clone G10 as the immunogen), and left on a stirrer for 72 hours at 4 °C. After this, the sections were washed 4x15 minutes in PBT at room temperature and incubated in secondary antibody 1:400 (goat anti-chicken AF488, Thermo Fisher Scientific Cat# A-11039, RRID:AB_2534096, and goat anti-mouse AF546, Thermo Fisher Scientific Cat# A-11003, RRID:AB_2534071 ) overnight at 4 °C. Thick sections containing cells filled with biocytin were also stained with streptavidin conjugated to a fluorescent tag (1:600; AF488/633, Thermo Fisher Scientific Cat# S32354, RRID: AB_2315383/ Thermo Fisher Scientific Cat# S-21375, RRID: AB_2313500) during the secondary antibody incubation. After secondary antibody incubation, all sections were washed 4x15 minutes in PB at room temperature before being dehydrated in increasing ethanol concentrations (30, 50, 70, 90, 100, 100%, 10 min each) at room temperature before incubation in a 1:1 mixture of 100% ethanol and methyl salicylate (VWR chemicals), this rendered the slices completely transparent. Finally, slices were stored in methyl salicylate. Staining against GFP and reelin was performed simultaneously in sections were both antigens were present. Thin (40 μm) sections from MEC13-53A mice used for controls were stained according to the protocol described above, but the antibody incubations were shortened. The primary antibody incubation lasted 48 hours at 4 °C and the secondary antibody three hours in room temperature.
Slices selected for parvalbumin staining were first washed 4x15min in PB at room temperature, before being permeabilized 4x15min in PB with 1% TrX at room temperature. The slices were preincubated in PB + 1% TrX and 10% NGS at room temperature, before being incubated in primary antibody, rabbit anti-parvalbumin 1:1000 (Swant Cat# PV 25, RRID: AB_10000344/ Swant Cat# PV27, RRID: AB_2631173, recombinant rat parvalbumin as the immunogen), on a stirrer for 24 hours at 4 °C. Next, the slices were washed 4x15min in PB at room temperature, and incubated in secondary antibody, goat anti-rabbit AF633 1:400 (Thermo Fisher Scientific Cat# A-31576, RRID: AB_2536186). Finally, slices were dehydrated and cleared in methyl salicylate.

Laser scanning confocal microscopy

Thick 400 µm slices from patch clamp recordings were imaged with a laser scanning confocal microscope (Zeiss Meta 510/ Zeiss LSM 880, Carl Zeiss). The slices were mounted in custom made metal well slides with methyl salicylate and cover slipped. AF405 was excited with a 405 diode laser (emission: BP 420-480/410 - 483), AF488 and GFP was excited by the 488 line of an Argon laser (emission: BP 505-550/ 490 - 543), AF546 by a DPSS 568 laser line (emission: BP 575-615 IR/ 570 - 623) and AF633 by a HeNe 633 laser line (emission: LP 650 /635 - 735). The main beam splitter for the Zeiss Meta 510 was a HFT 405/488/561/633/KP 725 and for the LSM 880 an MBS 488/561/633. First, overview images of the tissue were taken with low magnification (Plan-Apochromat 10x, NA 0.45 and Plan-Apochromat 20x, NA 0.8) to get the overall morphology of the cells, and to confirm their location in LII of LEC. After this, the slices were imaged with high magnification (Plan-Apochromat 40x oil, NA 1.3 and Plan-Apochromat 63x oil, NA 1.4) to perform morphological analyses of each cell. All images were acquired with 8 bit depth. Both overview images and high magnification images were obtained as z-stacks that included the whole extent of each recorded cell to recover the full cell morphology. For double staining experiments, confocal images through the soma of each cell were taken to
determine overlap of staining. Images for figures were exported to JPG files using the ZEN Black and
Blue software belonging to the confocal microscopes, and the contrast and brightness of the images
were adjusted in Adobe Photoshop (Adobe Photoshop CC, Adobe Systems Software). 3D
reconstructions of cells were made in the reconstruction software AMIRA (FEI, Thermo Fisher,
Hillsboro, OR, USA) using the custom skeleton toolbox (Schmitt et al., 2004; Evers et al., 2005).

**Fluorescent slide scanner images**

Mounted 40 µm thin sections were imaged using an Axio ScanZ.1 fluorescent scanner, equipped with
a 20x objective, Colibri.2 LED light source and a quadruple emission filter (Plan-Apochromat 20x, NA
0.8, ex. 488/546, em. 405/488/546/633, Carl Zeiss Microscopy). Images for figures were exported to
JPG files using the ZEN Blue software belonging to the scanner system, and contrast and brightness of
the images was adjusted in Adobe Photoshop.

**Morphological analysis of LEC LII recorded cells**

Morphological classification of the principal cells recorded in our study was done by visually inspecting
z-stacks and projection views of raw images acquired by confocal microscopy, at 20x, 40x or 63x
magnification. The morphology of each neuron was evaluated independently by two people, and
classified in accordance with previous descriptions of LEC LII principal cells (Tahvildari and Alonso,
2005; Canto and Witter, 2012a). Fan cells were recognized by a large, round soma, an extensive
dendritic tree fanning towards the pial surface and a lack of or rudimentary, basal dendritic tree.
Multipolar cells were large cells with a round soma, with dendrites that radiated in all directions.
Pyramidal cells had a triangular soma oriented perpendicular to the pial surface, and a large apical
dendrite reaching towards the pial surface before splitting into multiple branches. Pyramidal cells had
basal dendritic trees of varying extents. Oblique pyramidal cells had pyramidal cell morphology but
were tilted approximately 45° to the pial surface such that the primary apical dendrites extended at
an angle to the pial surface. If we were unable to determine the morphology of a principal neuron, it was excluded from the dataset.

