1 Research Paper

In vivo low-intensity magnetic pulses durably alter neocortical neuron excitability and spontaneous activity

6 Short title: LI-rTMS alters cortical excitability in vivo

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39 Abstract

Magnetic brain stimulation is a promising treatment for neurological and psychiatric disorders. However, 40 a better understanding of its effects at the individual neuron level is essential to improve its clinical 41 application. We combined focal low-intensity repetitive transcranial magnetic stimulation (LI-rTMS) to 42 the rat somatosensory cortex with intracellular recordings of subjacent pyramidal neurons in vivo. 43 44 Continuous 10 Hz LI-rTMS reliably evoked firing at ~4-5 Hz during the stimulation period and induced durable attenuation of synaptic activity and spontaneous firing in cortical neurons, through membrane 45 hyperpolarization and a reduced intrinsic excitability. However, inducing firing in individual neurons by 46 repeated intracellular current injection did not reproduce LI-rTMS effects on neuronal properties. These 47 data provide novel understanding of mechanisms underlying magnetic brain stimulation showing that, in 48 49 addition to inducing biochemical plasticity, even weak magnetic fields can activate neurons and enduringly modulate their excitability. 50

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54 Introduction

Repetitive transcranial magnetic stimulation (rTMS) is a promising tool for the treatment of neurological 55 and psychiatric diseases. However, a better understanding of the cellular mechanisms underlying rTMS 56 effects is needed to optimize current treatment protocols, whose outcomes remain variable (Bhattacharya 57 et al., 2021; Cirillo et al., 2017; Rossini et al., 2015). Clinical application of magnetic stimulation delivers 58 repeated high-intensity (0.5–2 Tesla, T) magnetic pulses to a targeted cortical area to induce an electric 59 field that activates cortical neurons and initiates activity-dependent plasticity (Bhattacharya et al., 2021; 60 Cirillo et al., 2017). Such plasticity is confirmed in rodents, where high-intensity magnetic stimulation 61 62 in vitro induces synaptic plasticity in hippocampal networks (Lenz and Vlachos, 2016; Vlachos et al., 63 2012) and modulates neocortical inhibitory interneurons (Hoppenrath et al., 2016; Mix et al., 2015).

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65 Besides high-field magnetic stimulation, whose duration and frequency of application is limited by technical and safety reasons (Rossini et al., 2015), new protocols using low-intensity (milliTesla, mT) 66 magnetic fields are emerging. Human studies show that low-intensity stimulation to the entire brain can 67 modulate EEG activity and cerebral function (Cook et al., 2009; Di Lazzaro et al., 2013; Rohan et al., 68 2014). In addition, focal application of these weak stimuli (low-intensity rTMS; LI-rTMS) in rodents 69 facilitates the reorganisation of abnormal circuits (Makowiecki et al., 2014; Rodger et al., 2012), 70 stimulates axonal growth and synaptogenesis (Dufor et al., 2019), and modifies cellular calcium 71 signalling and gene expression (Dufor et al., 2019; Grehl et al., 2015). However, low-intensity 72 stimulation is thought to generate electric fields weaker than those considered able to initiate action 73 potential (AP) firing (Chan and Nicholson, 1986; Grehl et al., 2016; Radman et al., 2009). Thus, it 74 remains unclear how LI-rTMS alters the properties of single neurons to contribute to structural and 75 functional changes. Moreover, human cortical responses to magnetic stimulation critically depend on the 76 ongoing activity of the stimulated brain region (Silvanto and Pascual-Leone, 2008). It is therefore 77

- necessary to identify the impact of LI-rTMS on the electrical properties of cortical neurons *in vivo*, *i.e.*
- ⁷⁹ in a physiologically relevant experimental condition that preserves neuronal connectivity.
- 80

We developed an *in vivo* model to deliver focal low-intensity (10 mT) rTMS to the somatosensory cortex 81 (S1) of sedated rodents during intracellular recording of underlying pyramidal neurons. We provide 82 direct evidence that LI-rTMS reliably elicits action potentials (APs) in S1 pyramidal cells. Also, 10 83 minutes of continuous 10 Hz LI-rTMS produces long lasting (\geq 40 minutes post-stimulation) 84 modifications in intrinsic membrane properties and spontaneous activity. These after-effects include 85 dampening of resting membrane properties, membrane hyperpolarization, and weaker background 86 synaptic and firing activity. This was accompanied by altered transfer function, characterized by 87 increased minimal current required to induce neuronal firing. These LI-rTMS-mediated effects were not 88 89 reproduced by 10 minutes of AP firing, induced by intracellular injection of threshold current pulses at 90 10 Hz. Our data show that, like high-intensity stimulation, LI-rTMS activates neurons and durably 91 modulates their activity and excitability. This changes the paradigm for brain stimulation with weak 92 magnetic fields and increases the range of potentially therapeutic stimulation parameters for clinical use.

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95 **Results**

To study the immediate and time-dependent effects of LI-rTMS on cortical activity, we paired *in vivo* intracellular recordings of S1 neurons (n=41 from 31 rats) with ECoG monitoring (Fig. 1, A–C). Some recorded neurons were filled with neurobiotin (n=19) and confirmed histologically as pyramidal neurons located in layer 5 (n=11) or 2/3 (n=8; Fig. 1B). All recorded neurons displayed the characteristic firing of S1 pyramidal cells (Mahon and Charpier, 2012; Zhu and Connors, 1999): regular spiking (n=27; Fig. 3B) or intrinsic bursts of APs (n=14) following suprathreshold current pulse injection (Fig. 3B, fig. S1). Since no differences in LI-rTMS effects were observed between pyramidal neurons located in layers 2/3
or layer 5, we pooled the corresponding results.

