

PP2A-dependent internalisation of GABA_B receptors in somatostatin interneurons regulates function and plasticity.

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Figures: 8 (4 supplementary)

Tables: 0 (2 supplementary)

1 **Abstract:**

2 Cortical circuits rely on a precise balance of inhibitory and excitatory neurotransmission to
3 encode information reliably and prevent pathology. Metabotropic GABA_B receptors
4 (GABA_BRs) are key regulators of inhibitory signalling in mammalian neurons. In
5 GABAergic interneurons (INs), GABA_BR activation reduces inhibition overall, leading to
6 disinhibitory mechanisms. In the hippocampus, somatostatin-expressing (SST) INs form a
7 major subtype that provides feedback inhibition to the distal dendrites of principal cells
8 (PCs) and other INs. Plasticity of SST INs is well established as a mechanism controlling
9 hippocampal circuit function through both inhibitory and disinhibitory pathways and
10 depends on metabotropic glutamate receptors (mGluRs) and GABA_BRs. However,
11 whether activation of GABA_BRs induces metaplastic changes in SST INs, and how this
12 influences circuit function and behaviour, remains unclear.

13 Here, we combined quantitative SDS-digested freeze-fracture replica immunoelectron
14 microscopy, ex vivo electrophysiology, in vivo behavioural testing, and pharmacological
15 manipulation of GABA_BRs. We show that receptor activation directly regulates SST IN
16 plasticity via protein phosphatase 2A (PP2A)-dependent internalisation. GABA_BR
17 activation not only controls its own surface expression but also regulates membrane levels
18 of mGluR1 α and high-voltage-activated Ca_v1.2 (L-type) Ca²⁺ channels. This GABA_BR-
19 dependent metaplasticity shifts circuit plasticity toward greater enhancement of long-range
20 inputs to the CA1 region and disrupts contextual memory formation. These findings
21 demonstrate that receptor-mediated surface dynamics in SST INs are critical for
22 maintaining physiological neurotransmission and proper hippocampal microcircuit function.

23

24 **Introduction**

25 The appropriate function of cortical microcircuits relies on dynamically balanced excitatory
26 and inhibitory synaptic transmission, which maintains the neural code and contributes to
27 memory formation¹⁻³. How these circuits respond to changing activity level, or indeed
28 sustained pharmacological activity, is crucial to understand how the brain functions at rest
29 and how neuropathological conditions lead to altered behaviour and cognition.

30 Synaptic inhibition is produced, in large part, by a heterogenous population of local
31 interneurons (INs). These INs release GABA from their presynaptic terminals, which
32 activates both synaptic and extrasynaptic ionotropic GABA_A receptors (GABA_ARs) and

33 metabotropic GABA_B receptors (GABA_BRs), on excitatory principal cells and INs alike ⁴.
34 Such inhibition of INs reduces their activity, leading to reduced local GABA release, and
35 thus a reduction of inhibitory control – known as disinhibition. Dynamic interactions
36 between inhibition and disinhibition are critical to ensure the timing and strength of
37 neuronal activity leading to complex behaviour. To date, most studied forms of disinhibition
38 focus on direct synaptic GABA_AR-mediated effects, which can regulate circuit level
39 plasticity in hippocampus ⁵ and neocortex ⁶. While the direct effect of GABA_BR signalling
40 on INs has been well described by ourselves and others ⁷⁻¹⁴, how their activation shapes
41 the long-term excitability of inhibitory cells and influence circuit activity remains less well
42 understood.

43 In CA1 of the hippocampus, a major type of IN are those which express the neuropeptide
44 somatostatin (SST)¹⁵ which have their somatodendritic domain within *stratum*
45 *oriens/alveus* (*str. O/A*) receiving inputs from local pyramidal cells (PCs)^{10,16,17}, with axons
46 ramifying in *str. lacunosum-moleculare* (*Str. L-M*). Thus, SST INs provide powerful
47 feedback inhibition into the CA1 circuit¹⁸. Therefore, the activation of SST INs leads to
48 complex circuit functions. First, they directly inhibit distal dendrites of CA1 PCs co-
49 terminating with entorhinal cortex inputs to strongly inhibit spatial information¹⁹⁻²¹. Second,
50 SST INs disinhibit proximal dendritic domains and thus gate contextual inputs from CA3²².
51 As such, long-term potentiation of SST INs leads to a functional re-arrangement of the
52 hippocampal information transfer, favouring CA3 inputs ^{22,23} and are potentially able to
53 alter the circuit dynamics of the hippocampus over behaviourally relevant time-scales ²⁴.
54 We have shown that GABA_BRs on SST INs acutely inhibit the induction of long-term
55 potentiation (LTP)¹¹ and their local circuit integration ¹⁰, but do not directly hyperpolarise
56 SST INs. Given the high affinity of GABA for GABA_BRs and their plethora of accessory
57 proteins, they undergo rapid desensitization ^{8,25-29} and agonist dependent internalisation
58 ^{30,31}. Not only this, but GABA_BRs are known to interact with metabotropic glutamate
59 receptors (mGluRs), preferentially with type 1α receptors (mGluR1α), in other GABAergic
60 cells in the cerebellum ^{32,33}. Considering the high density of both mGluR1α, and
61 GABA_BRs on SST INs ^{34,35} it is plausible that receptor cross-talk also exists in these cells.
62 Whether GABA_BR signalling is susceptible to rapid internalisation in SST INs, and how this
63 affects cell function and the local circuit remains unknown.

64 We hypothesised that GABA_BR activation leads to internalisation of itself, favouring greater
65 plasticity in SST INs. To determine this, we performed quantitative SDS-digested freeze-
66 fracture replica immunogold labelling (SDS-FRL) electron microscopy, *ex vivo*

67 electrophysiological recordings from CA1 of the mouse hippocampus following prolonged
68 application of the selective GABA_BR agonist baclofen, and behavioural characterisation.
69 We found, in opposition to our hypothesis, that after sustained GABA_BR activation,
70 synaptic plasticity of SST INs was reduced, independent of direct receptor activation. This
71 reduction in plasticity was due to concomitant protein phosphatase 2A (PP2A)-mediated
72 internalisation of GABA_BRs, Ca_v1.2 (L-type) voltage-gated Ca²⁺ channels (VGCCs) and
73 mGluR1α – both of which are required for LTP of SST INs. Finally, we show that baclofen
74 administration leads to circuit level strengthening of long-range inputs to CA1, which are
75 dependent on SST IN plasticity, and an inability to form contextual memories.

76

77

78 Results

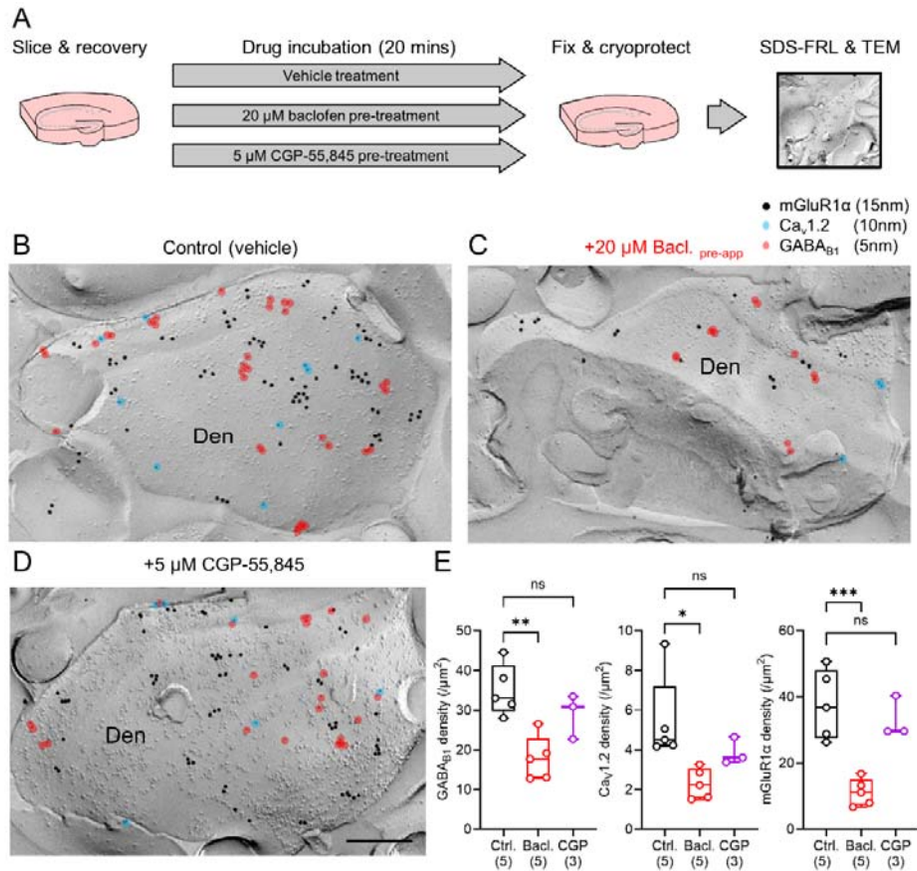
79 In this study we tested the hypothesis that sustained baclofen application leads to
80 internalisation of GABA_BRs, which favours enhanced plasticity of SST INs following
81 receptor activation. For this, we performed detailed quantitative immunoelectron
82 microscopy, electrophysiology, and behaviour in adult mice.

83 *Native GABA_BRs, mGluR1 α and L-type calcium channels are down-regulated in SST IN*
84 *membranes following baclofen pre-application, which requires PP2A*

85 We have previously shown that mouse SST IN dendrites possess a high density of
86 GABA_{B1}, Ca_v1.2, and mGluR1 α ¹¹. The latter two proteins are required for associative
87 theta-burst stimulus (aTBS) LTP of SST INs^{36,37}, and is inhibited by GABA_BRs¹¹. To
88 determine whether pre-application of the GABA_BR agonist baclofen (20 μ M) altered
89 membrane localization of GABA_BRs as compared to mGluR1 α and Ca_v1.2, we performed
90 SDS-FRL immunoelectron microscopy analysis from acute mouse hippocampal slices pre-
91 treated with baclofen (n=5 mice), compared slices treated with vehicle (n=5 mice) or the
92 GABA_BR antagonist CGP-55,845 (5 μ M, n=3 mice; Figure 1A). All SDS-FRL data are
93 shown as animal average values, with statistics shown based on linear mixed-effects
94 modelling to account for inter-animal variability. Data from each dendritic profile within
95 animal is shown in Figure S1.