In the few cases where the principal cell main axon was cut closer than 100 µm from the soma, possible synaptic connections arising from the activity of this cell were not included for further analysis. The inter-somatic distance of the recorded cells was measured as the Euclidian distance between the cell bodies.

**Experimental design and statistical analyses**

For our experiments, we used slices from 49 rats and 4 mice for whole-cell current clamp recordings, as well as 15 transgenic MEC13-53A enhancer strain mice for whole-cell voltage clamp recordings. 8 transgenic animals were also to control the specificity of the transgene expression of the MEC13-53A mouse line. The number of slices and cells recorded from each animal varied. All recordings were made in LII of the dorsolateral LEC close to the rhinal fissure. All statistical analyses were carried out using IBM SPSS statistics 25. Data were tested for normality using Q-Q plots, distribution histograms and the Shapiro-Wilk test before conducting statistical tests. All of the tested data were non-normally distributed, and some data sets were too small to perform a reliable test for normality and homogeneity of variance. We used the Mann-Whitney U test when comparing variables between two groups, and for comparisons of multiple measures, we performed a Kruskal-Wallis H test. Data was considered significant at p ≤ 0.05. Median and 25th-75th percentiles are reported for all data distributions.

**RESULTS**

**Polysynaptic connectivity between fan cells in layer II of LEC**

To probe the synaptic connectivity of LII principal cells, we performed simultaneous whole-cell current clamp recordings from up to four cells located in dorsolateral LEC in acute semicoronal slices.
of putative principal cells (inter-somatic distance; median, 42.95 µm; 25\textsuperscript{th}-75\textsuperscript{th} percentiles, 29.23-61.04 µm) were selected for recording. Our recordings included all four main principal cell types in LII of LEC, defined by somato-dendritic morphology acquired by post hoc visualization of the recorded cells. There are conflicting reports about the correlation between morphological principal cell type identity and electrophysiological properties in LEC LII (Tahvildari and Alonso, 2005; Canto and Witter, 2012a; Leitner et al., 2016). This potential conflict likely reflects two possible sampling variations. One could be a variability in intrinsic membrane properties of cells positioned differently along the medial to lateral extent of the LEC (Canto and Witter, 2012a). The other might reflect the chemical nature of the neuron, where intrinsic membrane properties differ between RE+ cells and calbindin positive cells, but not between different morphological classes of RE+ principal cells (Leitner et al., 2016). Since we recorded mainly from RE+ neurons, we chose to use dendritic features to determine the principal cell type. To describe the local connectivity between the different groups of principal cells accurately, recorded cells with incompletely recovered morphologies were excluded. The data set contains recordings from 292 principal cells from 48 Long Evans rats and 15 principal cells from four C57 mice. We tested in total 551 principal cell to principal cell connections, derived from 517 connections in the Long Evans data set and 34 connections in the C57 data set.

Most recorded principal cells (n=189/307) had morphologies typical of fan cells (Figure 1A). These cells usually fired late and few spikes during weak depolarizing current steps, and showed spike frequency adaptation during stronger depolarizing stimuli (Figure 1B). In all experiments, we evoked action potentials by injecting short depolarizing current pulses sequentially in each recorded cell, while observing the resting membrane potentials of the other cells to screen for synaptic connections (Figure 1C). We observed a diversity of postsynaptic potentials in individual fan cells upon activation of neighboring fan cells. A single fan cell pair (n=1/215 or 0.5%) was presumably monosynaptically connected through a short-latency depolarizing connection (latency < 3 ms), but depolarizing
postsynaptic potentials of longer latencies were observed more frequently (n=4/215, or 1.9%; Figures 1C, 1D). We also encountered hyperpolarizing potentials (n=7/215, or 3.3%) of variable latencies in fan cells when activating another nearby fan cell at high frequencies (40-70 Hz; Figures 1E, 1F), or when activating multiple fan cells simultaneously. We interpret these potentials as disynaptic inhibitory postsynaptic potentials (IPSPs) arising from the activation of intermediate inhibitory interneurons. In addition, one connection between fan cells could not with certainty be categorized as mono- or polysynaptic, and was thus marked as unclassifiable. Amplitudes of fan to fan postsynaptic potentials were invariably less than 1 mV following 10 Hz or 70 Hz presynaptic stimulation (Figure 1G).

Taken together, our recordings indicate that putative disynaptic connectivity is the preferred mode of communication between fan cells.