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105 LI-rTMS decreases spontaneous neuronal activity

First, we verified the stability of S1 cortical neuron recordings by measuring the variability of electrical 106 membrane properties for 20 minutes preceding LI-rTMS (Fig. 1C, fig. S1). During this control period, 107 the LI-rTMS coil was positioned above S1, but no stimulation was delivered. There were no significant 108 changes in neuronal spontaneous activity or membrane excitability during this baseline period (fig. S1; 109 see Supplementary Information for details). To analyze the time-dependent effects of LI-rTMS (n=16 110 111 experiments), we divided recordings into five consecutive periods: 1) pre-stimulation (Pre), the 10 minutes immediately before LI-rTMS; 2) 10 minutes LI-rTMS at 10 Hz; and 3) three successive post-112 113 stimulation times: 10-20 min (P1), 20-30 min (P2) and 40-50 min (P3) after the start of LI-rTMS (Fig. 114 1C). Before stimulation, spontaneous intracellular activity showed broadband frequency, small 115 amplitude synaptic fluctuations, and a Gaussian-like distribution of membrane potential values (Vm) around -72.1 \pm 4.8 mV (mean \pm SD; *n*=16 neurons) (Fig. 1C Pre, Fig. 2A-B). In half of the recorded 116 neurons, background synaptic activity caused spontaneous firing at 1.8 ± 2.3 Hz (n=8) with high temporal 117 variability (CV2 ISIs = 1.2 ± 0.3 , *n*=8) (Figs. 1C, 2A). Consistent with this intracellular activity, the 118 ECoG showed desynchronized, relatively fast, small-amplitude cortical waves (Fig. 1C), as classically 119 120 observed under sufentanil sedation (Altwegg-Boussac et al., 2017, 2014).

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In all tested neurons, 10 minutes continuous 10 Hz LI-rTMS induced membrane hyperpolarization compared to Pre (Fig. 2A-C), which was present immediately after LI-rTMS (P1: -2.3 \pm 2.0 mV, *n*=16 neurons; *P*<0.01) and persisted until the end of recording (P3: -2.3 \pm 1.9 mV, *n*=9 neurons; *P*<0.01) (Figs. 1Cb, 2A-C), resulting in a negative shift of the Vm distribution during the post-stimulation periods 126 (Fig. 2B left). Background synaptic activity amplitude (Vm standard deviation) was also reduced after 127 LI-rTMS in both active and silent neurons (Fig. 2A), reaching a 10.8 ± 12.2 % decrease at P3 as compared 128 to the pre-stimulation period (*P*<0.01, *n*=9 neurons) (Fig. 2A-C). This was associated with reduced power 129 of synaptic activity for frequencies below 10 Hz (Fig. 2A, B). This dampening of synaptic noise 130 amplitude was unexpected because sustained Vm hyperpolarization should increase synaptic current 131 driving force and therefore synaptic potentials, particularly given the lack of inward membrane 132 rectification in the voltage-current (*V-I*) relationships in our recorded cells (fig. S1Bb).

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Consistent with attenuated intracellular synaptic events and cell hyperpolarization, the spontaneous firing of these cortical neurons was markedly reduced after LI-rTMS. Five of the 8 neurons active prior to LIrTMS became completely silent by P2, while firing of the 3 remaining cells decreased to 0.07 ± 0.13 Hz at P3 (Fig. 2A-C). This failure of cell firing was not due to altered ionic AP mechanisms, as their amplitude, half-width duration and threshold voltage did not change (Fig. 2B right, table S1); it more likely reflected the decreased background synaptic drive.

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141 LI-rTMS alters neuronal integrative properties and input-output relationship

We next tested if LI-rTMS altered the excitability of S1 pyramidal neurons by changing their integrative properties, *i.e.* their ability to process subthreshold membrane currents, and their input-output relationship before and after stimulation.

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We first examined changes in membrane input resistance (Rm) and membrane time constant (τ m), two parameters that govern, respectively, the ability of synaptic currents to change Vm and the efficacy with which a neuron summates synaptic potentials over time. Rm values, extrapolated from the slope of the *V-I* curves (fig. S1B), were already significantly reduced at P1 (compared to Pre: -14.1 ± 6.6 %, *n* =11 neurons; P < 0.001), a decrease that persisted for up to 40 min after the end of stimulation (n=9 neurons; P < 0.01) (Fig. 3A, C). The decrease in Rm was not caused by LI-rTMS-mediated changes in the morphology of recorded neurons, whose anatomical properties were comparable to those classically described for S1 pyramidal cells (Brecht et al., 2003; Mahon and Charpier, 2012; Manns et al., 2004) (Fig. 1B). τ m values also showed a persistent decrease that closely matched the reduction in Rm (Fig. 3C). The attenuation of Rm and τ m by LI-rTMS may contribute to the smaller amplitude of spontaneous synaptic potentials and their ineffective summation to elicit APs.

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158 As changes in Vm and Rm can affect the reactivity of a neuron to its inputs (Altwegg-Boussac et al., 159 2014; Silver, 2010), we looked for possible LI-rTMS-induced modifications in the input-output 160 relationship. We injected positive current pulses of increasing intensity during the different recording 161 periods and counted the number of evoked APs, generating frequency-current (F-I) curves (fig S1Cb, 162 Fig. 3B). After LI-rTMS, F-I curves progressively shifted to the right, leading to a ~50% increase of the 163 threshold current values by P3 (n=7 neurons; compared to Pre, P<0.05) (Fig. 3B, C). This reduced 164 neuronal responsiveness to small excitatory inputs was accompanied by relatively stable neuronal gain, defined as the slope of the linear portion of F-I curves, throughout the post-stimulation period (P = 0.23) 165 (Fig. 3C). 166

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168 Low-intensity magnetic pulses trigger APs

