Postsynaptic GABABRs Inhibit L-Type Calcium Channels and Abolish Long-Term Potentiation in Hippocampal Somatostatin Interneurons

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**Cell Reports**

**Postsynaptic GABA<sub>B</sub>Rs Inhibit L-Type Calcium Channels and Abolish Long-Term Potentiation in Hippocampal Somatostatin Interneurons**

**Highlights**
- GABA<sub>B</sub> receptors do not activate Kir3-currents in CA1 somatostatin interneurons
- In somatostatin interneurons, GABA<sub>B</sub> receptors inhibit dendritic L-type Ca<sup>2+</sup> channels
- Ca<sub>V</sub>1.2 channels co-cluster with GABA<sub>B1</sub> on somatostatin interneuron dendrites
- GABA<sub>B</sub> activation abolishes long-term potentiation in somatostatin interneurons

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**In Brief**
Booker et al. show that GABA<sub>B</sub> receptors are highly expressed on somatostatin interneuron dendrites. Rather than activating Kir3 channels, they preferentially co-cluster with, and negatively couple to, L-type calcium channels inhibiting long-term potentiation at excitatory inputs.

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Postsynaptic GABA_B Rs Inhibit L-Type Calcium Channels and Abolish Long-Term Potentiation in Hippocampal Somatostatin Interneurons

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SUMMARY

Inhibition provided by local GABAergic interneurons (INs) activates ionotropic GABA_A and metabotropic GABA_B receptors (GABA_B Rs). Despite GABA_B Rs representing a major source of inhibition, little is known of their function in distinct IN subtypes. Here, we show that, while the archetypal dendritic-inhibitory somatostatin-expressing INs (SOM-INs) possess high levels of GABA_B R on their somatodendritic surface, they fail to produce significant postsynaptic inhibitory currents. Instead, GABA_B Rs selectively inhibit dendritic Ca_{V1.2} (L-type) Ca^{2+} channels on SOM-IN dendrites, leading to reduced calcium influx and loss of long-term potentiation at excitatory input synapses onto these INs. These data provide a mechanism by which GABA_B Rs can contribute to disinhibition and control the efficacy of extrinsic inputs to hippocampal networks.

INTRODUCTION

Maintained balance of excitation and inhibition controlled by feedforward and feedback interneurons (INs) is essential for appropriate function of cortical networks. Despite recruitment of local INs being critical to this balance, the contributing cellular mechanisms remain largely unexplored. Somatostatin (SOM) expressing INs constitute a dominant feedback inhibitory element in cortical circuits. In hippocampal CA1, SOM-INs are characterized by a somato-dendritic domain confined to stratum (str.) oriens and an axon providing inhibition to distal dendrites of pyramidal cells (PCs) in str. lacunosum-moleculare, as such they are referred to as O-LM cells (McBain et al., 1994; Katona et al., 1999; Müller and Remy, 2014). SOM-INs gate extrinsic cortical inputs into CA1 (Leão et al., 2012) and contribute to the generation of network oscillations at theta frequencies (Gloveil et al., 2005; Klausberger and Somogyi, 2008), with known roles in neuropathology (de Lanerolle et al., 1989; Dugladze et al., 2007; Wang et al., 2011). SOM-INs are recruited by recurrent input from CA1 PCs, involving ionotropic α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA) (Topolnik et al., 2005; Lamsa et al., 2007), N-methyl-D-aspartate (NMDA) (Standaert et al., 1996), and group 1 metabotropic glutamate receptors (mGluRs), particularly mGluR1α (Baude et al., 1993; McBain et al., 1994; Topolnik et al., 2006). Group I mGluRs on SOM-INs activate Ca_{V1.2} (L-type) high voltage-gated Ca^{2+} channels (VGCCs), promoting synaptic plasticity at excitatory inputs (Topolnik et al., 2009; Nicholson and Kullmann, 2014).