Our multi-cell recordings routinely contained one or more of the other LII principal cell types, allowing us to explore the number and amplitudes of synaptic connections of fan cells with these cells (Figure 2A). Presumed monosynaptic connections were detected between fan cells and pyramidal cells (fan to pyramidal: n=2/29, or 6.9%; pyramidal to fan: n=1/31, or 3.2%) and a long-latency potential, putative polysynaptic, was found from a multiform to a fan cell (n=1/28, or 3.6%). Finally, we tested the interconnectivity of the other principal cell types (Figure 2B). Presumed monosynaptic connectivity was found between cells belonging to the same principal cell class (multiform cells: n=1/22, or 4.5%; pyramidal cells: n=1/34, or 2.9%; oblique pyramidal cells: n=1/14, or 7.1%), and on a single occasion a long latency depolarizing potential was detected in a pair of simultaneously recorded multiform cells (n=1/22, or 4.5%). Furthermore, presumed monosynaptic connections were present from pyramidal to multiform cells (n=2/18, or 11.1%), and onto oblique pyramidal cells from a multiform (n=1/9, or 11.1%) or a pyramidal cell (n=1/14, or 7.1%).

Three main groups of interneurons in layer II of LEC
Noticing that long-latency synaptic events sometimes occurred when recording from pairs of fan cells, we next aimed to explore the connectivity between principal cells and inhibitory interneurons in LEC LII. We recorded small-sized somata of putative interneurons situated in close proximity to simultaneously recorded principal cells. Interneurons were distinguishable from principal cells by their electrophysiological properties, particularly by their larger spike afterhyperpolarizations and higher firing rates (Kawaguchi et al., 1987; Jones and Buhl, 1993; hu et al., 2010). Morphological recovery of the recorded cells revealed that these cells were typical interneurons with extensive local axonal trees and very low dendritic spine densities, or absence of spines. We used a k-means cluster analysis to classify LEC LII interneurons into fast-spiking cells (FS), non-fast spiking cells (nFS) and low threshold spiking cells (LTS) based on electrophysiological properties (Figure 3).

Fast-spiking cells were the most frequently encountered interneuron type in our data (n=29/61; Figures 3A, 3B). They had high maximum firing rates and short action potential half-widths, two features which distinguished them from the two other interneuron classes (Figures 3A-3C, Table 1, P < 0.0001, Kruskal-Wallis H test). A selection of FS cells were tested immunohistochemically, and a subset of these (n=7/17, or 41.2%) stained positive for the calcium-binding protein parvalbumin (PV, Figure 4), a common marker for FS cells (Hu et al., 2014). Low threshold spiking cells (n=13/61) could be separated from nFS cells (n=19/61) based on high input resistance, together with their tendencies to show sag potentials and large rebound potentials that occasionally reached firing threshold after hyperpolarizing current steps (Figures 3A, 3D, Table 1, P < 0.0001, Kruskal-Wallis H test).

Fan cells receive extensive intralaminar inputs from fast-spiking and non-fast spiking inhibitory cells

After establishing that LEC LII contains FS, nFS and LTS cells, we explored their connectivity with fan cells (Figure 5). We found robust connectivity onto fan cells from nearby FS (Figures 5A-5C) and nFS
cells (Figures 5D-5F). The most prevalent connectivity was from FS cells to fan cells (n=17/28, or 60.7%, Figures 5A-5C, 5H), and connections from fan cells to FS cells were also common (n=5/26, or 19.2%; Figures 5G, 5H). Fan cells were frequently targeted by nFS cells (n=6/21, or 28.6%; Figures 5D-5F, 5H), but connectivity in the opposite direction was sparse (n=1/18, or 5.6%; Figure 5H). We found one reciprocal connection in LTS and fan cell pairs (LTS to fan, n=1/11, or 9.1%; fan to LTS, n=1/7, or 14.3%; Figure 5H).

We also tested the connectivity of the three interneuron classes with the other principal cell types in LII of LEC (Figure 6). FS cells had connections to pyramidal (n=3/11, or 27.3%), oblique pyramidal (n=4/8, or 50%) and multiform cells (n=2/6, or 33.3%), and received connections from pyramidal (n=1/11, or 9.1%), oblique pyramidal (n=1/7, or 14.3%) and multiform cells (n=1/6, or 16.7%). Cells of the nFS class had connections onto pyramidal cells (n=1/5, or 20%) and oblique pyramidal cells (n=4/8, or 50%), and connections from pyramidal and oblique pyramidal cells onto nFS cells were also frequent (n=2/5, or 40% and n=3/5, or 60%, respectively). LTS cells was the interneuron group that had the sparsest connectivity with principal cells. We did not find connections from LTS cells to multiform, oblique pyramidal or pyramidal cells, but detected a single connection onto an LTS cell from an oblique pyramidal cell (n=1/7, or 14.3%).