To identify the mechanisms initiating the long-lasting inhibitory effects of LI-rTMS on cortical activity and integrative properties, we investigated the immediate effects of LI-rTMS during the stimulation period. A practical difficulty arises from the brief (~1.6 ms) large-amplitude artifacts generated by each LI-rTMS pulse (Figs. 1, 4, fig. S2), which obscures the early part of the neuronal response. However, we noticed in all recorded neurons (n=16) that the terminal part of many artifacts was rapidly followed by a biphasic voltage waveform resembling an intracellular AP (Fig. 4A, fig. S2A). We thus removed the 175 stimulation artifacts off-line from traces displaying an apparent AP waveform (see Methods, and Figs. 4, 5 and fig. S2 for details). The artifact-free recordings revealed intracellular waveforms whose shape, 176 peak amplitude and kinetics matched those of spontaneous APs recorded from the same neurons (Fig. 177 4Ab). The mean latency of AP-like events, calculated from the end of the stimulus to the peak of the 178 waveform, was 0.6 ± 0.1 ms (n=16 neurons) and the percentage of individual magnetic pulses 179 180 immediately preceding these waveforms was 24 ± 31 % (*n*=16 neurons; Fig 4B, bottom left), with a high cell-to-cell variability (from 0.3 to 99.8 %). Given that magnetic intensity decreases by distance-squared, 181 182 we tested whether this inter-neuronal variability depended on the depth of recorded cells; there was no correlation ($r^2 = 0.03$). Notably, these AP-like events are not non-biological artifacts resulting from direct 183 action of the magnetic field on the recording electrode, as they did not occur in control experiments in 184 185 which a model cell was used to simulate the recording of a pyramidal neuron (fig. S2B).

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To confirm that the high-voltage waveforms were indeed APs, we tested their sensitivity to sustained 187 Vm hyperpolarization - which prevents firing by moving the neuron's Vm further from the AP threshold 188 - or to the blockade of voltage-gated sodium channels by the lidocaine analog OX-314. In the absence of 189 LI-rTMS, spontaneous firing frequency which was 1.8 ± 1.6 Hz at rest (Vm = -71.9 \pm 7.0 mV, *n*=6 tested 190 neurons), declined with imposed DC hyperpolarization (from -0.1 to -2 nA), and fell to zero when Vm 191 reached -81.2 \pm 8.1 mV (*n*=6 neurons) (Fig. 4B, top). Vm hyperpolarization was associated with 192 increased amplitude of synaptic fluctuations, as expected from a greater driving force of excitatory 193 synaptic currents (Fig. 4B, top right). Hyperpolarizing these same cortical neurons during short periods 194 of LI-rTMS similarly decreased the rate of the AP-like events, from 4.0 ± 2.7 Hz at rest (*n*=6 neurons) 195 to a residual rate of 0.04 \pm 0.05 Hz at the most negative membrane polarization tested (Vm = -81.2 \pm 8.1 196 mV, *n*=6 neurons) (Fig. 4B, bottom). 197

199 Second, to test whether the putative LI-rTMS-induced APs had the same ionic mechanisms as spontaneous APs, we perfused the intracellularly recorded neurons with QX-314 (n=9) and compared 200 the drug-related changes in amplitude of both types of AP. In each neuron, measurements were made 201 during alternating control and stimulation periods of 5–160 seconds. The peak potential of spontaneous 202 APs was progressively reduced (P < 0.05) by the QX314 (Fig. 4C) and this time-dependent reduction was 203 accompanied by a mean Vm depolarization of 5–10 mV, as classically observed in QX-314-filled cortical 204 neurons (Benardo et al., 1982; Connors and Prince, 1982). The decrease in peak values of LI-rTMS-205 induced waveforms and spontaneous APs was almost identical for any given time after the onset of QX-206 207 314 diffusion (P>0.2 for each recording time), reaching ~ -14–20 mV in the neurons that could be 208 recorded until 30–40 min after cell impalement (n=5) (Fig. 4C). Because the QX314-reduced APs were 209 occasionally difficult to distinguish from spontaneous depolarized synaptic potentials in recordings of 210 long duration (Fig. 4Ca, arrows), we also examined the effects of QX-314 on APs evoked by depolarizing 211 current steps (0.2–0.4 nA, n=9 neurons) (Fig. 4Cb, Cc). Again, the time-course of the reduction in 212 current-induced AP peak potentials was similar to that measured for LI-rTMS-induced APs (P>0.1 for 213 each recording time) (Fig. 4Cb, Cc).

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215 The LI-rTMS-induced APs occurred in all recorded cortical neurons, regardless of their firing frequency prior to stimulation (Fig. 5A-C). This led to a significant increase in the overall firing rate (including 216 both spontaneous and LI-rTMS-evoked APs) throughout the LI-rTMS period (n=16 neurons; P<0.05) 217 (Fig. 5B). However, the rate of spontaneous APs, when measured separately, was decreased by almost 218 45% during LI-rTMS (Pre: 1.8 ± 2.3 Hz, n=8 active neurons, LI-rTMS: 0.6 ± 0.8 Hz, n=8 neurons; $P < 1.8 \pm 2.3$ H 219 0.01) (Fig. 5B). The induced APs did not result from a LI-rTMS-mediated increase in synaptic activity 220 221 or sustained membrane depolarization since artifact-free records did not show significant change in the amplitude of Vm fluctuations in association with the firing induced by magnetic pulses (Figs. 4, 5, and 222 fig. S2A). However, their short latency and ability to be evoked from relatively hyperpolarized membrane 223

potentials indicate that LI-rTMS-evoked APs probably resulted from activation of an axonal site located
 close to the soma.

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These results demonstrate for the first time that 10 Hz LI-rTMS can elicit AP firing in pyramidal neurons, with kinetic and ionic mechanisms similar to spontaneous APs, despite inducing an electric field estimated to be at least 100 times lower than that used in high-intensity rTMS. They also demonstrate that 10 minutes continuous 10 Hz LI-rTMS produces robust and long-lasting inhibitory effects on the activity and integrative properties of individual pyramidal neurons.