While glutamatergic mechanisms have been well characterized, inhibitory control of SOM-INs is less well understood (Tyan et al., 2014). In particular, little is known regarding the effects of metabotropic GABA_B Rs, despite GABA_B1 subunits being highly expressed at SOM-IN somata (Sloviter et al., 1999). In this study, we used whole-cell recording, 2-photon Ca^{2+}-imaging, and high-resolution quantitative SDS-digested freeze-fracture replica (SDS-FRL) immunoelectron microscopy to examine postsynaptic GABA_B R function and localization in SOM-INs.

RESULTS

To determine whether SOM-INs possess functional GABA_B Rs, we performed whole-cell recordings from rat acute hippocampal slices. CA1 SOM-INs were located in str. oriens/alic买到 us with horizontal dendrites (Figure 1A) and responded with a large voltage "sag" to hyperpolarizing currents and minimally adapting action potential (AP) trains to depolarizing currents (Figure 1A, inset). All INs tested were immunoreactive for SOM (155 INs), of which 64 (41.3%) were identified as O-LM cells and 3 (1.9%) were bistratified INs. The remaining 88 (56.8%) were not morphologically identified due to the axon being cut, but were included in further
CA1 PCs were recorded as controls, given their well described functional GABA<sub>B</sub>R expression (Lüscher et al., 1997).

**GABA<sub>B</sub>R-Mediated IPSCs Are Small in SOM-INs**

Hippocampal neurons possess slow GABA<sub>B</sub>R-mediated inhibitory postsynaptic currents (sIPSCs), elicited by extracellular stimulation (Degro et al., 2015). In SOM-INs, trains of stimuli (5 at 200 Hz) to str. oriens in the presence of GABA<sub>A</sub>, NMDA, and AMPA receptor blockers produced very small or no sIPSC (Figures 1B, upper, and 1C). The mean sIPSC amplitude was 1.8 ± 0.5 pA (17 cells) and when present was blocked by the GABAB<sub>R</sub> antagonist CGP-55,845 (CGP, 10 cells). In CA1 PCs, sIPSCs were markedly larger with a mean amplitude of 28.3 ± 8.8 pA (6 cells, U<sub>17,6</sub> = 0; p < 0.0001, Mann-Whitney test) (Figures 1B and 1C). In summary, the GABA<sub>B</sub>R-mediated inhibitory conductance in SOM-INs is an order of magnitude lower than CA1 PCs, indicating that GABA<sub>B</sub>R/Kir3 signaling does not significantly contribute to SOM-IN inhibition.

**GABA<sub>B</sub>Rs Strongly Inhibit Dendritic L-Type VGCCs**

The absence of GABA<sub>B</sub>R-mediated currents in SOM-INs suggests that the receptors may signal through an alternative effector, such as high voltage-gated CaV1.2 (L-type) Ca<sup>2+</sup> channels (Chalifoux and Carter, 2011) known to contribute to signaling and plasticity in SOM-INs (Topolnik et al., 2006). To determine whether GABA<sub>B</sub>Rs inhibit CaV1.2 in SOM-INs, we performed 2-photon imaging of IN dendrites filled with a morphometric and a Ca<sup>2+</sup>-indicator dye and evoked short trains of back-propagating APs (bAPs, 4 at 200 Hz) (Figure 2A). Imaging a primary dendrite with rapid line-scans, we observed large Ca<sup>2+</sup>-transients in response to bAPs (Figures 2B and 2C), which had a ΔF/F of 31.5% ± 3.7% (17 cells). These transients were stable for the 20-min recording (4 cells) (Figure S3A) and blocked by CdCl<sub>2</sub> (5 mM, 3 cells) (Figure S3B) and by a marked reduction in input resistance from 130 ± 14 MΩ to 95 ± 11 MΩ in CA1 PCs (10 cells) consistent with channel opening. In contrast, only a small change from 207 ± 21 MΩ to 198 ± 25 MΩ was observed in SOM-INs (14 cells; U<sub>14,10</sub> = 4.0; p < 0.0001, Mann-Whitney test) (Figure 1F). This finding was further validated by briefly activating GABA<sub>B</sub>Rs through uncaging of GABA over the dendrites of SOM-INs and CA1 PCs (Figure S2A), which resulted in currents of 6.4 ± 2.6 pA (5 cells) and 93.7 ± 23.4 pA (6 cells), respectively (U<sub>5,6</sub> = 0; p = 0.0043, Mann-Whitney test) (Figures S2B and S2C). In summary, the GABA<sub>B</sub>R-mediated inhibitory conductance in SOM-INs is an order of magnitude lower than CA1 PCs, indicating that GABA<sub>B</sub>R/Kir3 signaling does not significantly contribute to SOM-IN inhibition.
(10 cells) resulted in a 27.8% ± 6.5% reduction in the Ca2+ response (t(6) = 4.25; p = 0.002, Wilcoxon test), which recovered in CGP (5 μM; t(4) = 0.08; p = 0.94, Wilcoxon test) (Figures 2B and 2C).