96 Under control conditions (Figure 1B), putative SST IN dendrites were identified based on
97 high levels of surface labelling for mGluR1 α ³⁴, with an average density of 37.6 ± 4.7
98 particles/ μ m² (Figure 1B, E). Consistent with our previous data in rats¹¹, mGluR1 α -positive
99 dendrites displayed high-density GABA_{B1} subunit labelling of 35.1 ± 2.9 particles/ μ m² and
100 moderate expression of the L-type channel subunit Ca_v1.2 at 5.46 ± 0.98 particles/ μ m²
101 (Figure 1B, E). Compared to control slices, baclofen pre-application led to a 49% reduction
102 in density to 17.9 ± 2.5 particles/ μ m² for GABA_{B1} ($t_{(5,5)}=5.07$, $p=0.004$, Tukey test; Figure
103 1C, E). We observed a 58% reduction in Ca_v1.2 to 2.3 ± 0.3 particles/ μ m² ($t_{(5,5)}=5.36$,
104 $p=0.0009$, Tukey test) and a 70% reduction of mGluR1 α labelling 11.2 ± 1.8 particles/ μ m²
105 ($t_{(5,5)}=6.36$, $p=0.0004$, Tukey test) (Figure 1C, E). Application of the GABA_BR antagonist
106 CGP-55,845 did not decrease density of GABA_{B1} ($t_{(5,3)}=1.97$, $p=0.18$, Tukey test), Ca_v1.2
107 ($t_{(5,3)}=2.01$, $p=0.17$, Tukey test) or mGluR1 α ($t_{(5,3)}=0.69$, $p=0.78$, Tukey test) relative to
108 control levels (Figure 1D, E). This effect on effector channels appeared confined to SST
109 INs, as measurement of CA1 PC dendrites in the same replicas revealed that while
110 baclofen pre-application reduced GABA_{B1} density by 49.9% ($F_{(5,5)}=45.22$, $p=4.57 \times 10^{-10}$,

111 LMM, Supplementary Figure 2) $Ca_v1.2$ was unchanged ($F_{(3,3)}=0.009$, $p=0.93$, LMM;
 112 Supplementary Figure 2). $mGluR1\alpha$ was not tested as this protein is not expressed
 113 abundantly on CA1 PCs. Together, these data reveal that $GABA_B$ R activation leads to its



114 own reduction from the membrane of SST INs, but also leads to down-regulation of $Ca_v1.2$
 115 and $mGluR1\alpha$.

116 *Figure 1 Sustained $GABA_B$ R activation leads to reduction of membrane associated*
 117 *$GABA_{B1}$, $Ca_v1.2$, and $mGluR1\alpha$ proteins from putative SST-INs. A* Schematic of the
 118 experimental overview, indicating treatment and experimental sample times. **B** Example
 119 electron micrographs of SDS-FRL for $mGluR1\alpha$ (15 nm immunoparticles), $Ca_v1.2$ (10 nm,
 120 blue overlay) and $GABA_{B1}$ (5 nm, red overlay) on a putative SST IN dendrite in *str. O/A* of
 121 CA1 in a vehicle slice. The same labelling but performed in brain slices following 20
 122 minutes baclofen pre-application (20 μ M, **C**) or CGP-55,845 (5 μ M, **D**). **E** Quantification of
 123 labelling density for $GABA_{B1}$ (left), $Ca_v1.2$ (middle) and $mGluR1\alpha$ (right), in slices treated
 124 with vehicle (Ctrl., black, 5 mice), baclofen pre-application (Bacl., red, 5 mice), or CGP-
 125 55,845 (CGP, magenta, 3 mice). All data are shown as boxplots depicting the 25-75%
 126 range with median, maximum range and data from individual mice overlaid as open

127 circles. Statistics shown as: ns – $p > 0.05$, * - $p < 0.05$, ** - $p < 0.01$, *** - $p < 0.001$; all from
128 post hoc Tukey tests following LMM analysis. Scale bar represents 200 nm.

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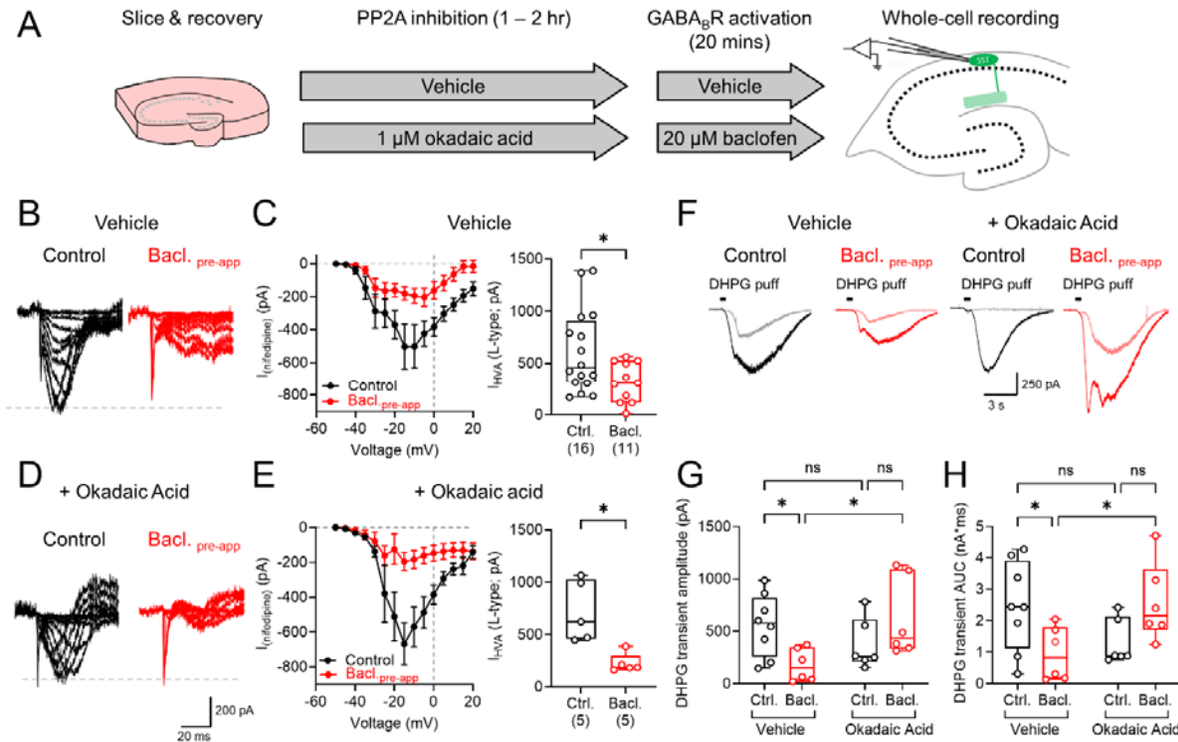
130 *Baclofen pre-application leads to reduced Cav1.2 and group 1 mGluR-mediated currents*
131 *in SST INs.*

132 To determine whether baclofen pre-application regulates the function of Cav1.2 and
133 mGluR1 α in SST INs, we next performed whole-cell patch clamp recordings from identified
134 cells using targeted pharmacology to interrogate receptor internalisation (Figure 2A), as
135 GABA_BR internalisation has been shown to be dependent on increased activity of
136 PP2A^{26,31,38}.

137 As Cav1.2 containing VGCCs are a major target of GABA_BR mediated inhibition in SST
138 INs¹¹, we first performed whole-cell recordings using a Cs-gluconate based internal
139 solution containing QX-314 (5 mM) to block voltage activated K⁺ and Na⁺ channels,
140 respectively, in the presence of CNQX (10 μ M), DL-AP5 (50 μ M) and gabazine (10 μ M) to
141 block ionotropic receptor signalling. High-voltage activated Ca²⁺ responses were
142 measured from -60 mV voltage-clamp, in which 5 mV depolarising current steps were
143 applied, followed by bath application of the selective L-type VGCC blocker nifedipine
144 (10 μ M; Figure 2B). The average subtracted L-type current measured across a range of
145 depolarising stimuli revealed large amplitude currents in SST INs recorded under control
146 conditions, which were substantially reduced, but with a similar voltage-dependency in
147 slices pre-application with baclofen. We found that baclofen pre-application reduced the
148 peak L-type current measured by 37% ($U_{(16,11)} = 50$; $P = 0.032$, Mann-Whitney test, Figure
149 2C). We performed the same recordings, but in slices that had been incubated with the
150 PP2A inhibitor okadaic acid (OA, 1 μ M; Figure 2D). Blocking PP2A activity had minimal
151 effect on current-voltage response or peak L-type currents following baclofen pre-
152 application ($U_{(5,5)} = 0$; $P = 0.008$, Mann-Whitney; Figure 2E).

153 For mGluR1 signalling, we recorded SST INs in whole-cell configuration then puff applied
154 the selective, high-affinity group I mGluR agonist S-DHPG (50 μ M, 200 ms, 10 PSI) within
155 100-200 μ m of the cell body. S-DHPG puffs resulted in large and slow depolarising
156 currents in SST INs, which were partially sensitive to the mGluR1 α selective antagonist
157 LY367,385 (100 μ M). Measurement of the same S-DHPG induced currents in slices that
158 were pre-application with baclofen possessed typically smaller currents (Figure 2F).
159 Quantification of peak S-DHPG currents revealed that baclofen pre-application reduced

160 the amplitude ($F(1,22)=7.99$, $P=0.01$, 2-way ANOVA [interaction], Figure 2G) and the
 161 charge transfer ($F(1,22)=9.59$, $P=0.005$, 2-way ANOVA [interaction], Figure 2H), effects
 162 which were not present when slices were incubated with OA prior to baclofen pre-
 163 application.



164 *Figure 2: Ca_v1.2 and group 1 mGluR-mediated currents are reduced following baclofen*
 165 *pre-application.* **A** Schematic of recording conditions, highlighting treatment periods and
 166 recording times. **B** Example nifedipine-sensitive VGCC current responses recorded in SST
 167 INs following depolarising stimulation from -60 mV. Dashed line indicates control response
 168 amplitude for comparison. **C** Quantification of voltage-dependency of nifedipine-sensitive
 169 VGCC currents (left) and peak nifedipine-sensitive currents (right). **D** Example nifedipine-
 170 subtracted isolated VGCC current responses in SST INs pre-application with 1 μM okadaic
 171 acid. **E** Quantification of voltage-dependency of nifedipine-sensitive VGCC currents (left)
 172 and peak nifedipine-sensitive currents (right) in okadaic acid pre-application SST INs. **F**
 173 Example voltage responses following 50 μM S-DHPG puff application to SST INs,
 174 recorded under control conditions (black) or following baclofen pre-application (red),
 175 following either vehicle or 1 μM okadaic acid pre-application. **G** Quantification of absolute
 176 current amplitudes following S-DHPG puff. **H** Quantification of the area-under-curve (AUC)
 177 for S-DHPG puff induced transients. Data are shown as either mean ± SEM (current-
 178 voltage plots) or box-plots showing 25-75% box with median, with maximum range shown.