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233 Action potentials evoked by intracellular current injection do not reproduce LI-rTMS effects

234 In order to determine whether the LI-rTMS-induced firing was responsible for the changes in cortical 235 neuron activity and responsiveness, we examined, in another group of S1 neurons, the impact of an AP discharge evoked by intracellular injection of current steps. Single cortical neurons (n=7) were 236 237 challenged with repetitive depolarizing current pulse stimulation (rCPS) delivered intracellularly at10 238 Hz for 10 min (Fig. 6A). Intensity of current steps was adjusted in each neuron to obtain a proportion of suprathreshold stimulations (~30%) and an overall firing frequency comparable to those produced by LI-239 rTMS (5.9 \pm 4.6 Hz, *n*=7 neurons, including both spontaneous and evoked APs; *P*>0.4) (Fig. 6B). The 240 effects of rCPS were compared to a subgroup of magnetically stimulated neurons (n=9) with similar 241 baseline membrane electrical properties, including mean Vm, spontaneous firing frequency and Rm 242 (n=9; P>0.1 for each parameter). 243

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The first obvious difference between effects of current-evoked APs (rCPS) and LI-rTMS was that rCPS did not produce neuronal hyperpolarization during the post-stimulation period (Fig. 6C, left and fig. S3D). This was accompanied by stable Vm fluctuations and spontaneous firing frequency (Pre *versus* post-rCPS: *P*>0.2 for each parameter) (fig. S3A, D). Furthermore, comparison of *V-I* curves before and

after rCPS showed no significant modification of Rm (n=6 neurons; Pre, 17.2 ± 6.9 M Ω vs. Post, $17.9 \pm$ 249 7.5 MΩ; P = 0.14) or τm (*n*=6 neurons; Pre, 6.5 ± 1.8 ms, vs. Post, 6.3 ± 1.6 ms; P = 0.28) (Fig. 6C, 250 middle and fig. S3B, D). Finally, analysis of F-I curves revealed heterogeneous modification in the 251 sensitivity of cortical neurons to weak depolarizing inputs. Although average current threshold values 252 did not significantly change (P=0.4), most neurons (n=4 out of 6 tested cells) had a significantly 253 254 decreased firing threshold ($\Delta I_{th} = -0.14 \pm 0.07$ nA, n=4; P<0.05) following rCPS. This modulation of cortical responsiveness was not accompanied by modification in the neuronal gain (P>0.4) (Fig. 6C right 255 256 and fig. S3C, D). Thus, the firing of single cortical neurons at a similar frequency to that induced by magnetic stimulation (~ 5 Hz) is not sufficient to reproduce, on its own, the LI-rTMS-related changes in 257 neuronal excitability. 258

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260 Discussion

To our knowledge, this study is the first to investigate the immediate (online) and after- (offline) effects 261 of low-intensity rTMS on the electrical properties of single neurons in vivo. Because rTMS-induced 262 cortical responses depend on the ongoing brain state (Gersner et al., 2011; Silvanto and Pascual-Leone, 263 264 2008), we used experimental conditions which retain stable background cortical activity, similar to the waking state, during the long intracellular recordings (Altwegg-Boussac et al., 2014; Bruno and 265 Sakmann, 2006; Constantinople and Bruno, 2011). Our results indicate that 10 minutes continuous 10 266 267 Hz LI-rTMS alters the passive membrane properties of S1 pyramidal neurons, reduces their intrinsic excitability and attenuates their spontaneous firing for at least 40 min. Moreover, low-intensity magnetic 268 pulses triggered APs in all recorded neurons. These findings deepen our knowledge on the cellular 269 270 mechanisms underlying LI-rTMS-induced plasticity and may guide the development of new stimulation patterns for clinical treatment. 271

273 LI-rTMS induces action potentials in stimulated neurons

A major and unexpected finding of our study was that low intensity (10 mT) magnetic pulses elicited 274 APs in all recorded neurons (25% of pulses on average). This challenges the common assumption that 275 LI-rTMS only acts through subthreshold mechanisms because it induces electrical fields weaker than 276 those required for neuronal activation (Chan and Nicholson, 1986; Grehl et al., 2016; Radman et al., 277 2009). Our result is however coherent with weak electric fields inducing short-latency spiking in retinal 278 ganglion cells in vitro (Bonmassar et al., 2012) and medium-intensity (60 mT) magnetic pulses 279 transiently activating sodium currents to facilitate AP firing in S1 pyramidal neurons (Banerjee et al., 280 281 2017) in slices. The percentage of magnetic pulses initiating APs varied between cells, consistent with 282 variable pyramidal neuron sensitivity to magnetic and electric fields, depending on their location, morphology and intrinsic excitability (Radman et al., 2009; Rotem and Moses, 2008) as well as 283 284 fluctuations in their spontaneous firing (Pasley et al., 2009).

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The mechanisms underlying magnetic pulse-induced neuronal activation remain unclear. In highintensity TMS, trans-synaptic or direct neuronal depolarization are thought to initiate APs in the axon (Müller-Dahlhaus and Vlachos, 2013; Pashut et al., 2014; and references therein). Here, LI-rTMSinduced APs had short latency, were not associated with increased subthreshold synaptic events, and could be triggered from a hyperpolarized Vm, suggesting direct activation of the axon initial segment, whose high concentration of voltage-gated sodium channels confers a lower AP threshold (Eccles, 2013; Kole et al., 2008).

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294 Continuous 10 Hz LI-rTMS durably decreases cortical excitability and activity

295 The main after-effects of LI-rTMS included smaller amplitude background synaptic noise, substantial

296 membrane hyperpolarization and marked reduction or cessation of spontaneous firing. These changes in

297 spontaneous activity were accompanied by attenuated neuronal integrative properties (correlated 298 decrease in Rm and τ m) and reduced intrinsic excitability (higher minimal current firing threshold).