**Fan cells receive different types of inhibition from fast-spiking and non-fast spiking cells**

We noticed that stimulation of FS and nFS cells lead to IPSPs in fan cells varying in strength, temporal dynamic and polarity. Driving FS cells usually induced short-latency (median, 2.10 ms; 25th – 75th percentiles, 1.55-2.60 ms) IPSPs in fan cells at all stimulation frequencies (n=14/17 connections; Figure 7A), whereas driving nFS cells resulted in small, slow IPSPs that were usually only evident at high frequency stimulation (n=4/6 connections; Figure 7A). High frequency activation (70 Hz) of FS cells produced fast, transient responses compared to those evoked by nFS cell stimulation (latency to peak; FS, median, 136.00 ms; 25th – 75th percentiles, 102.6-175.45 ms and nFS, median, 394.25 ms; 25th –
75th percentiles, 225.15-601.38 ms; Mann Whitney U test, U=2.00 P=0.000079; Figure 7B; half-width; FS median, 241.90 ms; 25th – 75th percentiles, 221.20-296.00 ms and nFS, median, 470.7 ms; 25th – 75th percentiles, 271.53-667.48 ms; Mann Whitney U test; U=15.00, P=0.01; Figure 7C). Furthermore, larger IPSP amplitudes were observed in postsynaptic fan cells following FS stimulation than after nFS stimulation (70 Hz stimulation; FS, median, 2.0 mV; 25th – 75th percentiles, 0.63-3.75 mV and nFS, median, 0.44 mV; 25th – 75th percentiles, 0.36-0.82 mV; Mann Whitney U test, U=15.5, P=0.01; Figure 7D). Interestingly, the IPSPs evoked by FS cell activity differed in polarity from IPSPs derived from nFS cell stimulation. In all cases, FS cell stimulation lead to depolarizing rather than hyperpolarizing events in the postsynaptic fan cell (Figure 7A). This depolarizing effect could be explained by the reversal potential for chloride, which for our biocytin containing internal solution was -47.0 mV, and thus more depolarized than the resting membrane potential of the fan cells (-70.3 mV, -72.6 - -68.0 mV in FS cell recordings and -70.0 mV, -71.5 - -66.5 mV in nFS cell recordings; median, 25th-75th percentiles). Application of the GABA_A antagonist bicuculline abolished FS induced IPSPs in all tested cases (n=5), and the IPSPs were partially recovered after bicuculline removal, confirming that they were GABA_A dependent responses (Figure 7E). Activation of nFS cells produced slow, lasting hyperpolarizing events, sometimes preceded by a brief depolarizing event. Given the depolarized chloride reversal potential in our experiments, this biphasic response likely stems from different ionic mechanisms, potentially from GABA_A and GABA_B mediated currents (Nicoll et al., 1990). We encountered connections between nFS cells and fan cells only sporadically, and could test these connections pharmacologically on a single occasion only. In this case, the IPSP was partially sensitive to bicuculline. More specifically, the fast depolarizing potential was blocked, whereas the slower hyperpolarizing component was less affected (Figure 7F). Taken together, our data indicate that FS and nFS cells exert two different inhibitory effects on fan cells in LII of LEC.

All classes of layer II principal cells project to the dentate gyrus and CA3
In consideration of the low number of connections found in our multi-cell recordings, we aimed to optimize the detection of synaptic connections by introducing an oChIEF variant of the light-activating protein channelrhodopsin2 exclusively in EC LII excitatory RE+ cells. To this end, we used the MEC13-53A mouse line, which carries the tetracyclin-controlled transcriptional activation (tTA) factor in a subpopulation of RE+ cells (Blankvoort et al., 2018) and injected a tTA dependent adeno-associated virus carrying a fused oChIEF-citrine protein (AAV2/9-TRE-tight-oChIEF-Citrine) into the dorsal DG/CA3. This led to specific retrograde labeling of LII DG/CA3-projecting cells dorsally in EC (Figure 8A). We first performed whole-cell recordings in acute semicoronal slices to characterize the DG/CA3-projecting transgenic cell population. Fluorescent cells were patched and voltage clamped at -50 mV during photostimulation with brief light pulses (473 nm light, 1 ms duration). Post hoc morphological recovery of the recorded cells revealed that DG/CA3-projecting cells encompass fan (n=10/20), multiform (n=6/20), pyramidal (n=2/20) and oblique pyramidal cells (n=2/20) (Figures 8B, 8C). This observation is supported by immunohistochemical data from the rat, showing that RE+ neurons are found within all classes of principal cells in LEC LII (Fjeld, 2015). Reelin has previously been shown to be a reliable marker for DG/CA3-projecting cells both in MEC and LEC (Varga et al., 2010; Leitner et al., 2016). OChIEF expressing cells displayed inward currents with sub-millisecond onset latency (median, 0.44 ms; 25th – 75th percentiles, 0.38 – 0.46 ms; Figures 8C, 8D). Inward current amplitudes were variable across cells (median, -211.1 pA; 25th – 75th percentiles, -477.6 - - 90.5 pA) but all occurred with negligible delay following laser stimulation (Figures 8C-8E). Short latency currents were resistant to glutamatergic (DNQX + APV, n=6 cells from three animals) and GABAergic (bicuculline, n=6 from three animals) synaptic blockers (data not shown), indicating that they were direct photocurrents resulting from oChIEF activation. Our findings demonstrate that RE+ cells are present within all morphological classes of LII principal cells. Moreover, all morphological groups are present in the DG/CA3-projecting population of LEC LII cells, though with a preferential origin from fan cells. The network of layer II reelin immunoreactive cells is dominated by disynaptic inhibition.
After the morphological characterization of DG/CA3-projecting cells, we next set out to map the local functional connectivity of these cells onto neighboring fan cells using an optogenetic approach. As MEC LII RE+ cells also project to the DG/CA3 we first wanted to check whether projections from these cells were inducing activity in LII of LEC. To test this, we injected the retrograde AAV2/9-TRE-tight-oChIEF-Citrine directly into LEC. This approach revealed a near complete lack of virally transduced MEC cells, demonstrating that transgenically labeled LII MEC cells do not project to LEC (Figure 9). Hence, AAV2/9-TRE-tight-oChIEF-Citrine delivery into dorsal DG/CA3 enabled optogenetic manipulation restricted to local LEC LII circuitry.