To confirm that L-type VGCCs contribute to Ca2+-transients, we applied the selective blocker nifedipine (10 μM), resulting in a 13.2% ± 4.2% reduction in the signal (t(4) = 3.20; p = 0.033, Wilcoxon test) (Figure 2D), comparable to baclofen effect (t(10,5) = 1.35; p = 0.22, one-way ANOVA). Moreover, co-application of baclofen and nifedipine did not further reduce the Ca2+ signal (t(10,9) = 1.02; p = 0.68) (Figures 2D and 2E), independent of whether the co-application followed an initial baclofen (t(4,4) = 1.05; p = 0.62, Holm-Sidak’s post-test) or nifedipine application (t(6,6) = 1.39; p = 0.22, Holm-Sidak’s post-test). This mutual occlusion indicates that GABABRs predominantly inhibit L-type VGCCs in SOM-IN dendrites. This result was verified using 1-photon imaging, using the same pharmacological treatment (Figure S4). Furthermore, GABABR inhibition of Ca2+-transients was maintained into adulthood (Figures 2F–2H).

Figure 2. GABABRs Inhibit L-Type VGCC-Mediated Ca2+-Transients in SOM-IN Dendrites
(A) A 2-photon image of a SOM-IN filled with Alexa Fluor 594 (morphometric dye) and BAPTA-OG1 (100 μM for Ca2+-imaging) and the imaged proximal dendritic segment (inset). Top right: confocal images confirm immunolabeling for SOM (green, left) at the soma (blue, right). Bottom right: voltage response of the cell to hyper- to depolarizing current pulses. Scale bars, 5 μm (left inset); 10 μm (right insets). (B) Trains of bAPs (top) evoked by brief pulses (1 nA, 1 ms, 200 Hz) in the SOM-IN resulted in Ca2+-transients (bottom) under control conditions (left, gray), in baclofen (10 μM, middle, in black) and CGP (5 μM, right, in black).

(C) Summary of peak Ca2+-transients during control (Ctrl), baclofen (Bac), and CGP in 5 SOM-INs from juvenile rats (P17–P25).

(D) Ca2+-transient amplitudes during control condition, application of nifedipine (Nif) and co-application of nifedipine and baclofen (Bac+Nif) from 5 SOM-INs.

(E) Bar chart of normalized Ca2+-transients during application of Bac, Nif, Nif+Bac, and CGP in juvenile rats. Bar charts show mean ± SEM. Data from individual cells is superimposed on bars (open circles); numbers of tested cells are in parentheses. ns, p > 0.05; *p < 0.05; **p < 0.01. See also Figures S3 and S4.

(G) Summary of peak Ca2+-transients during control (Ctrl), in baclofen (Bac) and CGP in 9 SOM-INs from adult rats (P50–P60).

(H) Bar chart of normalized Ca2+-transient amplitudes in Bac and CGP in juvenile rats. Bar charts show mean ± SEM. Data from individual cells is superimposed on bars (open circles); numbers of tested cells are in parentheses. ns, p > 0.05; *p < 0.05; **p < 0.01. See also Figures S3 and S4.