179 Individual cell data is shown overlaid. Statistics shown as: ns – $p > 0.05$, * - $p < 0.05$; from 2-
180 way ANOVA and Mann Whitney non-parametric tests

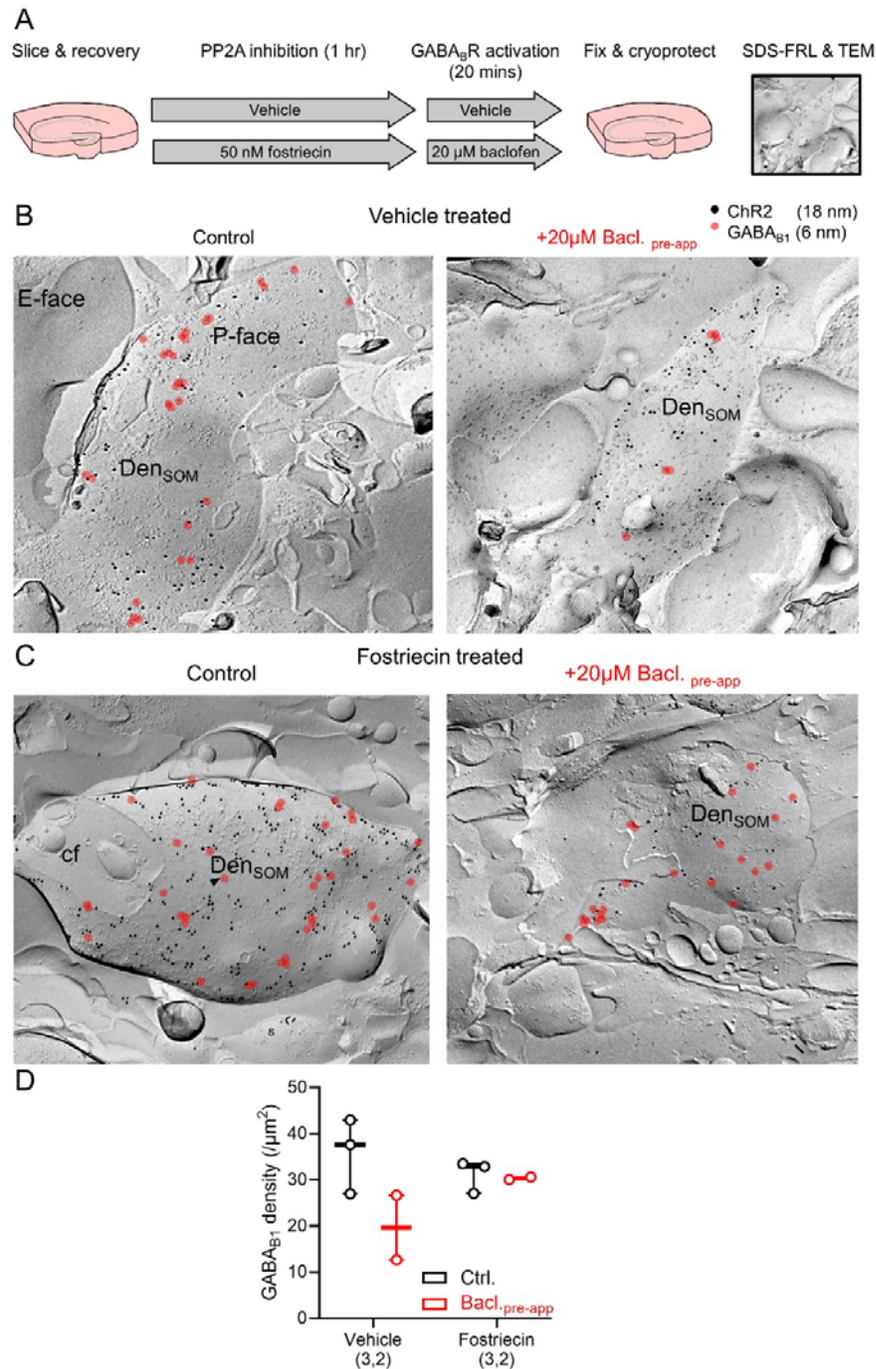
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183 *Baclofen pre-application dependent reductions in $Ca_v1.2$ and $mGluR1\alpha$ are mediated by*
184 *PP2A.*

185 To confirm surface down-regulation of $GABA_B$ Rs was indeed dependent on PP2A in SST
186 INs, we utilised a mouse line expressing a Channelrhodopsin2 fused to yellow
187 fluorescence protein (ChR2/YFP) under the SST promotor to allow selective SDS-FRL
188 from identified SST IN dendrites¹⁰. Initially, we prepared slices from WT mice treated with
189 the PP2A inhibitor OA. However, in these slices we observed minimal to no labelling for
190 $mGluR1\alpha$, $GABA_{B1}$ or $Ca_v1.2$ labelling in putative SST IN and CA1 PC dendritic profiles
191 (Figure S3). This effect was observed across >5 independent experiments.

192 As such, we next used the the small molecule PP2A inhibitor fostriecin (50 nM) which
193 slices were incubated with for 1 hour prior to baclofen pre-application (Figure 3A). In *str.*
194 *O/A* from the ChR2/YFP mice, we observed strong immunolabeling for YFP (33.7 ± 1.07
195 particles/ μm^2 , Figure 3B). In vehicle slices, we observed a 45% reduction of $GABA_{B1}$
196 labelling following baclofen pre-treatment (16.7 ± 1.02 particles/ μm^2 , $t_{(3,2)}=3.45$, $p=0.037$,
197 Kenward-Roger test, Figure 3B, D), consistent with observations from $mGluR1\alpha$ labelled
198 dendrites (Figure 1). Incubation with fostriecin prevented the baclofen pre-treatment
199 mediated reduction in $GABA_{B1}$ surface density (28.3 ± 3.2 particles/ μm^2 $t_{(3,2)}=0.88$, $p=0.44$,
200 Kenward-Roger test, Figure 3C, D), confirming that PP2A inhibition prevents $GABA_B$ R
201 internalisation.



202 *Figure 3: Inhibition of PP2A prevents internalisation of GABA_BRs in SST INs following their*
 203 *activation.* **A** Schematic of experimental conditions, highlighting treatment periods and
 204 fixation times **B** Example electron micrographs of SDS-FRL for SST-cre dependent
 205 expression of ChR2/YFP (18 nm immunoparticles) and GABA_{B1} (6 nm, red overlay) in *str.*
 206 *O/A* of CA1 of slices with vehicle or baclofen pre-application (20 μM). **C** The same labelling
 207 but performed in slices incubated with 50 nM fostriecin for 1 hour prior to baclofen pre-
 208 application. **D** Quantification of GABA_{B1} density in vehicle slices (Ctrl., black, 3 mice) or
 209 those pre-application with baclofen (Bacl., red, 2 mice) with and without fostriecin

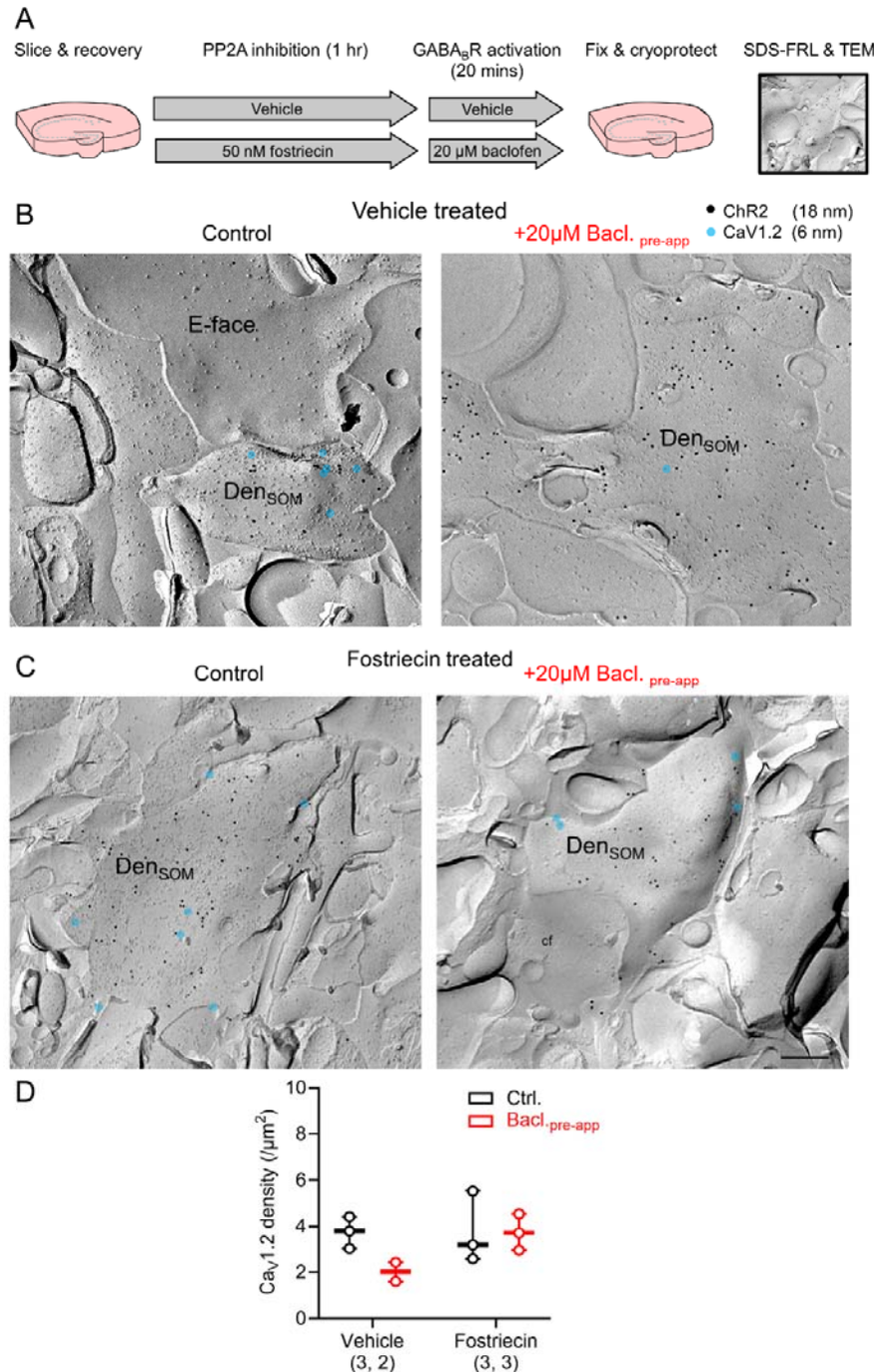
210 incubation. All data are shown as boxplots depicting the 25-75% range with median,
211 maximum range shown, and average density calculated from individual mice results
212 overlaid-as open circles. Scale bar represents 200 nm.

213

214 To determine whether GABA_BR-dependent internalisation of L-type calcium channels also
215 requires PP2A-dependent mechanisms, we performed the same experiments but labelling
216 for Ca_v1.2 (Figure 4A). In replicas from vehicle treated slices we observed Ca_v1.2 labelling
217 of 3.7 ± 0.3 particles/ μm^2 , which was 46% reduced in baclofen pre-application slices
218 (1.8 ± 0.3 particles/ μm^2 $t_{(3,2)}=3.32$, $p=0.0497$, Kenward-Roger test, Figure 4B, D),
219 consistent with with mGluR1 α labelling experiments (Figure 1). Fostriecin pre-incubation
220 (Figure 3C) prevented this reduction in Ca_v1.2 labelling following baclofen treatment
221 (3.5 ± 0.4 particles/ μm^2 $t_{(3,2)}=1.06$, $p=0.326$, Kenward-Roger test, Figure 3C, D), confirming
222 that PP2A inhibition prevents Ca_v1.2 channel internalisation.

223 These data confirm that baclofen induced GABA_BR internalisation requires PP2A in SST
224 INs in *str. O/A*, and that inhibiting this internalisation also prevents loss of Ca_v1.2 from
225 their dendritic membranes.

226



227

228 *Figure 4: Inhibition of PP2A prevents internalisation of Ca_v1.2 in SST INs following*
 229 *GABA_BR activation.* **A** Schematic of experimental conditions, highlighting treatment
 230 periods and fixation times. **B** Example electron micrographs of SDS-FRL for SST-cre
 231 dependent expression of ChR2/YFP (18 nm immunoparticles) and Ca_v1.2 (6 nm, blue
 232 overlay) in *str. O/A* of CA1 of slices with vehicle or baclofen (20 μ M). **C** The same labelling
 233 but performed in slices pre-application with 50 nM fostriecin for 1 hour prior to baclofen

234 application. **D** Quantification of Ca_v1.2 density in slices treated with vehicle (Ctrl., black, 3
235 mice) or baclofen (Bacl., red, 2 mice) with and without fostriecin pre-application. All data
236 are is shown as boxplots depicting the 25-75% range with median, maximum range
237 shown, and average density calculated from individual mice results overlaid—as open
238 circles. Scale bar represents 200 nm.