Following such injections into the DG/CA3 we targeted large cell bodies in LEC LII that did not express oChIEF but were located near citrine-labeled fibers to investigate the connectivity of LEC LII cells (Figures 10A, 10B). During laser scanning photostimulation of surrounding oChIEF expressing fibers, recorded fan cells (n=33/56) were alternately voltage clamped at potentials close to the excitatory or inhibitory reversal potentials (≈0 mV or ≈-49 mV, respectively) to isolate excitatory and inhibitory postsynaptic currents (EPSC and IPSC, respectively). These experiments revealed that fan cells receive strong inhibitory input (outward currents; Figures 10B-10D) derived from the activity of nearby LEC LII excitatory cells, and that this inhibition occasionally is accompanied by excitatory events (inward currents; Figures 10B-10D). These currents had considerably longer latencies than those observed in oChIEF expressing cells, indicating that they derived from synaptic activity (inward current, 8.45 ms, 7.84 - 9.23 ms; outward current, 9.32 ms, 8.28 – 10.19 ms; median, 25th – 75th percentiles). Inhibitory events were detected in more than half (n=19/32, or 59%) of all recorded fan cells (Figure 10C). We did not record fan cells that exclusively got excitatory inputs, but fan cells (n=6/24, or 25%) that solely displayed inhibitory currents in response to photostimulation were present (Figures 10B, 10D).

Given that we did not observe any putative GABAergic cells among the recorded oChIEF expressing cells (Figures 9A, 9B), the outward currents observed in our data were likely disynaptic inhibitory
events driven by the concurrent activation of oChIEF excitatory fibers. In line with a disynaptic
coupling, we observed in fan cells receiving convergent excitatory and inhibitory input (n=6) a short,
but frequent, delayed onset for IPSCs relative to EPSCs evoked from the same laser stimulation spot
(Figures 10E, 10F). To prepare for a pharmacological test of these inhibitory connections, we sought
to increase the efficacy of optogenetic stimulation by injecting AAV2/1-TRE-tight-oChIEF-Citrine
directly into the LEC (Figure 8). In these experiments, we confirmed the disynaptic inhibitory nature
by demonstrating that outward currents could be reversibly blocked by bath application of either
bicuculline (n=11 cells from five animals; Figure 10G) or DNQX/APV (n=10 cells from four animals;
Figure 10H).

We occasionally recorded from other principal cell types in LII after virus injection in the DG/CA3. Our
findings show that multiform (n=10/17), oblique pyramidal (n=2/2) and pyramidal cells (n=2/4) all
receive local inhibitory input when driving other RE+ LII excitatory cells. Excitatory inputs to multiform
(n=5/12) and pyramidal cells (n=1/2) were also present, whereas oblique pyramidal cells were
incompletely tested for excitatory connectivity.

Collectively, our voltage clamp data confirm that fan cells receive sparse excitatory inputs from
surrounding LII RE+ excitatory cells, which mainly include fan cells but also pyramidal, oblique
pyramidal and multiform cells. However, the prevailing local input to fan cells is disynaptic inhibition,
which is observed more frequently than excitatory inputs. This demonstrates that the RE+ principal
cell circuit in LEC LII is dominated by local recurrent inhibition.

**DISCUSSION**

Our data collected from a combination of optogenetic stimulation experiments and multi-cell
recordings point to sparse excitatory connectivity in the local network of fan cells. Several studies have
recently emphasized low or nearly absent excitatory connectivity between pairs of stellate cells and
between pairs of pyramidal cells in LII of MEC (Dhillon and Jones, 2000; Couey et al., 2013; Pastoll et al., 2013; Fuchs et al., 2016; Winterer et al., 2017), and between pyramidal cells in the superficial layers of the presubiculum (Peng et al., 2017). These observations collectively indicate that sparse excitatory connectivity is a recurring feature in LII of the parahippocampal region, present within at least four cell populations across MEC, LEC and the presubiculum. This is in contrast to previous reports of higher connectivity among pyramidal cells in LIII (between 5-10%) and LV (around 12%) of MEC (Dhillon and Jones, 2000) (Winterer et al., 2017). Furthermore, data on micronetworks in L2 of primary sensory cortices show a higher probability of connectivity between principal cells (around 9%) (Lefort et al., 2009; Jouhanneau et al., 2015; Seeman et al., 2018).

Putative disynaptic connections were encountered more frequently than monosynaptic connections in our simultaneous recordings of fan cells. In line with this observation, optogenetic activation of LEC LII excitatory cells routinely elicited disynaptic inhibitory events in a majority of the recorded fan cells, suggesting a strong indirect functional connection mediated by interneurons. This is similar to MEC stellate cells where local excitatory connections primarily target neighboring interneurons, leading stellate cells to interact primarily through the activity of these cells (Couey et al., 2013). Although such couplings have not been unequivocally demonstrated for pyramidal cells in MEC and the presubiculum, their prevalent connectivity with local inhibitory cells suggests that similar functional interactions also apply to these cell groups (Fuchs et al., 2016; Peng et al., 2017; Nassar et al., 2018).