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The likely primary origin of these effects, whether altered synaptic events or intrinsic cellular properties, 300 can be deduced from our data. Long-term depression of synaptic inputs onto the recorded neurons, which 301 could explain the observed reduction in synaptic activity, would decrease global membrane conductance 302 (due to smaller excitatory synaptic conductance) and thus increase Rm and Tm values (Destexhe et al., 303 2003) – the opposite of our findings. Rather, we suggest that LI-rTMS-mediated inhibition of neuronal 304 activity originates from alteration to cortical neuron integrative properties. Indeed, the initial reduction 305 in Rm and τm , which attenuates neuronal responsiveness to their synaptic inputs, would explain both the 306 307 reduced amplitude of Vm fluctuations and membrane hyperpolarization. Given the inverse correlation linking current firing threshold to Vm in cortical neurons (Altwegg-Boussac et al., 2014; Carton-Leclercq 308 309 et al., 2021), the membrane hyperpolarization would in turn explain the rightward shift of input-output 310 relations and the reduced intrinsic excitability. Finally, all these cellular modifications will combine to 311 reduce the efficiency with which cortical neurons process synaptic currents and generate APs and, thus, 312 explain the progressive decrease of spontaneous firing. This reduced firing in individual neurons will contribute to global reduction of synaptic activity flow within the stimulated cortical network, which will 313 further reduce synaptic and firing activity. Furthermore, increased activity of neocortical GABAergic 314 315 interneurons could also contribute to reducing pyramidal cell activity, by producing a robust shunting effect on pyramidal neuronal membrane (Tremblay et al., 2016) and lowering Rm (due to larger 316 inhibitory synaptic conductance). 317

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The precise subcellular mechanisms through which LI-rTMS exerts its effects warrant further investigation. We hypothesize that increased neuronal firing during stimulation opens voltage-gated 321 calcium channels, increasing intracellular calcium and activating calcium-dependent enzymatic cascades that regulate ionic conductances (Delord et al., 2007; Xu and Kang, 2005). In line with this hypothesis, 322 pyramidal neuron firing induced by medium-intensity (60 mT) rMS in vitro is associated with increased 323 cytosolic calcium (Banerjee et al., 2017). Moreover, our data showing changes in Rm and current firing 324 threshold together with stable neuronal gain after LI-rTMS suggest that the long-lasting decrease in 325 neuronal excitability is due to modulation of potassium currents activated around resting Vm (Naudé et 326 al., 2012). This suggestion is supported by previous reports, in which similar depression of intrinsic cell 327 excitability was attributed to calcium-dependent upregulation of slowly-inactivating potassium currents 328 329 (Li et al., 2004; Lombardo et al., 2018).

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Our results also indicate that induced APs, and their associated changes in calcium concentration, may 331 332 not be sufficient to trigger the time-dependent post-LI-rTMS effects. Simply replicating the level of LIrTMS-induced AP firing by repeated intracellular injection of current pulses in single neurons did not 333 significantly modify resting and active membrane properties. This suggests that electric field-induction 334 is only one of several mechanisms activated by electromagnetic pulses. We have previously shown that 335 cryptochrome magnetoreceptors are required for neuroplasticity effects of LI-rTMS (Dufor et al., 2019) 336 and modulate reactive oxygen species (ROS) (Sherrard et al., 2018). In cell redox physiology, ROS 337 activate ryanodine receptors to release calcium from intracellular stores (Görlach et al., 2015), a 338 phenomenon we have demonstrated in cortical neurons in response to LI-rTMS (Grehl et al., 2015). It 339 is therefore likely that both voltage-gated calcium channel activation and calcium release from 340 intracellular stores are required for the long-term modulation of pyramidal neuron activity. 341

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343 Relevance of LI-rTMS to human rTMS treatment

The LI-rTMS-induced inhibitory effects on neuronal excitability and firing is surprising in light of former studies using high-intensity rTMS. According to these reports, high-frequency magnetic stimulation (\geq

5 Hz) should increase cortical excitability (Bhattacharya et al., 2021; Cirillo et al., 2017; Huang et al.,
2005), potentiate excitatory synaptic transmission and/or upregulate the expression of neurochemical
markers of synaptic plasticity (Cirillo et al., 2017; Gersner et al., 2011; Lenz et al., 2015; Vlachos et al.,
2012).

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However, there are important differences between those reports and our study, which can explain the 351 different outcomes. First, in some of these studies, rTMS effects on synaptic function were examined ex 352 vivo after repeated daily sessions of high-frequency rMS (Cirillo et al., 2017; Gersner et al., 2011), which 353 354 is not the case here. Second, in studies where acute effects of magnetic stimulation were measured, rMS 355 was usually delivered directly to a brain slice or organotypic culture and not to an intact brain as in our in vivo experimental condition (Lenz et al., 2015; Tang et al., 2016; Vlachos et al., 2012). For example, 356 357 in mouse S1 slices, 3 minutes of medium-intensity (85 mT) intermittent theta-burst magnetic stimulation reduced AP voltage threshold in pyramidal neurons and increased their current-evoked firing, without 358 359 changing resting membrane properties (Tang et al., 2016). Discordance between results obtained in vitro 360 and in living brains has already been reported: opposite changes in synaptic efficacy can be induced in vivo and in vitro at the same synapses by the same stimulation paradigm (Mahon et al., 2004). Thus 361 differences between previous observations and our current in vivo findings reinforce that rTMS outcomes 362 depend on ongoing activity in the targeted neural networks during the stimulation (Gersner et al., 2011; 363 Silvanto and Pascual-Leone, 2008). Finally, most previous studies used magnetic fields at least ten times 364 stronger than the one used in our study. High-intensity, high-frequency stimulation protocols require 365 regular pauses to allow the coil to cool - resulting in short periods (a few seconds) of stimulation 366 separated by pauses without stimulation (see Bhattacharya et al., 2021; Cirillo et al., 2017 for reviews), 367 an intermittent pattern in contrast to our continuous 10 Hz. Theta-burst stimulation, for example, induces 368 opposite effects if applied continuously versus intermittently (Huang et al., 2005). Thus, LI-rTMS effects 369

370 on cell excitability are likely to depend on stimulation parameters, similar to the pattern-dependence of

LI-rTMS effects on structural and biochemical plasticity (Dufor et al., 2019; Grehl et al., 2015).

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Although the functional consequences of LI-rTMS-mediated depression of cortical excitability for S1 network function and behavior remain unknown, our LI-rTMS protocol could be useful in epileptic disorders, which are often associated with cortical hyperexcitability and paroxysmal network synchronization (Kimiskidis et al., 2014). An understanding of the immediate and lasting effects on neuronal activity, measured in real time, is an important advance for the development and evaluation of stimulation patterns in future clinical treatments.