(G) GABAARs and CaV1.2 Channels Preferentially Cluster on Dendritic Shafts of mGluR1α-Expressing Cells
Our data indicate that GABAARs modulate L-type VGCCs but not Kir3 channels in SOM-INs. Therefore, we next examined the distribution, density, and spatial relationship of GABAARs, Kir3, and CaV1.2 channel on SOM-IN dendrites by quantitative SDS-FRL electron microscopy, using mGluR1α as a surface marker for SOM-INs (Baude et al., 1993). Immunoreactivity for the GABAAR α1 subunit was consistently observed at mGluR1α-positive dendrites (Figure 3A), with a density of 49.1 ± 4.5 particles/μm²
(35 dendrites from 3 animals), higher than that of CA1 PCs in the same replicas (28.5 ± 3.2 particles/μm², 36 dendrites, U(34,35) = 287.0; p < 0.0001, Mann-Whitney test) (Figure 3B). In contrast, Kir3 channel subunit density was 8.6 ± 1.1 particles/μm² on mGluR1α dendrites (39 dendrites from 3 animals) (Figure 3C), 50% lower than on neighboring PCs (16.2 ± 1.6 particles/μm², 45 dendrites, U(38,44) = 459.0; p = 0.0001, Mann-Whitney test) (Figure 3D), explaining the small GABA B1-mediated currents in SOM-INs.

Next, we determined the surface expression of Ca V1.2, which was observed on mGluR1α-positive dendrites (Figure 3E) with a density of 14.3 ± 1.5 particles/μm² (37 dendrites from 3 animals) (Figure 3F), over 3-fold higher than on PCs (4.3 ± 0.4 particles/μm², 30 dendrites, U(36,29) = 77.0; p < 0.0001). Finally, to examine the spatial relationship between GABA B1 and CaV1.2 we performed triple labeling for mGluR1α, GABA B1 and Ca V1.2 (Figure 3G) and measured the proximity of Ca V1.2 particles to the closest GABA B1 particle. This analysis revealed that 51% of Ca V1.2 subunit-containing channels were located within 100 nm of a GABA B1 subunit (Figure 3H). Thus, GABA B1 and Ca V1.2 subunits are present at high density and colocalize on SOM-IN dendrites.

**Postsynaptic GABA B1 Activation Inhibits Synaptic Plasticity in SOM-INs**

Long-term potentiation (LTP) at excitatory synapses onto SOM-INs critically depends on L-type VGCC activation (Topolnik et al., 2009). Therefore, we asked if associative LTP in SOM-INs is sensitive to GABA B1 activation. EPSC amplitudes were potentiated to 163.8% ± 17.3% (measured at 20–25 min; t(6) = 3.99; p = 0.007, t test, 7 cells) (Figure 4A) following LTP induction in SOM-INs. When baclofen was pre-applied, the same stimulus did not potentiate EPSCs (mean EPSC amplitude: 95.8% ± 10.8% of baseline; t(5) = 0.47; p = 0.66, t test, Wilcoxon test, 6 cells) (Figures 4B and 4F). A comparable GABA B1-mediated inhibition of LTP was observed in adult rats (Figures 4C, 4D, and 4F). To confirm that the LTP observed was dependent on L-type VGCCs, as previously shown (Topolnik et al., 2009), we pre-applied nifedipine (10 μM) to 6 SOM-INs. As expected, this manipulation fully abolished LTP (EPSC: 89.2 ± 11.0% of baseline; t(5) = 1.07; p = 0.33, t test) (Figures 4E and 4F). These data,
thus, demonstrate that activation of postsynaptic GABA\(_B\)Rs, via inhibition of Ca\(_V1.2\) Ca\(^{2+}\) channels, abolishes postsynaptic LTP induction at excitatory synapses onto SOM-INs.