Hence, our data point to a fan cell local circuit architecture with a similar fundamental wiring motif as the MEC stellate cell, and possibly pyramidal cells in MEC and the presubiculum. Even though inhibitory disynaptic connectivity seems widespread both for MEC stellate and LEC fan cells, the underlying interneuron networks accountable for this inhibition might involve different cell types and modes of action. Our data suggest an important role for FS interneurons in mediating communication within the fan cell network through quick and robust presumed perisomatic inhibition. This input is
accompanied by slow, but sustained, inhibitory synaptic inputs provided by nFS cells, similar in nature to inhibitory inputs targeting distal dendritic compartments (Miles et al., 1996; Silberberg and Markram, 2007). Similar to fan cells, MEC stellate cells are primarily targeted by FS interneurons (Couey et al., 2013; Pastoll et al., 2013; Fuchs et al., 2016). However, stellate cells receive additional inhibition mainly from LTS cells, and not from nFS cells (Fuchs et al., 2016), pointing to a potential important difference in inhibitory inputs between fan and stellate cells.

Although we now know that LEC and MEC develop from different parts of the pallium (Medina et al., 2017), fan cells and stellate cells have been considered counterparts in LII of LEC and MEC respectively, since both are the dominant cell type contributing massively to the projections to DG, CA3 and CA2, and share morphological and immunohistochemical features (Germroth et al., 1989; Tahvildari and Alonso, 2005; Fuchs et al., 2016; Leitner et al., 2016). Our data add to the similarities between these cell types by showing that fan cells are interconnected in a circuit that shares the inhibitory motif characteristic of the MEC stellate cell network (Couey et al., 2013). Notwithstanding the observed differences in inhibitory components participating in the two respective networks, the overall similar coupling of these cell types has important implications for our understanding of ongoing computations underlying EC physiology. More specifically, although there are alternative models for the generation of the grid pattern (Witter and Moser, 2006; Burgess et al., 2007; Kropff and Treves, 2008; Islam and Yamaguchi, 2009; Naumann et al., 2016), an inhibition-dominated continuous attractor network receiving velocity-modulated head-direction inputs has been proposed to be sufficient to generate grid cell firing when navigating in space (Bonnevie et al., 2013; Couey et al., 2013; Pastoll et al., 2013). Support for this premise is present in the networks of MEC and presubiculum, two regions where principal cells are embedded in an inhibition-dominated network (Couey et al., 2013; Peng et al., 2017; Nassar et al., 2018) and contain grid cells (Hafting et al., 2005; Boccara et al., 2010). Our data show that a comparable network structure exists in an adjacent area where grid cells have not been
described (Hargreaves et al., 2005). It is thus conceivable that, given the appropriate afferent input, neurons in LII of LEC will show hexagonal, or at least regularly repeating, firing patterns, domain along dimensions defined by their inputs. In contrast to the pure spatial representation observed in MEC, periodic patterns might arise in LEC to represent complex features of experience (Tsao et al., 2013; Knierim et al., 2014; Constantinescu et al., 2016; Tsao et al., 2018).

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**Figure legends**

![Diagram A](image1)

![Diagram B](image2)

![Diagram C](image3)

![Diagram D](image4)

![Diagram E](image5)

![Diagram F](image6)