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In summary, our findings that even weak magnetic fields can activate neurons and modulate their excitability, change our understanding of low-intensity magnetic brain stimulation. We show that lowintensity approaches have some underlying processes in common with already established high-intensity clinical treatments, but without the constraints of special high-voltage equipment. This knowledge not only expands the range of stimulation protocols (pulse frequency and patterns), and therefore future potential clinical treatments, but also renders LI-rTMS treatments available beyond the hospital setting.

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388 Materials and Methods

389 Animals, their preparation and surgery

All animal experiments received approval (APAFIS no. 18003-2019051019017280) from the *Charles Darwin* Animal Experimentation Ethics Committee (C2EA-05), in accordance with the guidelines of the European Union (directive 2010/603/EU). Experiments were conducted *in vivo* on adult Sprague Dawley rats (n = 31) of either sex (Charles River Laboratories, France). Under isoflurane (3.5%; Centravet, France) anesthesia, animals underwent tracheotomy for artificial ventilation (room air, 85 breaths.min⁻¹, 395 2.6 ml.cycle⁻¹) under neuromuscular blockade (gallamine triethiodide, 40 mg/2h i.m; Sigma-Aldrich, France). They were placed in a stereotaxic frame and small craniotomies were made above the left barrel 396 field of S1 (1.2–1.3 mm posterior to bregma, 4.1–6.5 mm lateral to midline; Paxinos and Watson, 1986) 397 to allow simultaneous recording of ECoG and intracellular activity of subjacent pyramidal neurons. After 398 craniotomy, isoflurane was discontinued, but sedation and analgesia were maintained by repeated 399 injection of sufentanil (3 µg/kg every 30 min, i.p.; Piramal) and local infiltration of lidocaine (2%, 400 Centravet) in incision and pressure points. Sedation with sufentanil (Niemegeers et al., 1976) was chosen 401 because it induces a globally stable background ECoG activity characterized by low amplitude 402 403 desynchronized cortical waves resembling those encountered during the waking state (Altwegg-Boussac 404 et al., 2017, 2014; Bruno and Sakmann, 2006; Constantinople and Bruno, 2011; Depaulis et al., 2016). 405 It thus avoids anesthesia-related modulation of cortical responses to magnetic stimulation, as observed 406 in both animal and human studies (Gersner et al., 2011; Silvanto and Pascual-Leone, 2008). Physiological 407 parameters were continuously monitored during recordings (Altwegg-Boussac et al., 2017; Schramm et 408 al., 2020). Each animal formed its own control, with LI-rTMS effects compared to baseline measurements obtained before the magnetic stimulation. After the recordings, animals were euthanized 409 with euthasol (40%, i.p.; Centravet) for histological examination (see below). 410

411

412 In vivo multi-scale electrophysiological recordings

Intracellular recordings were performed with glass micropipettes filled with 2-M potassium acetate (50-90 MΩ). Current-clamp recordings were amplified using an Axoclamp 900A amplifier (Molecular Devices, Union City, CA, USA) operating in bridge mode, filtered between *DC* and 30 kHz and digitized at 10 kHz (CED 1401plus, Spike2 software version 7.06; Cambridge Electronic Design). Intracellularly recorded neurons (n = 41), identified as pyramidal cells by their morphological and electrophysiological characteristics (Steriade, 2004), were located in layers 2/3 and 5 at depths ranging between 800–3300 µm below the cortical surface (Paxinos and Watson, 1986; Wilent and Contreras, 2004). In some experiments (n = 8), we added to the pipette solution 50–100 mM of QX-314 (Tocris Biosciences), a voltage-gated sodium channels blocker known to suppress APs in a time-dependent manner (Connors and Prince, 1982; Wilson and Kawaguchi, 1996).

Spontaneous ECoG activity was recorded with a low-impedance ($\sim 60 \text{ k}\Omega$) silver electrode placed on the dura above S1 and a reference electrode placed on a contralateral temporal muscle. Surface cortical signals were amplified using a differential *AC* amplifier (Model 1700; A-M Systems), filtered at 1 Hz– 1 kHz, and digitized at 3 kHz.

427

428 Stimulation protocols

Continuous 10Hz LI-rTMS was applied for 10 minutes (6000 pulses) using a miniaturized round coil 429 connected to an electromagnetic pulse generator (E-Lab5, Global Energy Medicine, Perth, Australia), 430 which generated 300-us monophasic pulses of 10mT intensity, as previously described (Makowiecki et 431 al., 2014; Rodger et al., 2012). The coil was secured to a micromanipulator and placed above S1 (Fig. 432 1A). Brain-to-coil distance was minimized (~2 mm) while assuring contact-free stimulation 433 perpendicular to the cortical surface. We verified that LI-rTMS caused no electromagnetic interference 434 in the recording leads and electrodes, with negative control experiments in which a model cell (CLAMP-435 1U Model Cell, Molecular Devices, Union City, USA) in 'cell mode' was attached to the headstage 436 amplifier (fig. S2B). 437

Repetitive current pulse stimulation consisted of intracellular injections of depolarizing square current
pulses (40 ms of duration) at 10 Hz for 10 min. The intensity of the current pulses (0.1–0.5 nA) was
adjusted for each neuron to evoke an AP every 3–5 current pulses, mimicking the frequency of LI-rTMSinduced APs.

442

443 Intracellular and ECoG signal analysis

Spontaneous activity, membrane excitability and integrative properties of cortical neurons were assessed 444 before and after LI-rTMS. Average membrane potential (Vm) was calculated as the mean of the 445 distribution of spontaneous subthreshold membrane potential values and the magnitude of Vm 446 fluctuations was quantified as the standard deviation (σ) of the distribution (σ Vm). Time-frequency maps 447 illustrating the spectral content of synaptic activity over time were constructed using a multitaper 448 decomposition analysis with a sliding time window of 3 seconds (Oostenveld et al., 2011), after removing 449 APs which were replaced by a piecewise cubic hermite polynomial interpolation. Power spectra were 450 computed by averaging the time-frequency decomposition over a 10-minute period. 451

Mean spontaneous firing rate and Vm values were measured from recordings of at least 1 min. The regularity of neuronal firing was calculated by the CV2 method: here $CV2 = (2|\Delta t_{i+1} - \Delta t_i|)/(\Delta t_{i+1} + \Delta t_i)$, and $\Delta t_i = t_i - t_{i-1}$ is the inter-spike-interval (Altwegg-Boussac et al., 2014). AP properties were measured from spontaneously occurring APs. AP voltage threshold was defined as the Vm at which dVm.dt⁻¹ first exceeds 10 mV.ms⁻¹ (Mahon and Charpier, 2012). AP amplitude was calculated from threshold to peak potential and AP half-width was the duration of the waveform at half-amplitude.