239

240 *Sustained GABA_BR activation leads to reduced plasticity of SST INs that is mediated by*
241 *receptor internalisation*

242 We have previously shown that aTBS induced LTP in SST INs can be prevented by
243 acutely activating GABA_BRs¹¹. However, our data suggests that application of baclofen
244 leads to reduced LTP machinery from SST INs. To determine whether sustained activation
245 of GABA_BRs shape the long-term plasticity landscape of SST INs, we first performed
246 whole-cell patch-clamp recordings in *str. O/A* from adult male and female transgenic mice
247 expressing YFP (Venus variant) under the vGAT promoter (Figure 5A). We recorded 108
248 *str. O/A* INs for the current study, those that were recovered (n=70) typically had horizontal
249 dendrites confined to *str. Oriens (str. O)*, with axons projecting to *Str. L-M* and were
250 uniformly SST immunoreactive (Figure 5B). Recorded SST INs displayed prominent I_h-
251 mediated sag potentials, and possessed a medium to high rate of action potential (AP)
252 discharge (Figure 5C), which was lower after baclofen pre-application; due to reduced
253 input resistance and elevated rheobase (Supplementary Table 1).

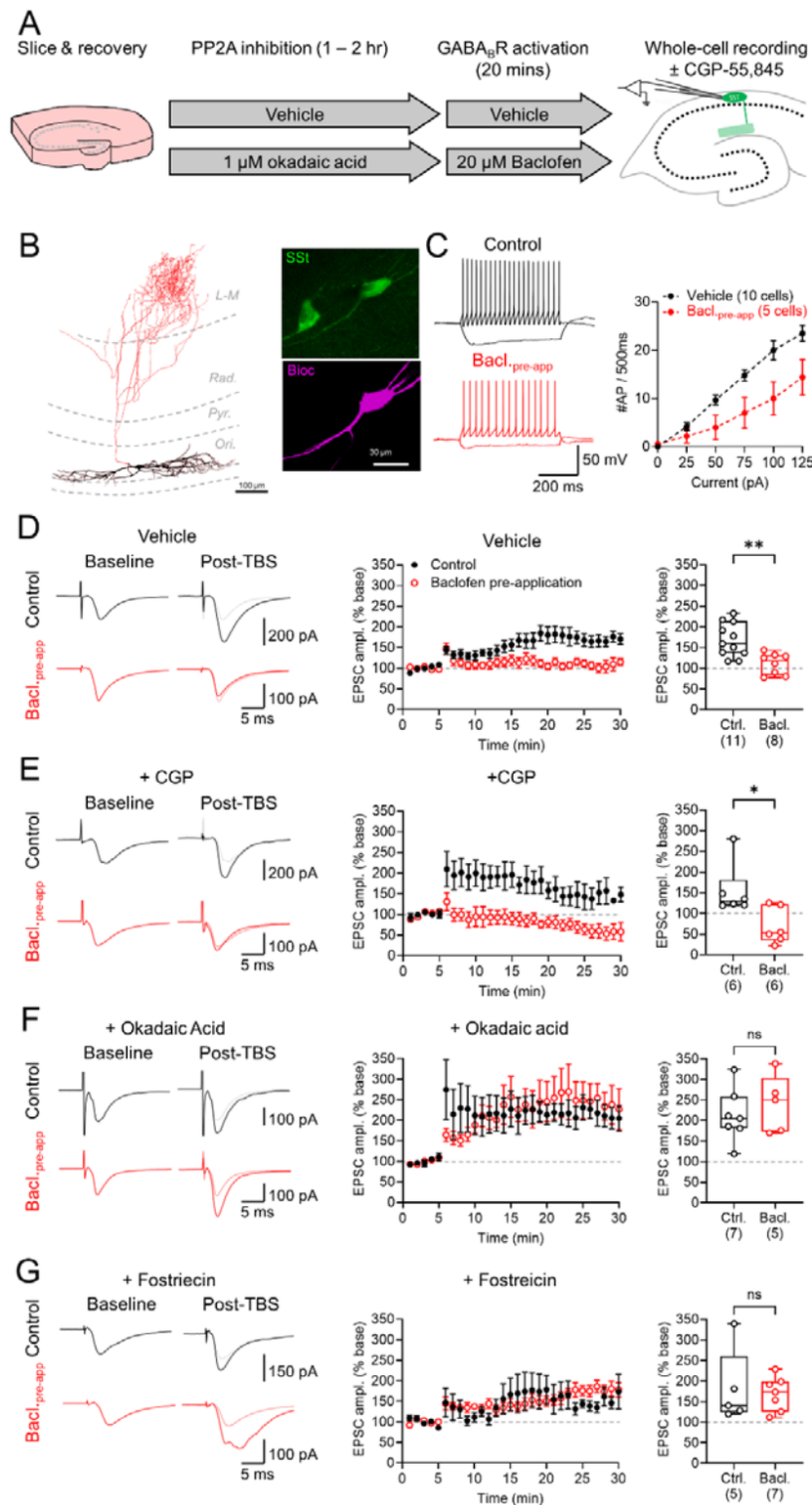
254 Under vehicle control conditions, we observed excitatory postsynaptic currents (EPSCs)
255 resulting from *alveus* stimulation of 197.8 ± 17.6 pA (13.3 ± 3.9 V stimulation, n=11 SST
256 INs), which following aTBS-LTP induction led to a facilitation of $68.5 \pm 12.9\%$ above
257 baseline levels (p=0.001, Wilcoxon matched-pairs test). In baclofen pre-application slices,
258 we observed baseline EPSCs with an average amplitude of 195.6 ± 54.6 pA (8.5 ± 2.9 V
259 stimulation, n= 8 SST INs), which did not differ from vehicle recordings in amplitude ($U_{(11, 8)} = 28$, P=0.21, Mann-Whitney) or stimulation strength ($U_{(11, 8)} = 33$, P=0.38, Mann-Whitney). An aTBS-LTP induction in baclofen pre-application slices failed to increase EPSC amplitude above baseline levels ($11.3 \pm 9.6\%$ above baseline, p=0.38, Wilcoxon matched-pairs test), and which was lower than for vehicle conditions ($U_{(11,8)} = 10$; p=0.004, Mann-Whitney). To confirm that this loss of aTBS-LTP was not due to residual direct activation of GABA_BRs we performed the same experiments, but in which the GABA_BR antagonist CGP-55,845 (CGP, 5 μM) was applied to the bath through-out recording

267 (Figure 5E). In the presence of CGP, we still failed to induce aTBS-LTP in baclofen pre-
268 application slices ($31.3 \pm 19.6\%$ below baseline, $p=0.31$, Wilcoxon matched-pairs test),
269 despite persistent induction in vehicle treated slices ($54.5 \pm 28.0\%$ above baseline
270 $p=0.031$, Wilcoxon matched-pairs test; $U_{(6,6)}=5$; $p=0.041$ Mann Whitey); confirming that
271 aTBS-LTP inhibition was not due to direct receptor activation.

272 As postsynaptic GABA_BRs undergo PP2A dependent internalisation (Figure 3)^{26,30,31,38},
273 we next asked if the impaired aTBS-LTP in SST INs was due to internalisation of the
274 receptor. For these experiments, we incubated slices with OA (1 μ M) a for 60-120 minutes
275 prior to vehicle or baclofen pre-application. Following OA incubation, we continued to
276 observe reliable aTBS-LTP in both vehicle slices ($112.1 \pm 26.4\%$ above baseline, $p=0.016$
277 Wilcoxon paired test) and baclofen pre-application slices in SST INs ($140.1 \pm 31.3\%$ above
278 baseline, $p=0.029$ Wilcoxon paired test), which were not different ($U_{(7,5)} = 15$; $p=0.76$,
279 Mann-Whitney test; Figure 5F). As our SDS-FRL labelling was not possible with OA
280 (Supplementary Figure 3), we next confirmed that fostriecin (50 nM) produced similar
281 effects to OA. After fostriecin incubation (60-90 minutes), we observed a tendency towards
282 aTBS-LTP in vehicle slices ($81.8 \pm 45.6\%$ above baseline, $p=0.063$ Wilcoxon paired test)
283 and robust LTP in baclofen pre-application slices ($69.0 \pm 17.0\%$ of baseline, $p=0.016$
284 Wilcoxon paired test), which was not different from vehicle ($U_{(5,7)}=16$, $p=0.876$, Mann-
285 Whitney test; Figure 5G).

286 Combined, these data indicate that sustained GABA_BR activation impairs aTBS-LTP
287 induction in SST INs, which is independent of direct receptor activation and which relies on
288 internalisation of the receptor via PP2A-dependent phosphorylation.

289



290 *Figure 5: LTP of local inputs to SST INs is abolished by prolonged baclofen application.* **A**
 291 Schematic of recording conditions, highlighting treatment periods and recording times. **B**
 292 Example reconstruction of a SST IN recorded from *str. O/A* of the adult mouse CA1 region,
 293 showing somatodendritic (black) and axonal (red) ramifications. Inset immunoreactivity for

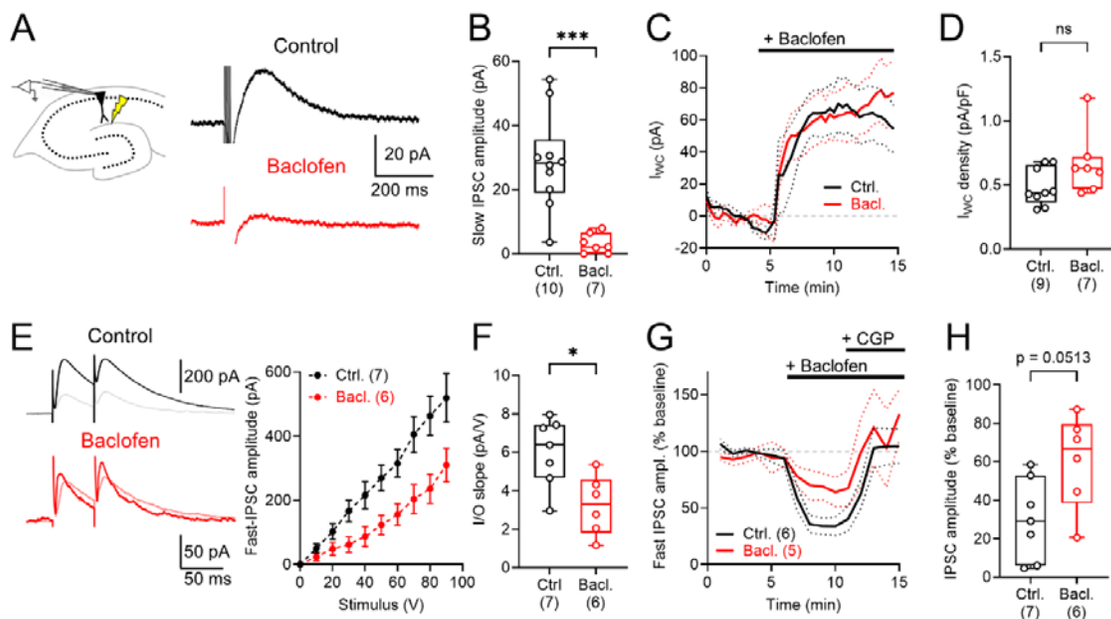
294 SST (green) and biocytin (purple). **C** Voltage responses in response hyper- to depolarising
295 current steps (-500 to +500 pA, 100 pA steps, 500 ms duration) from SST INs recorded in
296 following vehicle (black) and baclofen-pre-application (Bacl._{pre-app}; red). Right,
297 quantification of the action potential output of SST INs recorded under both conditions. **D**
298 Example EPSC traces from vehicle (upper, black) and 20 μ M baclofen pre-application
299 (lower, red) measured under control conditions; for aTBS-LTP traces baseline EPSCs are
300 shown underlain (grey and pink). Middle, time-course of plot of EPSC amplitude before
301 and after aTBS-LTP induction for vehicle (black) and baclofen pre-application (red); 100%
302 baseline level is shown for reference (grey dashed line). Right, bar-chart of EPSC
303 amplitude change post aTBS-LTP (25-30 minutes) compared to baseline for vehicle
304 (black) and baclofen pre-application (red); 100% baseline level indicated (grey dashed
305 line). **E** aTBS-LTP recordings from SST INs performed in the presence of 5 μ M CGP-
306 55,845 (CGP) according to the same scheme as **D**. **F** aTBS-LTP following pre-application
307 of slices with 1 μ M okadaic acid for 1 hour with subsequent vehicle or baclofen pre-
308 application. **G** aTBS-LTP in slices pre-application with 50 nM fostriecin for 1 hour prior to
309 vehicle or baclofen pre-application. All data are shown as either mean \pm SEM (time-course
310 plots) or box-plots showing 25-75% box with median, with maximum range shown and
311 individual cell results. Statistics shown as: ns – $p > 0.05$, * - $p < 0.05$, ** - $p < 0.01$ from Mann-
312 Whitney non-parametric tests.