**DISCUSSION**

In the present study, we provide compelling evidence that GABA\(_B\)Rs are present on dendritic membranes of CA1 SOM-INs, but do not activate the canonical Kir3 signaling cascade. Rather, GABA\(_B\)Rs cluster with and inhibit L-type VGCCs, reducing dendritic calcium influx and blocking LTP at excitatory synapses onto SOM-INs. This effect will preclude synaptic strengthening during network activation, a mechanism by which GABA\(_B\)Rs can contribute to a long-term alteration of excitation and inhibition balance in the network.

**Small GABA\(_B\)R-Mediated Inhibitory Currents in SOM-INs**

The major GABA\(_B\)R signaling in postsynaptic compartments has long been considered to involve Kir3 channels (Lüscher et al., 1997; Kaupmann et al., 1998; Degro et al., 2015). In contrast, while we find that GABA\(_B\)Rs and Kir3 channels are present on SOM-IN dendrites, only very small K\(^+\) currents were produced, partially explained by a lower Kir3 channel expression. In a network context, the small GABA\(_B\)R currents in SOM-INs are consistent with observations in other dendritic-targeting IN types: parvalbumin bistriatified cells (Booker et al., 2013) and cholecystokinin INs (Booker et al., 2017) and may be a common principle for dendritic-targeting INs. This divergence between perisomatic and dendritic inhibitory INs implies that GABA\(_B\)R activation shifts inhibition between the two target compartments.

**Colocalization and Negative Coupling of GABA\(_B\)Rs and VGCCs in SOM-INs**

We provide evidence that postsynaptic GABA\(_B\)Rs preferentially signal through and inhibit VGCCs in SOM-INs. The high
expression of CaV1.2 and nifedipine-sensitive Ca²⁺-transients further indicate that these channels substantially contribute to Ca²⁺ influx in SOM-IN dendrites. In fact, GABA_BRs have been shown to inhibit VGCCs, as an alternative postsynaptic effector in PC dendrites (Sabatini and Svoboda, 2000; Chalifoux and Carter, 2011; Pérez-García et al., 2013). The co-clustering of the GABA_Bs subunits with the CaV1.2 (L-type) VGCCs subunit is a structural correlate of this interaction in SOM-IN dendrites and may reflect a tight functional coupling through a membrane delimited G_Vo-TY interaction (Pérez-García et al., 2013). Whether these mechanisms also apply to dendritic spines remains an open question.

**GABA_BR Signaling Abolishes Synaptic Plasticity in SOM-INs**

By negatively coupling to L-type VGCCs, GABA_BRs block the induction of LTP in SOM-INs, adding to the wide repertoire of molecular mechanisms involved in synaptic plasticity in these INs (Topolnik et al., 2006; Nicholson and Kullmann, 2014; Vasuta et al., 2015). The form of plasticity is dependent on the activity pattern (Lamsa et al., 2007) that is plausibly translated into a differential activation of glutamate and downstream signaling cascades (Topolnik et al., 2005, 2006, 2009; Oren et al., 2009). Indeed, L-type VGCCs potentiation by group I mGluRs promotes LTP in SOM-INs (Topolnik et al., 2009). In cerebellar Purkinje cells, GABA_BRs facilitate mGluR1 activation (Hirono et al., 2001). In SOM-INs, the two receptors converge on L-type VGCCs, but exert opposing actions: GABA_BRs intercept mGluR-mediated signaling by inhibiting L-type channels and thereby abolish LTP induction.