458 Dynamic changes of Vm, σ Vm and spontaneous firing frequency were assessed over successive windows 459 of 30 seconds. Recording periods during which cortical excitability was assessed (see below) were 460 removed from this analysis and replaced by a linear interpolation.

Rm was calculated as the slope of the linear portion of voltage-current (*V-I*) plots generated following repeated injections of hyperpolarizing current of increasing intensity (200 ms at 2 Hz, 10–20 trials per intensity, -0.2 to -0.8 nA) or from the average voltage deflection induced by hyperpolarizing current pulses of -0.4 nA (200 ms, \geq 10 trials). τ m was derived from an exponential decay fit applied to the initial phase of the mean current-induced hyperpolarization. Neuronal transfer function (*F-I* relationship), which describes the cell output firing as a function of injected depolarizing current pulses of increasing intensity (200 ms at 2 Hz, 10–20 trials per intensity, 0.1–1.5 nA), was quantified before and after LI- 468 rTMS. Linear regression of *F*–*I* curves yielded the threshold current for AP generation (I_{th} ; *x*-intercept) 469 and neuronal gain (γ; slope) (Altwegg-Boussac et al., 2014; Mahon and Charpier, 2012).

470

471 LI-rTMS artifact removal

To assess changes in cortical activity during the 10 minutes of LI-rTMS, we removed brief highamplitude electrical artifacts induced by the magnetic pulses. We removed the recorded signal for 1.6 ms following each pulse onset and preserved continuity by linear interpolation (fig. S2), as described for human stimulation experiments (Amengual et al., 2017; Stengel et al., 2021). We validated this approach, by adding a new channel containing artificially generated stimulation events at 10 Hz for 10 minutes to control recordings. These false artifacts were then removed with the same procedure and the resulting electrophysiological signal compared to the original to ensure their similarity (Amengual et al., 2017).

479

480 Neuronal labelling

Neurons were filled with neurobiotin (1.5% in the pipette solution; Vector Laboratories, Burlingame, USA) through repeated depolarizing current pulse injections over 10 minutes after the end of the poststimulation period, as previously described (Mahon and Charpier, 2012). Rats were euthanized with euthasol and transcardially perfused with 0.3% glutaraldehyde-4% paraformaldehyde in PBS (0.1 M, pH 7.4; VWR Chemicals, France), and the neurobiotin-filled neurons revealed as previously described (Williams et al., 2016).

487

488 Statistical analysis

489 Data were explored for normality, outliers and statistical test assumptions in Prism version 9.00 490 (GraphPad Inc., San Diego, USA). Group differences were assessed using paired two-tailed Student's t-491 test, one-way Repeated Measures ANOVA, or non-parametric Wilcoxon signed rank test and the Mann-492 Whitney rank sum test. The Benjamini-Hochberg test was used to correct for multiple comparisons

493 (Benjamini and Hochberg, 1995). Values are expressed as means \pm SD and significance level was P <

494 0.05.