313

314 *Prolonged activation of GABA_BRs impairs presynaptic release of GABA, but not*
315 *postsynaptic expression of GABA_BRs in CA1 PCs*

316 Assessing the wider function of GABA_BR signalling following sustained pharmacological
317 activation is critical to infer circuit wide effects. In CA1 PCs, following baclofen pre-
318 application we observed a tendency towards reduced GABA_{B1}, but unchanged Ca_v1.2
319 surface expression. We next determined if these effects altered inhibitory signalling in and
320 onto CA1 PCs. Therefore, we recorded GABA_BR-mediated slow IPSCs in the presence of
321 CNQX (10 μ M), DL-AP5 (50 μ M) and gabazine (10 μ M) to block ionotropic receptor
322 signalling (Figure 6A). We found that slow IPSCs were substantially reduced following
323 baclofen pre-application ($U_{(10,7)}=3$, $p=0.0007$, Mann-Whitney test; Figure 6B). However,
324 baclofen-mediated whole-cell currents (I_{WC}) were unaffected by baclofen pre-application
325 (Figure 6C), with normal peak current-density ($U_{(9,7)}= 15.5$, $p=0.10$, Mann-Whitney test;
326 Figure 6D). One explanation for this disconnect between synaptic and pharmacological
327 activation of GABA_BRs could be impaired GABA release. As such, we measured the

328 input/output (I/O) relationship of ionotropic GABA_AR-mediated IPSCs following *str. L-M*
 329 stimulation as a proxy for synaptic strength. We found that for the same stimulus, fast
 330 IPSCs were consistently lower in amplitude (Figure 6E), which upon quantification
 331 revealed 46% lower I/O slopes ($U_{(7,6)}=4$, $p=0.014$, Mann-Whitney test, Figure 6F). To
 332 determine whether baclofen pre-application alters presynaptic GABA_BR function, we next
 333 bath applied 10 μ M baclofen to recordings of fast IPSCs (Figure 6G). Slices with pre-
 334 application of baclofen tended to have lower GABA_BR presynaptic inhibition of IPSCs in
 335 *str. L-M* ($U_{(7,6)}=7$, $p=0.051$, Mann-Whitney test, Figure 6H). Finally, we compared the effect
 336 of baclofen pre-application on CA1 PC intrinsic excitability. Contrary to what we observed
 337 in SST INs, we find that CA1 PCs are largely unaffected by baclofen pre-application, with
 338 subtle effects on membrane capacitance and action potential kinetics observed (Table S2).
 339 Together these data suggest that prolonged GABA_BR activation has limited effects on the
 340 postsynaptic function of CA1 PCs, in particular we find no evidence for reduced functional
 341 GABA_BR expression. However, due to strong reductions in presynaptic GABA release,
 342 postsynaptic IPSCs are functionally reduced.



343

344 **Figure 6: Prolonged GABA_BR activation decreases GABA release in *str. L-M*, specific for**
 345 **presynaptic but not postsynaptic GABA_BR-mediated inhibition.** **A** Experimental schematic
 346 of recordings from CA1 PCs and example slow IPSCs recorded following 5x 200 Hz
 347 stimulation of *str. L-M* in the presence of CNQX (10 μ M), DL-AP5 (50 μ M) and gabazine

348 (10 μ M) from -65 mV voltage-clamp from control slices (black) and those pre-application
349 with 20 μ M baclofen (red). **B** quantification of slow IPSC amplitudes from control and
350 baclofen pre-application slices. **C** Time course of whole-cell current (I_{WC}) recorded from -
351 65 mV following bath application of 10 μ M baclofen in control and baclofen pre-application
352 slices. **D** Quantification of I_{WC} current density, normalised to cell capacitance. **E** Fast
353 IPSCs evoked following paired stimulation of *str. L-M* from 0 mV using Cs-gluconate
354 internal solution in the presence of CNQX (10 μ M), DL-AP5 (50 μ M) from control and
355 baclofen pre-application slices. Traces recorded following bath application of 10 μ M
356 baclofen are shown underlain (grey/pink). Right, input-output (I/O) plot of fast IPSC
357 amplitude from both groups. **F** Quantification of I/O slope for fast IPSCs. **G** Time course of
358 fast IPSC amplitude before, and following bath application of 10 μ M baclofen and 5 μ M
359 CGP-55,845 (CGP) in control and baclofen pre-application slices. **H** Quantification of
360 baclofen mediated inhibition of fast IPSC amplitudes, expressed as % of baseline. Data is
361 shown as either mean \pm SEM (**C**, **E**, **G**) or box-plots showing 25-75% box with median,
362 with maximum range (**B**, **D**, **F**, **H**). Individual cell data is shown overlaid. Statistics shown
363 as: ns – $p > 0.05$, * - $p < 0.05$, *** - $p < 0.001$; from Mann Whitney non-parametric tests.

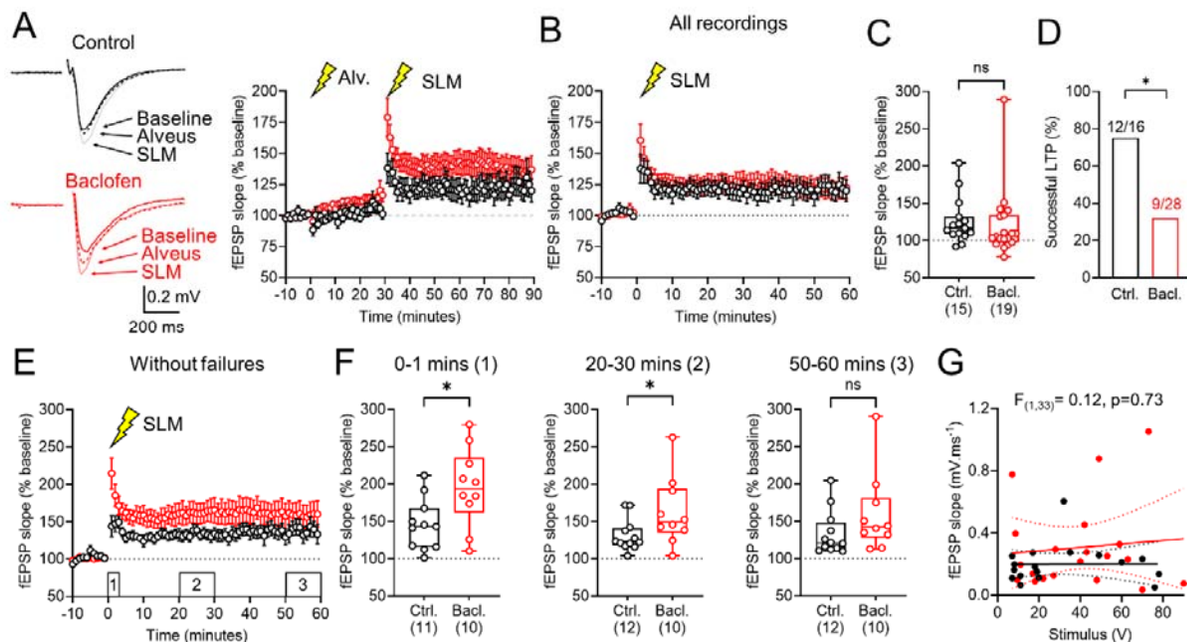
364

365 *GABA_BR-dependent internalisation of LTP proteins in SST INs leads to enhanced*
366 *temporoammonic plasticity onto CA1 PCs*

367 SST INs are known to control the ability of CA1 PCs to undergo plasticity, notably in their
368 distal dendrites in *str. L-M* aligned with entorhinal cortex inputs²³. To determine whether
369 prolonged activation of GABA_BRs, and subsequent loss of SST IN plasticity alter
370 temporoammonic LTP we performed extracellular field recordings from *str. L-M* of CA1, in
371 which we induced plasticity at SST INs via repetitive stimulation of the alveus, followed by
372 high frequency stimulation (HFS, 1x 100 Hz) to the temporoammonic pathway in vehicle or
373 those with pre-application of 20 μ M baclofen for 20 minutes (Figure 7A). Following HFS to
374 *str. L-M* we observed increased fEPSP amplitudes when measured in all recordings from
375 both treatment groups (Figure 7B), which when measured at 50-60 minutes post induction
376 were $25.6 \pm 7.8\%$ above baseline in vehicle control recordings, and which did not differ in
377 baclofen pre-application slices ($20.5 \pm 10.4\%$; $U_{(15,19)}=104$, $p=0.19$, Mann-Whitney test,
378 Figure 7C). However, we noted that a higher proportion of slices pre-application with
379 baclofen failed to undergo LTP ($>10\%$ above baseline) as compared to vehicle controls
380 ($\chi^2=7.5$, $p=0.0062$ Chi square test; Figure 7D). Therefore, we next analysed only those
381 recordings in which LTP was observed (Figure 7E). Further quantification of successful

382 LTP recordings (Figure 7F) revealed that post-tetanic potentiation (0-1 minute post HFS)
 383 was 34% higher in baclofen pre-application slices ($U_{(11,10)}=25$, $p=0.018$, Mann-Whitney
 384 test) and fEPSPs measured at 20-30 minutes (consistent with whole-cell recordings) were
 385 also 24% higher than control slices ($U_{(12,10)}=27$, $p=0.015$, Mann-Whitney test), while LTP
 386 measured at 50-60 minutes was higher, albeit not significantly so ($U_{(12,10)}=37$, $p=0.07$,
 387 Mann-Whitney test). We observed no difference in fEPSP slope relative to stimulus
 388 strength between control and baclofen pre-application slices (Figure 7G). These effects
 389 were likely specific to SST INs, as performing the same recordings but in the presence of
 390 the mGluR1 α specific antagonist LY367,385 (100 μ M), prevented any baclofen pre-
 391 application effects on temporoammonic LTP (Figure S4). This also excludes the effects of
 392 baclofen pre-application on GABAergic release and CA1 PC excitability as being causal to
 393 this effect.