![Diagram G](image7)
Figure 1. Polysynaptic connectivity between fan cells. (A) Reconstruction of three LII fan cells (black, magenta and blue) and one LII non-fast spiking (nFS, green) cell recorded simultaneously in an acute semicoronal slice through LEC. Scale bar=20 µm. (B) Electrophysiological profiles of the cells in A resulting from hyperpolarizing and depolarizing current steps. The three fan cells all show similar characteristic responses that differ from responses of the nFS cell. Traces are color coded according to the color of the cells in A. (C) Average membrane potential traces showing a single depolarizing postsynaptic potential in the blue fan cell arising from an action potential in the magenta fan cell (second panel). Average membrane potential is indicated for each cell. The connectivity protocol is illustrated with dashed and solid lines indicating unconnected and connected cells, respectively. Traces are color coded referring to the cells in A. (D) Enlarged time scale of the connection in C. Average trace (blue) superimposed on individual traces (grey) showing a long delay between the presynaptic spike (magenta) and the onset of the postsynaptic potential. Dotted line indicates the peak of the presynaptic action potential. (E) Fan cells are connected with neighboring fan cells through disynaptic inhibitory connections. Shown is a recorded cluster (different from the cluster shown in A) containing three fan cells (cells 1, 2 and 4) and one pyramidal cell (cell 3). Brief inhibitory postsynaptic potentials were detected in fan cell 2 (black) and fan cell 4 (black) in response to a presynaptic 70 Hz action potential train in fan cell 1 (magenta). Average traces (black) are superimposed on individual traces (grey). Average membrane potential is indicated for each cell. Cells were recorded and filled with K-glucuronate containing different spectral variants of Alexa Fluor dyes. (F) Fan cells are interconnected through slow disynaptic inhibitory connections. Displayed is a cluster of four fan cells (cells 1, 2, 3 and 4). Long-latency, slow inhibitory postsynaptic potentials were detected in fan cell 2 (black) and fan cell 3 (black) in response to a presynaptic 70 Hz action potential train in fan cell 1 (magenta). Average traces (black) are superimposed on individual traces (grey). Average membrane potential is indicated for each cell. Cells were recorded and filled with K-glucuronate internal solution containing biocytin. (G) Postsynaptic potential amplitudes recorded in pairs of fan cells. Amplitudes are shown in response to single stimuli (10Hz, left) and high-frequency presynaptic stimulation (70Hz, right).
Figure 2. Connectivity between principal cells in layer II of LEC. (A) Connectivity between fan cells, and between fan cells and the other classes of principal cells in LII of LEC. Boxes indicate the number of observed connections out of the number of tested connections, as well as postsynaptic potential amplitudes (in mV) following 70Hz presynaptic stimulation. Amplitudes of postsynaptic responses in fan-to-fan connections are shown in figure 1G. (B) Connectivity and average amplitudes of postsynaptic potentials following 70Hz stimulation between multiform, pyramidal and oblique pyramidal cells.
Figure 3. Three main groups of interneurons in layer II of LEC. (A) Examples of electrophysiological profiles from a fast-spiking cell (FS, red), a non-fast spiking cell (nFS, blue) and a low threshold spiking cell (LTS, yellow) in response to depolarizing and hyperpolarizing current step injections. The example traces show that the three cell classes typically display different firing properties after depolarizing current injections (two top traces) and that LTS cells have a characteristic sag potential and rebound spikes following a hyperpolarizing current step (third trace from the top). The bottom trace shows the injected currents. See Table 1 for electrophysiological properties of recorded interneurons. (B) Examples of dendritic morphology from the three interneuron classes. Scale bars are 20 µm for FS cells and 10 µm for nFS/LTS cells. (C) Maximum action potential (AP) firing rate (Hz) plotted against AP half-width (ms), showing that FS cells have shorter spikes and high firing rates compared to nFS and LTS cells. (D) Rebound potential (mV) plotted against sag ratio shows that these two features differentiate nFS cells from LTS cells.
Figure 4. Immunoreactivity of fast spiking cells to the calcium binding protein parvalbumin. (A) Whole-cell cluster recording containing a fan cell, multiform cell and a fast spiking interneuron. Scale bar=50 µm. (B) Reconstruction of the recorded fast spiking cell in A. Scale bar=40 µm (C) Electrophysiological profile of the cell in A and B shows a fast spiking phenotype. (D) Confocal images show the biocytin filled cell body (left) of the cell in A and B, immunostaining to parvalbumin (middle), and overlap between the biocytin filled soma and the parvalbumin signal (right). Scale bar=20 µm.
Figure 5. Fan cells receive extensive intralaminar inputs from fast-spiking and non-fast spiking interneurons. (A) Reconstructed cluster with three fan cells (black) and a fast-spiking (FS) cell (red), scale bar=20 μm. Inset shows position of the cluster in dorsal LEC LII, scale bar=50 μm. Rf=rhinal fissure. (B) Electrophysiological profiles of the cells in A. (C) Responses to elicited action potentials in the four recorded cells (stimulation represented along the diagonal from the top trace in the first panel) showing that the FS cell elicits a response in all three fan cells, but that neither of the fan cells produced a response in the other recorded cells. In some cases, small instantaneous deflections that are artefacts time-locked to the presynaptic current injections can be seen riding on top of the recorded membrane voltage traces (e.g. trace in column 1, row 2). (D) Like A, with two fan cells (black), one non-fast spiking cell (nFS, blue) and one oblique pyramidal cell (Op, grey), scale bars are 25 μm and 100 μm (inset). (E) Electrophysiological profiles of the cells in D. (F) Like C for cells in D/E. nFS cell was presynaptic to all of the other cells, but no other connections were detected. (G) Connectivity from fan cells onto FS cells. Shown are recorded FS postsynaptic potentials in response to a train of 10 action potentials (10Hz, left) and a train of 15+1 action potentials (70 Hz, right) elicited in a presynaptic fan cell. The average postsynaptic trace (red) is superimposed on individual sweeps (grey).
average synaptic amplitudes following 10Hz and 70Hz stimulation in all FS cells receiving input from fan cells (n=5). (H) Summary histogram and diagram of connection probabilities, boxes indicate the number of found connections out of the number of tested connections.