Inhibition of LTP in SOM-INs by GABA_BRs is in stark contrast to the facilitation of LTP by GABA_BR activation observed in PCs (Davies et al., 1991; Mott and Lewis, 1991). In fact, despite their inhibitory nature, GABA_BRs can produce disinhibitory effects in cortical networks due to a preferential inhibitory bias of IFS and their output synapses (Foster et al., 2013; Papatheodoropoulos, 2015). In SOM-INs, GABA_BRs do not produce hyperpolarization, but prevent the induction of LTP and thereby preclude an enhanced recruitment of the feedback circuit. Considering that the main output of SOM-INs is onto PC distal dendrites, this reduced recruitment will allow increased synaptic transmission onto CA1 PCs and may lead to a breakdown of the specificity of spatial information carried by entorhinal inputs onto CA1 PCs (Leão et al., 2012) via activation of GABA_BRs on nearby pre- and postsynaptic elements, as previously described in the neocortex (Urban-Ciecko et al., 2015). Indeed, prior studies have shown that GABA_BR activation is capable of impairing hippocampal-dependent spatial learning (McNamara and Skelton, 1996; Arofo et al., 1998), consistent with the importance of this circuit.

**EXPERIMENTAL PROCEDURES**

**Electrophysiological Recordings**

A full description of methods can be found in the Supplemental Information. In brief, 300–400 μm acute hippocampal slices were prepared from juvenile (17- to 25-day-old) and adult (50- to 60-day-old) male Wistar rats (Booker et al., 2014, 2017). All experiments were performed in accordance with institutional, local governmental (LaGeSo, Berlin T 0215/11; LaGeSo, Freiburg X-14/11H) and national guidelines (German Animal Welfare Act; ASPA, United Kingdom Home Office). Whole-cell recordings were made using pipettes filled with K-glucuronate-based solution at 32 °C ± 1 °C. GABA_B-mediated currents were measured in the presence of the ionotropic receptor blockers NBQX, CNQX or DNPX, DL-APV, and bicuculline, gabazine, or picrotoxin at a holding potential of −65 mV. Synaptic currents were elicited by a glass monopolar electrode in str. oriens.

For Ca²⁺-imaging, we used 2-photon microscopy with pipettes filled with intracellular solution containing BAPTA-OGB1 and a morphometric dye. Ca²⁺-transients were measured in proximal dendrites following trains of 4 APs to evoke Ca²⁺ influx, line-scans were recorded at ~200 Hz. Baclofen or nifedipine was applied to the bath; CGP was applied following baclofen to confirm receptor specificity.

LTP was induced at inputs to SOM-INs with EPSCs elicited by a monopolar electrode placed in the alveus. Theta-burst stimulation was paired with a postsynaptic depolarization to −20 mV, repeated 3 times at 30-s intervals. In a subset of experiments, baclofen or nifedipine were pre-applied to the bath, and the EPSC was titrated to match control recordings. All neurons were filled with biocytin during recordings, fixed overnight, labeled with streptavidin and antibodies to SOM, and imaged with confocal microscopy.

**Electron Microscopy**

Electron microscopic analysis was performed on 60-day-old wild-type Wistar rats (Althof et al., 2015). Coronal hippocampal sections were cut, cryoprotected, and blocks of str. oriens/str. alveus of CA1 were dissected and frozen under high-pressure. Samples were fractured and coated with carbon and platinum in a freeze-fracture replica machine. Replicas were then digested, washed, blocked, and then incubated with subunit-specific primary antibodies followed by incubation with gold-coupled secondary antibodies. Strongly mGluR1s-immunoreactive and CA1 PC dendrites in str. oriens were imaged and analyzed.

**Statistics**

All data are shown as mean ± SEM. Analysis was performed in GraphPad Prism 3.0 (GraphPad Software, CAUSA). For all electrophysiology data, “n” refers to the number of recorded cells; for electron microscopy “n” refers to the number of dendrites tested from 3–4 rats. Group data were compared with one-way ANOVA test combined with Holm’s-Sidak post-test. Analysis of unpaired and paired data was performed with Mann-Whitney or Wilcoxon matched-pairs tests, respectively. Significance was assumed if p < 0.05.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Supplemental Experimental Procedures and four figures and can be found with this article online at https://doi.org/10.1016/j.celrep.2017.12.021.

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**AUTHOR CONTRIBUTIONS**


**DECLARATION OF INTERESTS**

The authors declare no competing interests.
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