394 These data confirm that prolonged GABA_BR activation leading to a down-regulation in
 395 surface expression of GABA_{B1}, mGluR1 α , and Ca_v1.2 in SST INs has profound effects on



396 the ability of the CA1 microcircuit to encode inputs in *str. L-M*.

397 **Figure 7: Prior GABA_BR activation abolishes SST IN inhibition of short-term**
 398 **temporoammonic LTP.** **A** Example fEPSP responses recorded in *str. L-M* following
 399 temporoammonic stimulation, from control (black) and 20 μ M baclofen pre-application
 400 (red) slices. Traces are shown for initial baseline (solid line), following *alveus* LTP
 401 induction (dashed lines) and following *str. L-M* 1x 100 Hz HFS (grey/pink lines). Right,

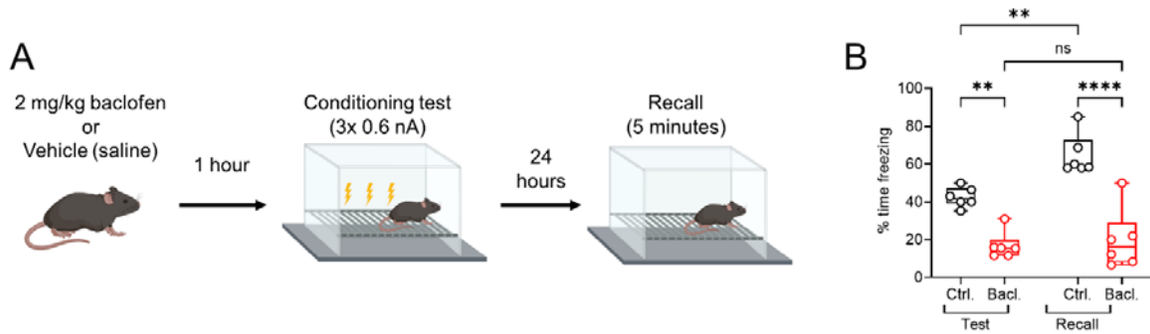
402 summary time-course of all fEPSP recordings normalised to the initial baseline, indicating
403 LTP induction points (lightning bolts). **B** Time-course of LTP induced in *str. L-M* (SLM,
404 lightning bolt), normalised to the 20-30 minutes baseline following *alveus* stimulation in all
405 recordings. **C** Magnitude of fEPSP facilitation at 50-60 minutes post *str. L-M* HFS in all
406 recordings. **D** Proportion of recordings in both control and baclofen pre-application that
407 successfully induced LTP (>10% facilitation at 50-60 minutes post HFS). **E** Time-course of
408 LTP induced in *str. L-M* normalised to the 20-30 minutes baseline following *alveus*
409 stimulation in recordings where LTP was induced. **F** Quantification of fEPSP facilitation in
410 control and baclofen pre-application slices at 0-1 (left), 20-30 (middle), and 50-60 (right)
411 minutes post *str. L-M* HFS. **G** comparison of fEPSP slope and stimulus strength for all
412 recordings in both groups. Data is shown as either mean \pm SEM (**A, B, E**) or box-plots
413 showing 25-75% box with median, with maximum range (**C, F**), proportions (**D**), or as
414 individual data (**g**). Individual cell data is shown overlaid. Statistics shown as: ns – $p > 0.05$,
415 * - $p < 0.05$; from Mann Whitney non-parametric tests.

416

417 *Activation of GABA_BRs prevents acquisition of contextual fear memories*

418 Finally, as SST IN plasticity has been causally linked to contextual fear conditioning when
419 administered during training³⁹, we asked if earlier baclofen administration impairs fear
420 learning. For this, we administered mice with 2 mg/kg baclofen (or saline controls, both
421 intraperitoneal injection) 1 hour prior to fear conditioning. Mice were then administered 3
422 foot-shocks (0.4 nA, 1-minute intervals, 5 minutes total) in the conditioning arena, and the
423 proportion of time spent freezing measured. To determine contextual memory was
424 retained by mice, they were returned to the same arena 24 hours later for 5 minutes
425 (Figure 8A). Overall, saline control mice froze $42 \pm 2\%$ of the time during the initial test,
426 increasing to $65 \pm 4\%$ freezing upon recall ($p = 0.004$, Holm-Sidak test). By comparison,
427 mice treated with baclofen 1 hour before training froze $17 \pm 3\%$ of the time, which did not
428 increase upon recall ($20 \pm 7\%$, $p = 0.63$, Holm-Sidak test), but which was markedly lower
429 than saline controls ($p < 0.0001$, Holm Sidak test; Figure 8B). These data confirm that prior
430 administration of baclofen produces sustained impairments in contextual fear conditioning.

431 This data suggests that prolonged GABA_BR activation – as experience with in vivo
432 administration - likely has lasting effects on the ability of hippocampal circuits to perform
433 complex behavioural tasks and may provide important considerations for the use of
434 baclofen therapeutically.



435 *Figure 8: Prior GABA_BR activation impairs contextual memory formation.* **A** Schematic of
436 contextual fear conditioning experiments, in which mice were dosed with 2 mg/kg baclofen
437 1 hour prior to training. **B** Quantification of % time freezing during the test and recall
438 phases of conditional fear conditioning. Data is shown as box-plots showing 25-75% box
439 with median, with maximum range. Individual cell data is shown overlaid. Statistics shown
440 as: ns – $p > 0.05$, ** - $p < 0.01$, **** - $p < 0.0001$; from 2-way ANOVA with Holm-Sidak post-
441 tests.

442

443 Discussion

444 In this study we tested the hypothesis that sustained GABA_BR activation leads to receptor
445 internalisation in hippocampal SST INs, releasing inhibition of synaptic plasticity. We
446 disproved this hypothesis and show baclofen administration leads to internalisation of
447 GABA_BRs, Ca_v1.2 and also mGluR1 α in SST INs, which is mediated by PP2A-dependent
448 phosphorylation. This internalisation impairs the long-term ability of SST INs to undergo
449 synaptic plasticity, which in turn shifts the balance of plasticity at temporoammonic inputs
450 and impairs contextual fear memory. These data reveal a novel mechanism of long-term
451 disinhibition that has the potential to shape the plasticity landscape of the CA1 region over
452 behaviourally relevant time-scales.

453 *PP2A-dependent GABA_BR internalisation regulates SST IN activity*

454 SST INs are a critical element in cortical circuits, gating the strength of inputs, and their
455 plasticity, arriving from entorhinal cortex¹⁹, and associative inputs within the hippocampus
456 itself^{22,24}. These effects are mediated by a complex interplay of direct inhibitory actions on
457 postsynaptic CA1 PCs, via presynaptic inhibition of glutamatergic and GABAergic inputs
458 and through regulation of astrocyte function⁴⁰. Indeed, SST IN activity leads to profound
459 disinhibition of the hippocampal circuit to strengthen CA3 inputs²² and limit CA1 PC

460 dendritic function²⁴. Thus, plasticity of SST INs themselves may act to shift the balance of
461 CA1 function²³, which has a direct effect on CA3-dependent behaviours involving
462 contextual information⁴¹. Our work has shown that GABA_BRs acutely inhibit L-type
463 VGCCs comprising Ca_v1.2 subunits on SST INs to inhibit LTP¹¹, which led to our
464 hypothesis that GABA_BR internalisation would promote LTP induction, as this key
465 inhibitory mechanism was lost. We have failed to prove this hypothesis, with the opposite
466 appearing to be the case, due to coincident internalisation of the key LTP machinery,
467 namely mGluR1 α and L-type VGCCs; and which depends on activation of PP2A and that
468 has been implicated in GABA_BR phosphorylation in other brain areas^{26,30,31,38}.

469 Following GABA_BR activation, adenylyl cyclase activity is inhibited, leading to lower cAMP
470 levels and reduced PKA activation, which in turn limits the phosphorylation of Serine 892
471 on the GABA_{B2} subunit, which destabilises the receptor at the cell surface⁴². Additionally,
472 PP2A dephosphorylates Serine 783 on the GABA_{B1} subunit, which also destabilises the
473 receptor leading to internalisation²⁶. mGluR1 α is also a Type C G-protein coupled
474 receptor (GPCR), thus it is highly plausible that these receptors are targeted for
475 internalisation through the same intracellular cascade⁴³, particularly given PP2A has been
476 shown to regulate recycling of mGluR1 isoforms⁴⁴. For Ca_v1.2, the situation is more
477 complex, as it is internalised upon GABA_BR activation which appears to be prevented by
478 PP2A inhibition, while functional currents are not restored. Given the close proximity of
479 GABA_{B1} to Ca_v1.2 in SST IN dendrites¹¹ it may form part of the GABA_BR interactome in
480 SST INs, as is the case for other Ca_v proteins²⁹, thus function is impaired when GABA_BRs
481 are lost. However, Ca_v1.2 conductance is regulated by PKA phosphorylation⁴⁵, thus, loss
482 of GABA_BR-mediated inhibition may counterbalance a numerical loss of channels. Beyond
483 VGCCs, native GABA_BRs interact with a variety of transmembrane and cytosolic proteins
484 in pyramidal cells^{28,29}, if the same interactome is present in SST INs and whether these
485 proteins are similarly lost from these cells remain unexplored. Indeed, the functional
486 association of GABA_BRs with other postsynaptic effectors (e.g. Kir3 channels) may differ in
487 diverse INs^{8,9,12}. As INs make up only approximately 10% of neurons in the hippocampus
488⁴⁶, efforts to identify the GABA_BR interactome using bulk protein samples²⁹ have likely
489 underestimated cell-type specific effects. Future recent advances in single-cell proteomic
490 analysis⁴⁷ may allow detection of cell-type specific GABA_BR interactomes, crucial to
491 determine such function. Our data provides limited evidence of impaired postsynaptic
492 GABA_BR function in CA1 pyramidal cells under the same conditions, further suggesting
493 cell-type specificity.

494

495 *GABA_BR internalisation on SST INs impairs hippocampal circuit function*

496 It has long been appreciated that acute GABA_BR activation leads to paradoxical functions
497 at the circuit level, notably enhanced LTP^{48,49} and increase seizure activity⁵⁰, despite
498 directly hyperpolarising membranes^{51,52} and inhibiting pre- and postsynaptic Ca²⁺
499 channels^{53,54} on PCs. These effects point to a function beyond direct inhibition (favouring
500 a disinhibitory model), exemplified by GABA_BR inhibition on multiple IN types across the
501 hippocampus⁴. Our data extends this, providing evidence that prolonged GABA_BR
502 activation fundamentally shifts the ability of SST INs to undergo LTP on behaviourally
503 relevant timescales. Given that SST IN LTP has been well associated with numerous
504 behavioural functions, including spatial and contextual memory^{24,41,55-57}, this indicates a
505 potential role of GABA_BR in regulating this activity. We confirm that LTP in SST INs is
506 crucial to maintain transfer of information to CA1 via the TA path, as previously reported¹⁹,
507 the control of which is impaired when baclofen is pre-applied. Furthermore, we show that
508 GABA_BR activation prior to contextual fear conditioning is sufficient to prevent memory
509 encoding. While baclofen has been shown to impair fear conditioning previously³⁹, these
510 studies relied on administering the drug immediately (<15 minutes) before fear acquisition,
511 suggesting direct inhibitory effects. The pharmacokinetics of baclofen suggest peak brain
512 concentrations with 30 minutes of IP administration, which remain high for several hours
513⁵⁸. Therefore, it is nigh on impossible to estimate when (and if) loss of SST IN LTP
514 supersedes inhibition driven by GABA_BR inhibition. Future studies investigating GABA_BR
515 surface localisation on SST INs over hours and days post baclofen injection may reveal
516 the temporal resolution of this relationship, but are outside the scope of this project.