**Figure 6. Summary of connectivity between interneurons and non-fan principal cells.** (A) Paired recording from a fast-spiking (FS) cell and a multiform cell (MC; top), cell reconstructions (Red, FS cell; Black, multiform cell; middle) and summary diagram of connectivity between MCs and all classes of interneurons (INs) (bottom). Red and black traces represent membrane potential recordings from FS and principal cells, respectively. The diagram show an example recording of a unidirectional FS to MC connection after evoking 10 Hz spike trains in the recorded cells. Scale bars are 40 μm for the reconstructed neurons. (B) Like A for oblique pyramidal cells (Opy). The example recording shows a reciprocal connection between the FS and Opy cell. (C) Like A for pyramidal cells (Pyr). The example recording shows a reciprocal connection between the FS and Pyr cell.
Figure 7. Fan cells receive different types of inhibition from fast-spiking and non-fast spiking cells. (A) Inhibitory postsynaptic potentials (IPSPs) in fan cells evoked by increasing stimulation frequencies of fast-spiking (FS, left) and non-fast spiking (nFS, right) cells. Increasing action potential frequency in FS cells results in larger IPSPs in postsynaptic fan cells, whereas the nFS cell elicits IPSPs in the fan cell only during high intensity stimulation. Average traces (black) are superimposed on the individual traces (grey). Dashed lines indicate average resting potentials of recorded fan cells (left, -71.0 mV; right, -66.5 mV). (B) Shorter IPSP latency to peak after FS cell compared to nFS cell stimulation (Mann Whitney U test, U=2.00, P=0.000079). Box shows median (red line) and interquartile range, whiskers indicate min/max values. (C) Same as B, but for IPSP half-width (Mann Whitney U test, U=15.00, P=0.01). (D) Same as B, but for absolute IPSP amplitude (mV) at 70Hz stimulation (Mann Whitney U test, U=15.5, P=0.01). (E) Reversible block of FS to fan cell connections by the GABA_A antagonist bicuculline. Left panel, electrophysiological profiles of a FS and a fan cell. Right panel, stimulation (top) and IPSPs (bottom) before (black), during (blue) and after bicuculline (yellow) treatment. (F) Like E for a nFS cell, showing that the IPSP after nFS stimulation is partially abolished by bicuculline.
Figure 8. All excitatory cell types in LEC LII project to dentate gyrus/CA3 of the hippocampal formation. (A) Retrograde labeling is limited to LII excitatory cells after virus injection into dentate gyrus/CA3 of the MEC13-53A mouse line. Scale bar=200 µm. Insets: mCitrine signal of retrogradely labelled cells colocalize with reelin immunoreactive cells in superficial LII. Scale bar=50 µm. (B) Example cell reconstructions of recorded fan, multiform and pyramidal cells that express oChIEF and reelin. Immunostaining for reelin (yellow; left), citrine/GFP (green, middle) and biocytin-filled somata (cyan, right) are shown in the bottom panels for all cells. Scale bars=20 µm. (C) OChIEF mediated photocurrents recorded from the different LEC LII principal cell types. The fan, multiform and pyramidal cells correspond to the cells shown in B. All cells were voltage clamped at -50 mV. (D) OChIEF expressing cells (n=20) show fast activation after exposure to blue light. (E) Average photocurrent amplitudes of oChIEF expressing cells (n=18).
Figure 9. Reelin positive cells in MEC do not contribute to the activation of reelin cells in LEC. A) Injection of retrograde AAV virus in LEC yields strong viral expression in LEC layer II. B) Very sparse viral expression is seen in MEC following LEC retrograde virus injection. C) Fibers are almost exclusively present in the outer molecular layer (oML) of the dentate gyrus, which contains the projections from LEC. The middle molecular layer (mML), which is known to receive the projections of the MEC, is virtually without labelled axonal fibers. iML – inner molecular layer.
Figure 10. Optogenetic stimulation of LII reelin cells reveals a network dominated by inhibition. (A) Fan cells (magenta) situated near retrogradely labelled oChIEF positive cells (cyan). Scale bar=100 µm. (B) Voltage clamp recordings of the cells in A during illumination of oChIEF+ cells. Left and right panels show example traces from the corresponding left and right fan cell clusters in A, respectively. In the schematics, cyan and black colored cells represent oChIEF+ cells and oChIEF- fan cells, respectively. (C) Distribution of synaptic inputs onto fan cells after AAV injection in the DG/CA3. (D) Individual fan cells show inhibition or combined inhibition excitation, but not pure excitation, assessed following AAV injection in the DG/CA3. (E) Example average traces of outward (red) and inward (black) currents.
evoked from a single stimulation spot. Inset shows magnified view of the onset of postsynaptic currents, revealing a latency between outward and inward currents. (F) Histogram showing average latency between EPSC and IPSC measured from currents evoked at the same laser stimulation position. (G) Outward currents recorded following AAV injection into LEC are reversibly blocked by adding bicuculline, and partially restored upon wash out. Left panel shows voltage clamp traces. Average traces (red) are superimposed on the individual traces (grey). Right panel shows average current amplitudes before, during, and after bicuculline (n=11 cells from five animals). Individual data points are plotted together with population average amplitude (blue line). (H) Same as in G, but with the addition of DNQX/APV (n=10 cells from four animals) instead of bicuculline.

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<th>Fast-spiking cells n = 29</th>
<th>Non-fast spiking cells n = 19</th>
<th>Low threshold spiking cells n = 13</th>
<th>P-value Kruskal-Wallis H test</th>
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<td>AP half-width (ms)</td>
<td>0.75 (0.70-0.80)</td>
<td>1.10 (0.95-1.20)</td>
<td>1.15 (1.00-1.20)</td>
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<td>Max AP rate (Hz)</td>
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<td>54.00 (44.00-64.00)</td>
<td>66.00 (41.00-81.00)</td>
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<td>Sag ratio</td>
<td>0.99 (0.98-0.99)</td>
<td>0.98 (0.98-0.99)</td>
<td>0.91 (0.82-0.94)</td>
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<td>Rebound potential (mV)</td>
<td>1.73 (1.13-3.17)</td>
<td>1.31 (0.54-3.1)</td>
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<td>Input resistance (MO)</td>
<td>144.90 (120.29-181.97)</td>
<td>158.54 (126.97-199.22)</td>
<td>262.24 (178.32-281.77)</td>
<td>p = 0.0005 H(2) = 15.21</td>
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Table 1. Intrinsic electrophysiological properties of three distinct classes of LEC interneurons. Values are given as median and 25<sup>th</sup>-75<sup>th</sup> percentiles, and statistical differences between the groups are given as exact P-values from the Kruskal-wallis H test.