517

518 *Translational importance*

519 Baclofen is a common and important medicine, primarily prescribed for muscle spasm, but
520 also tested for alcohol dependency, neurodevelopmental disorders, among other
521 conditions (reviewed in⁵⁹), up to 100 mg/day; but is associated with many common
522 neurological side effects. Thus, understanding how baclofen affects cortical circuits may
523 inform its best use, particularly in conditions where GABAergic inhibition has also been
524 implicated.

525 One of the most common clinical features associated with baclofen administration is the
526 paradoxical induction of seizure activity, both in patients⁶⁰ and in rodent models⁵⁰.

527 Originally proposed as a potential anti-seizure medication ⁶¹, effects of baclofen on
528 hippocampal circuits has previously been described, leading to seizure like activity ⁶². The
529 baclofen-dependent loss of SST IN excitatory signalling we observe may favour long-term
530 reduction in local SST IN inhibition, which would be predicted to cause seizures ⁶³.
531 Interestingly, we recently showed higher GABA_BR signalling in PCs in patients who have
532 experienced seizures ⁶⁴. It is tempting to speculate that such increases may result from
533 reduced inhibitory tone, leading to compensatory upregulation of receptor expression.
534 Further study is required to determine if this is the case.

535 A recent off-label use for high-dose baclofen has been in treating addiction, particularly
536 alcoholism use disorder ⁶⁵. In rodents, GABA_BRs have been shown to be lost in the cortex
537 following chronic alcohol administration ⁶⁶ and ventral tegmental area (VTA) following
538 administration of amphetamine or cocaine ²⁶, albeit not tested in SST INs. However, it has
539 been shown that acute alcohol administration bidirectionally affects SST INs, with low
540 doses increasing activity and high doses attenuating activity ⁶⁷. SST INs are present in
541 cortical ^{15,68} and VTA ⁶⁹ circuits, and their activity is strongly downregulated in chronic
542 alcohol consuming mice, which may be corrected by inhibiting them directly⁷⁰. Our data
543 suggests that prolonged baclofen administration may prevent SST IN activity, akin to
544 chemogenetic approaches used previously, thus may reflect a mechanism by which
545 baclofen could produce therapeutic benefit.

546

547 *Limitations:*

548 We have identified some key limitations of our current study. First, while we show that
549 internalisation of GABA_BRs from SST IN dendritic membranes occurs following 20 minutes
550 of baclofen application, using our approach we were not able to identify the specific
551 temporal dynamics of this loss. If such effects occur rapidly (within seconds) or require
552 longer (minutes to hours) remains unclear. Nevertheless, baclofen as a therapy is often
553 administered at moderate to high doses (up to 1g/day) and has a CSF half-life of several
554 hours ⁵⁸, thus such temporal dynamics – whilst mechanistically interesting – are somewhat
555 negated by these pharmacokinetics. Further study identifying whether GABA_BR
556 internalisation on SST INs displays dose- and time-dependence would provide useful
557 insight to baclofen's therapeutic use. Furthermore, we have not identified how long it takes
558 SST INs to recover LTP following baclofen administration, which may also provide
559 important therapeutic information.

560 One unusual observation in our data was that okadaic acid appeared to occlude labelling
561 of GABA_{B1} in SDS-FRL experiments, both on SST INs and CA1 PCs (Figure S3). We do
562 not have a good explanation for this effect. One possibility is that SDS-FRL relies on
563 immunolabelling of receptors⁷¹ and okadaic acid as a large polypeptide interacts with an
564 epitope on PP2A close to its interaction site with GABA_{B1}. Such an interaction may mask
565 or occlude antibody binding, that would otherwise not be observed with the use of small
566 molecule pharmacology, such as fostriecin. A similar phenomenon has been proposed for
567 the binding of the amyloid precursor protein (APP) to GABA_{B1}, which has been indicated in
568 SDS-FRL to be diminished in APP overexpressing mice⁷², but in which mice GABA_BR
569 function is unaltered⁷³. These data indicate the importance of providing multiple lines of
570 evidence to support conclusions based on physiology or anatomy alone, which we have
571 provided through comparative use of okadaic acid and fostriecin that both selectively
572 inhibit PP2A⁷⁴.

573 Finally, our data suggest that in SST INs GABA_BRs likely interact with themselves, and
574 influence the function of – mGluR1 α , Ca_v1.2, and PP2A. Such proteins do not appear on
575 previously published interactomes of GABA_{B1} or GABA_{B2} in the hippocampus²⁹. Our data
576 suggest the existence of cell-type-specific interactomes, which may not be detectable at a
577 whole brain, or even region-specific level. Such an endeavour would almost certainly
578 identify cell-type specific differences even within neurochemically defined classes, as
579 highlighted by previous work^{4,12}. However, such analyses are beyond the scope of the
580 current study. Future work should seek to identify which proteins GABA_BRs interact with, in
581 a cell-type specific manner, and potentially if such interactions change with age or
582 neuropathology.

583

584 *Summary and outlook:*

585 Here we show in hippocampal SST INs that prolonged activation of GABA_BRs is sufficient
586 to cause internalisation of the receptor through PP2A-dependent mechanisms, which also
587 leads to loss of mGluR1 α and Ca_v1.2 to prevent synaptic plasticity at CA1 PC inputs.
588 These data provide compelling evidence of how the inhibitory molecule baclofen exerts a
589 paradoxical disinhibitory effect by reducing SST IN function, which has relevance for
590 hippocampal function and disease.

591

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598

599 **Author Contributions:** Conceptualisation: SAB, AK; Methodology: NS, SAB, MW, AS,
600 DL, AK; Validation: SAB, AK; Formal Analysis: SAB, AK, NS, MW, AS; Investigation: SAB,
601 NS, MW, DL, AS, RL; Writing original draft: SAB; Writing – review and editing: SAB, NS,
602 IV, AK; Supervision: SAB, IV, AK; Funding acquisition: SAB, IV, AK.

603

604 **Declaration of interests:** The authors declare no competing interests.

605

606 **Materials and methods:**

607 *Animals:*

608 All *ex vivo* electrophysiology was performed in brain slices prepared from male and female
609 mice (2-4 months old), with all procedures performed according to Home Office (ASPA,
610 2013) and The University of Edinburgh Ethical Board guidelines. Mice were maintained on
611 a C57/Bl6J^{CRL} background and expressed yellow-fluorescent protein (YFP) under the
612 vesicular GABA transporter (vGAT)⁷⁵. Mice were housed on a 12h light/dark cycle, with *ad*
613 *libitum* access to food and water.

614 For SDS-FRL, male mice (8-week-old, n=12) either wild-type or in which SST-INs
615 selectively expressed Channelrhodopsin2(ChR2)-YFP fusion protein were derived from
616 crossing SST-Cre and Ai32^{RCL-ChR2(H134R)/EYFP} transgenic mice. These mice and male
617 Wistar rats (8-week-old, n = 3) were used for quantitative immuno electron microscopic
618 analysis. For behavioural analysis, male C57/Bl6J mice (8–12 weeks old, n = 6 per group)
619 were used. Care and handling of the animals prior to and during the experimental
620 procedures followed European Union and national regulations (German Animal Welfare
621 Act) and all experiments were performed in accordance with institutional guidelines
622 (University of Freiburg, Germany) with permission from local authority (Freiburg, X21/04B).

623

624 *Acute brain slice preparation:*

625 Brain slices were prepared as previously described⁷⁶. Mice were terminally anaesthetised
626 with isoflurane, decapitated, then their brains rapidly dissected into ice-cold sucrose-
627 modified artificial cerebrospinal fluid (sucrose-ACSF; ACSF; in mM: 87 NaCl, 2.5 KCl, 25
628 NaHCO₃, 1.25 NaH₂PO₄, 25 glucose, 75 sucrose, 7 MgCl₂, 0.5 CaCl₂) which was
629 saturated with carbogen (95% O₂/5% CO₂). Brains were glued to a vibratome stage (Leica
630 VT1200S, Leica, Germany), then 300 μm (for whole-cell recordings) or 500 μm (for
631 extracellular recordings) horizontal slices containing the hippocampi were prepared.
632 Following slicing, brain slices were placed in either: a submerged holding chamber
633 containing sucrose-ACSF warmed to 35 °C for 30 min, then at room temperature; or on
634 small squares of filter paper placed in a liquid/gas interface chamber, containing recording
635 ACSF (in mM: 125 NaCl, 2.5 KCl, 25 NaHCO₃, 1.25 NaH₂PO₄, 25 glucose, 1 MgCl₂, 2
636 CaCl₂) and bubbled with carbogen.

637 *Whole-cell patch clamp recordings:*

638 Briefly, 400 μm horizontal hippocampal slices were prepared from 60-90 day-old C57/Bl6J
639 mice as previously described⁷⁷. For recording, slices were transferred to a submerged
640 chamber maintained at $31 \pm 1^\circ\text{C}$. Whole-cell recordings were made using pipettes filled
641 with either K-gluconate or Cs-gluconate based solutions. Voltage-clamp recordings were
642 performed from a potential of either -65 mV or -60 mV, while current-clamp recordings
643 were made from resting membrane potential. Intrinsic properties of recorded neurons were
644 characterized on their voltage response to hyper- to depolarizing current steps, in current-
645 clamp. HVA calcium transients, mGluR EPSCs, and GABA_BR-mediated currents were
646 measured in the presence of 10 μM CNQX, 50 μM DL-AP-5, and 50 μM picrotoxin.
647 GABA_AR IPSCs were elicited by placing a bipolar stimulating electrode in str. lacunosum-
648 moleculare approximately 200-300 μm rostral to the recorded neuron, in the presence of
649 DL- 10 μM CNQX, 50 μM DL-AP-5. LTP was induced at alveus inputs to SST INs with
650 EPSCs elicited by a bipolar stimulating electrode placed in the alveus, in the presence of
651 50 μM picrotoxin. Following baseline, associative LTP was induced by theta-burst
652 stimulation paired with a depolarization to -20 mV, repeated 3 times at 30 second intervals.
653 EPSCs were measured for 25 minutes following induction. For all stimulation, the pulse
654 duration was 200 μs delivered via a constant voltage generator. For HVA VGCC activation,
655 once a whole-cell patch-clamp recording was obtained using a Cs-Gluconate internal
656 solution (containing QX-314), the cell was allowed to dialyse for 5 minutes. Then at -60 mV
657 voltage-clamp, 3 families of depolarizing steps were applied (-60 to 20 mV, 5 mV steps,
658 500 ms duration). The L-type VGCC blocker nifedipine (20 μM) was then applied to the
659 bath for 3 minutes and 3 further families of depolarizing steps obtained. All traces were
660 then P/N subtracted based on the 5 mV step and the peak current measured. The peak
661 current under nifedipine was then subtracted from control. For mGluR EPSC experiments,
662 slow-EPSCs were elicited by focal puff application of the Group 1 mGluR agonist DPHG
663 (100 μM , 500 ms, 20 psi, diluted from stock in 150 mM NaCl) delivered via a patch pipette
664 placed \sim 50 μm from the cell body. The specificity of the mGluR current was assessed by
665 subsequent bath application of the mGluR1 α -specific antagonist LY-367,385 (100 μM).

666 For pre-application experiments, slices were transferred to a holding chamber containing
667 recording ACSF and 20 μM R-baclofen for 20 minutes, slices were then placed in the
668 recording chamber and rinsed with ACSF without R-baclofen for 4-6 minutes prior to
669 recording. To block PP2A activity, slices were transferred to a chamber containing ACSF
670 and 1 μM Okadaic Acid for at least 3 hours prior to recording or baclofen pre-application.
671 All other drugs were applied directly to the perfusing ACSF.

672

673 *Field recordings:*

674 For field excitatory postsynaptic potential (fEPSP) recordings, hippocampal slices were
675 transferred to an interface recording chamber perfused with carbogenated artificial
676 cerebrospinal fluid (ACSF) at 2–3 mL/min and maintained at 30 ± 1 °C. Recording
677 electrodes (1–3 M Ω) were pulled from borosilicate glass capillaries (1.5 mm outer / 0.86
678 mm inner diameter; Harvard Apparatus, UK) using a horizontal puller and filled with
679 recording ACSF. Slices were visualised with a wide-field microscope (Leica, Germany),
680 and electrodes were positioned in the str. L-M of CA1.

681 Extracellular fEPSPs were evoked using a paired-pulse protocol delivered via a bipolar
682 stimulating electrode placed in *str. L-M*, approximately 500 μ m to 1 mm from the recording
683 electrode, to activate the temporoammonic pathway. A second stimulating electrode was
684 positioned in parallel within the alveus of CA1. To activate GABA -B receptors, slices were
685 perfused with 20 μ M baclofen for 20 min, followed by a 20-min washout period.

686 A 10-min baseline was recorded under control conditions or following baclofen washout.
687 LTP was first induced via alveus stimulation using a high-frequency stimulation (HFS)
688 protocol consisting of three 100 Hz trains of 20 pulses delivered 30 s apart, or a protocol
689 consisting of five 100 Hz trains of 5 pulses, delivered 30 s apart. LTP was then induced in
690 *str. L-M* using an HFS protocol of 1x 100 Hz train of 100 pulses delivered 30 s apart.

691 Potentiation was monitored for 60 min following induction. LTP magnitude was quantified
692 as the mean fEPSP slope measured 50–60 min post-induction, normalised to the 10-min
693 baseline. LTP was considered successful if the 50–60 min slope exceeded baseline by
694 >10%. Recordings were excluded if baseline stability, assessed by comparing mean
695 slopes at 1–2 min and 9–10 min, varied \pm 10%. Signals were filtered online (1 Hz high-
696 pass, 500 Hz low-pass) and digitised at 10 kHz. Data were acquired and analysed offline
697 using WinLTP (v3.01, University of Bristol, UK).

698 *Neuronal visualization and immunohistochemistry:*

699 All neurons were filled with biocytin during recording, fixed overnight in 4% PFA and
700 labelled with streptavidin and antibodies to SST according to previous methods⁷⁸. Slices
701 were washed in PBS then blocked for 1 hour at room temperature (10% NGS, 0.3% Triton
702 X-100 and 0.05% NaN₃ in PBS). Following blocking slices were incubated with primary
703 antibodies at 4°C for 48h (1:1000 rabbit SST-14, T-4103, BMA Biomedicals, Switzerland;

704 5% NGS, 0.3% Triton X-100 and 0.05% NaN_3 in PBS), washed in PBS then transferred to
705 a secondary antibody solution containing fluorescent-conjugated streptavidin (1:500 goat
706 anti-rabbit AlexaFluor 488, 1:500 Streptavidin AlexaFluor 633, 3% NGS, 0.1% Triton X-
707 100, 0.05% NaN_3). Following labeling the slices were washed in 0.1M PB and mounted on
708 glass slides with Vectashield HardSet mounting medium (Vector Labs, UK). To identify
709 SST Ins Confocal image stacks of recorded neurons were taken on a Leica SP8 confocal
710 microscope using a 63x (1.4 NA) objective at 1024 × 1024 resolution (Z step size 1 μm).

711

712 *Electron microscopy:*

713 *Acute slice preparation and pharmacology for SDS-FRL*

714 Acute hippocampal slices were prepared as previously described ⁷⁶. Animals were sedated
715 with isoflurane then they were decapitated and the brains were rapidly dissected and
716 chilled in semi-frozen carbogenated (95% O_2 /5% CO_2) sucrose-substituted artificial
717 cerebrospinal fluid (sucrose-ACSF, in mM: 87 NaCl, 2.5 KCl, 25 NaHCO_3 , 1.25 NaH_2PO_4 ,
718 25 glucose, 75 sucrose, 7 MgCl_2 , 0.5 CaCl_2 , 1 Na-pyruvate, 1 Na-ascorbate). Transverse
719 hippocampal slices (200 μm thick) were cut on a vibratome (VT1200s, Leica, Germany)
720 and stored submerged in sucrose-ACSF warmed to 35°C for at least 30 min and
721 subsequently at room temperature (RT). Both solutions were equilibrated with 95% O_2 and
722 5% CO_2 gas mixture throughout experiments. Acute slices were divided into four groups
723 for pharmacological treatment: 1, incubated in ACSF (control, 30 min), 2, ACSF + GABA_BR
724 agonist baclofen (Bac, 20 μM , 20 min), 3, ACSF + protein phosphatase 2A inhibitor
725 fostriecin (FS, 50 nM, 90 min), 4, ACSF + fostriecin + baclofen (FS/Bac, FS 50 nM 90 min
726 + Bac 20 μM , 20 min). After pharmacological manipulations, slices were transferred into
727 fixative containing 1% paraformaldehyde and 15% saturated picric acid made up in 0.1 M
728 phosphate buffer (PB) overnight (O/N) at 4 °C.

729 SDS-FRL immunoelectron microscopy

730 After fixation, slices were cryoprotected in 30% glycerol in 0.1M PB O/N at 4°C ¹⁰. Blocks
731 containing stratum oriens-alveus were trimmed from the slices and frozen under high-
732 pressure (HPM 100, Leica). The frozen samples were fractured at -140°C and the
733 fractured faces were coated with carbon, (5 nm), platinum-carbon (2 nm) and carbon (18
734 nm) in a freeze-fracture replica machine (ACE 900, Leica). Replicas were digested at 80°C
735 in a solution containing 2.5% SDS and 20% sucrose diluted in 15 mM Tris buffer (TB, pH
736 8.3) for 20 hr. Subsequently, replicas were washed in washing buffer containing 0.05%

737 bovine serum albumin (BSA, Roth, Germany) and 0.1% Tween 20 in 50 mM Tris-buffered
738 saline (TBS) and then blocked in a solution containing 5% BSA and 0.1% Tween 20 in
739 TBS for 1 hr at RT. Afterwards, replicas prepared from SST-cre mice were incubated in the
740 following mixtures of primary antibodies in a solution containing 1% BSA and 0.1% Tween
741 20 made up in TBS for 3 days at 15°C: (i) GABA_{B1} (B17, rabbit, 10 µg/ml; Kulik et al.,
742 2002) and green fluorescence protein (GFP-1010, chicken 0.4 µg/ml, Aves Labs, Oregon;
743 Booker et al., 2020) or (ii) Ca_v1.2 (mouse, 50 µg/ml, NeuroMab Facility, California; Booker
744 et al., 2018) and GFP. Replicas were washed then reacted with 6 nm (GABA_{B1} and
745 Ca_v1.2) and 18 nm (GFP) gold particle-conjugated secondary antibodies (1:30, Jackson
746 ImmunoResearch Europe, Cambridgeshire) O/N at 15°C. The replicas prepared from WT
747 mice were incubated in Ca_v1.2 (mouse, 50 µg/ml, NeuroMab Facility, California), GABAB1
748 (B17, rabbit, 10 µg/ml) and metabotropic glutamate receptor 1α-subunit (mGluR1α, Guinea
749 pig, 0.4 µg/ml, Frontier Institute, Hokkaido; Booker et al., 2018) in a solution containing 1%
750 BSA and 0.1% Tween 20 made up in TBS for 3 days at 15°C. Replicas were washed then
751 reacted with 5 nm (GABA_{B1}) 10nm (Ca_v1.2) and 18 nm (mGluR1α) gold particle-
752 conjugated secondary antibodies (1:30, Jackson ImmunoResearch Europe,
753 Cambridgeshire) O/N at 15°C. Finally, replicas were washed in TBS then distilled water
754 and mounted on 100-mesh grids.

755 *Electron microscopy and quantitative analysis of protein density*

756 Labeled replicas were analyzed under an electron microscope (JEM 2100 Plus). Since all
757 antibodies target intracellular protein epitopes, immunoreactivity can be detected on the
758 plasma membrane's protoplasmic face (P-face) but not the exoplasmic face (E-face). The
759 surface density of receptors and ion channels was determined by dividing the absolute
760 number of particles labeling molecules of interest by the surface of the respective
761 segments of the dendritic shafts of GFP-positive SST-INs on replicas.

762

763 *Behavioural analysis:*

764 To investigate the role of GABA_BR activation in fear learning, we performed a fear
765 conditioning experiment. Male C57/Bl6J mice (8–12 weeks old, n = 6 per group) were
766 handled and habituated. On the day of the test, mice were given a two-minute baseline
767 period before receiving three-foot shocks (0.06mA) spaced one minute apart. Following
768 the last shock, the mice remained in the fear conditioning arena for an additional minute,
769 then they were transferred to the home cage. The entire training session lasted for approx.

770 five minutes. Mice were divided into test and control groups. Test group were injected with
771 baclofen (i.p., 2 mg/kg), while the control group received a vehicle (0.9% NaCl). One hour
772 following the baclofen or vehicle injection, freezing time was measured in the same context
773 as recall in order to access fear memory.

774 *Statistical analysis:*

775 For *ex vivo* electrophysiology, one or two cells or slices were recorded per animal per
776 treatment, limiting the ability to assess intra-animal variability. As such cells and slices
777 were treated as independent replicates and statistical analyses were conducted on these
778 data. Parametric and non-parametric tests were applied as appropriate, including two-way
779 ANOVA, Student's t-test, Mann–Whitney U-tests, and Wilcoxon signed-rank tests. For
780 SDS-FRL, where many dendritic profiles were sampled per animal replicate, we performed
781 linear-mixed effects modelling (LMM) of using the “lmer” function to limit the potential
782 inclusion of pseudoreplication. Animal was assigned as random effects, then type III
783 ANOVA (Satterthwaites method) was performed on the model to assess main effects (e.g.
784 drug treatment). If a statistical interaction between main effects was observed, then post-
785 hoc analysis was performed (Tukey post hoc tests). For all comparisons data is shown as
786 box-plots with median and 25–75% percentiles displayed in the box; minimum and
787 maximum extents are shown as whiskers, with individual data points shown from cell or
788 animal averages. For timecourses and current-voltage plots, data is shown as
789 mean \pm SEM. Statistical significance was defined as $p < 0.05$. Statistical tests and
790 graphing were performed using either GraphPad Prism (GraphPad Software v10.4.1, San
791 Diego, CA, United States) or R-Studio.

792

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