495

497 **References**

- Altwegg-Boussac T, Chavez M, Mahon S, Charpier S. 2014. Excitability and responsiveness of rat barrel
 cortex neurons in the presence and absence of spontaneous synaptic activity in vivo. *J Physiol* 500 592:3577–3595. doi:10.1113/jphysiol.2013.270561
- Altwegg-Boussac T, Schramm AE, Ballestero J, Grosselin F, Chavez M, Lecas S, Baulac M, Naccache 501 L, Demeret S, Navarro V, Mahon S, Charpier S. 2017. Cortical neurons and networks are dormant 502 but fully responsive during isoelectric brain state. Brain **140**:2381–2398. 503 doi:10.1093/brain/awx175 504
- Amengual JL, Vernet M, Adam C, Valero-Cabré A. 2017. Local entrainment of oscillatory activity
 induced by direct brain stimulation in humans. *Sci Rep* 7:41908. doi:10.1038/srep41908
- Banerjee J, Sorrell ME, Celnik PA, Pell G. 2017. Immediate Effects of Repetitive Magnetic Stimulation
 on Single Cortical Pyramidal Neurons. *PloS One* 12:e0170528. doi:10.1371/journal.pone.0170528
- Benardo LS, Masukawa LM, Prince DA. 1982. Electrophysiology of isolated hippocampal pyramidal
 dendrites. *J Neurosci* 2:1614–1622.
- Benjamini Y, Hochberg Y. 1995. Controlling the False Discovery Rate: A Practical and Powerful
 Approach to Multiple Testing. *J R Stat Soc Ser B Methodol* 57:289–300.
- Bhattacharya A, Mrudula K, Sreepada SS, Sathyaprabha TN, Pal PK, Chen R, Udupa K. 2021. An
 Overview of Noninvasive Brain Stimulation: Basic Principles and Clinical Applications. *Can J Neurol Sci* 48:1–14. doi:10.1017/cjn.2021.158
- Bonmassar G, Lee SW, Freeman DK, Polasek M, Fried SI, Gale JT. 2012. Microscopic magnetic
 stimulation of neural tissue. *Nat Commun* 3:921. doi:10.1038/ncomms1914
- Brecht M, Roth A, Sakmann B. 2003. Dynamic receptive fields of reconstructed pyramidal cells in layers 518 Physiol 519 3 and 2 of rat somatosensory barrel cortex. J**553**:243–265. doi:10.1113/jphysiol.2003.044222 520
- 521 Bruno RM, Sakmann B. 2006. Cortex is driven by weak but synchronously active thalamocortical 522 synapses. *Science* **312**:1622–1627. doi:10.1126/science.1124593
- 523 Carton-Leclercq A, Lecas S, Chavez M, Charpier S, Mahon S. 2021. Neuronal excitability and sensory
 524 responsiveness in the thalamo-cortical network in a novel rat model of isoelectric brain state. J
 525 Physiol 599:609–629. doi:10.1113/JP280266
- 526 Chan CY, Nicholson C. 1986. Modulation by applied electric fields of Purkinje and stellate cell activity
 527 in the isolated turtle cerebellum. *J Physiol* 371:89–114.
- Cirillo G, Di Pino G, Capone F, Ranieri F, Florio L, Todisco V, Tedeschi G, Funke K, Di Lazzaro V.
 2017. Neurobiological after-effects of non-invasive brain stimulation. *Brain Stimulat* 10:1–18.
 doi:10.1016/j.brs.2016.11.009
- Connors BW, Prince DA. 1982. Effects of local anesthetic QX-314 on the membrane properties of
 hippocampal pyramidal neurons. *J Pharmacol Exp Ther* 220:476–481.
- Constantinople CM, Bruno RM. 2011. Effects and mechanisms of wakefulness on local cortical
 networks. *Neuron* 69:1061–1068. doi:10.1016/j.neuron.2011.02.040
- Cook CM, Saucier DM, Thomas AW, Prato FS. 2009. Changes in human EEG alpha activity following
 exposure to two different pulsed magnetic field sequences. *Bioelectromagnetics* 30:9–20.
 doi:10.1002/bem.20434
- Delord B, Berry H, Guigon E, Genet S. 2007. A new principle for information storage in an enzymatic
 pathway model. *PLoS Comput Biol* 3:e124. doi:10.1371/journal.pcbi.0030124
- Depaulis A, David O, Charpier S. 2016. The genetic absence epilepsy rat from Strasbourg as a model to
 decipher the neuronal and network mechanisms of generalized idiopathic epilepsies. *J Neurosci Methods* 260:159–174. doi:10.1016/j.jneumeth.2015.05.022
- 543 Destexhe A, Rudolph M, Paré D. 2003. The high-conductance state of neocortical neurons in vivo. *Nat* 544 *Rev Neurosci* 4:739–751. doi:10.1038/nrn1198
- 545 Di Lazzaro V, Capone F, Apollonio F, Borea PA, Cadossi R, Fassina L, Grassi C, Liberti M, Paffi A,

- Parazzini M, Varani K, Ravazzani P. 2013. A consensus panel review of central nervous system
 effects of the exposure to low-intensity extremely low-frequency magnetic fields. *Brain Stimulat*6:469–476. doi:10.1016/j.brs.2013.01.004
- 549 Dufor T, Grehl S, Tang AD, Doulazmi M, Traoré M, Debray N, Dubacq C, Deng Z-D, Mariani J, Lohof
 550 AM, Sherrard RM. 2019. Neural circuit repair by low-intensity magnetic stimulation requires
 551 cellular magnetoreceptors and specific stimulation patterns. *Sci Adv* 5:eaav9847.
 552 doi:10.1126/sciadv.aav9847
- 553 Eccles JC. 2013. The Physiology of Synapses. Academic Press.
- Gersner R, Kravetz E, Feil J, Pell G, Zangen A. 2011. Long-Term Effects of Repetitive Transcranial
 Magnetic Stimulation on Markers for Neuroplasticity: Differential Outcomes in Anesthetized and
 Awake Animals. *J Neurosci* 31:7521. doi:10.1523/JNEUROSCI.6751-10.2011
- Görlach A, Bertram K, Hudecova S, Krizanova O. 2015. Calcium and ROS: A mutual interplay. *Redox Biol* 6:260–271. doi:10.1016/j.redox.2015.08.010
- Grehl S, Martina D, Goyenvalle C, Deng Z-D, Rodger J, Sherrard RM. 2016. In vitro Magnetic
 Stimulation: A Simple Stimulation Device to Deliver Defined Low Intensity Electromagnetic
 Fields. Front Neural Circuits 10:85. doi:10.3389/fncir.2016.00085
- Grehl S, Viola HM, Fuller-Carter PI, Carter KW, Dunlop SA, Hool LC, Sherrard RM, Rodger J. 2015.
 Cellular and molecular changes to cortical neurons following low intensity repetitive magnetic
 stimulation at different frequencies. *Brain Stimulat* 8:114–123. doi:10.1016/j.brs.2014.09.012
- Hoppenrath K, Härtig W, Funke K. 2016. Intermittent Theta-Burst Transcranial Magnetic Stimulation
 Alters Electrical Properties of Fast-Spiking Neocortical Interneurons in an Age-Dependent
 Fashion. *Front Neural Circuits.* 10:22. doi:10.3389/fncir.2016.00022
- Huang Y-Z, Edwards MJ, Rounis E, Bhatia KP, Rothwell JC. 2005. Theta burst stimulation of the human
 motor cortex. *Neuron* 45:201–206. doi:10.1016/j.neuron.2004.12.033
- Kimiskidis VK, Valentin A, Kälviäinen R. 2014. Transcranial magnetic stimulation for the diagnosis and
 treatment of epilepsy. *Curr Opin Neurol* 27:236–241. doi:10.1097/WCO.000000000000071
- Kole MHP, Ilschner SU, Kampa BM, Williams SR, Ruben PC, Stuart GJ. 2008. Action potential
 generation requires a high sodium channel density in the axon initial segment. *Nat Neurosci*11:178–186. doi:10.1038/nn2040
- Lenz M, Platschek S, Priesemann V, Becker D, Willems LM, Ziemann U, Deller T, Müller-Dahlhaus F,
 Jedlicka P, Vlachos A. 2015. Repetitive magnetic stimulation induces plasticity of excitatory
 postsynapses on proximal dendrites of cultured mouse CA1 pyramidal neurons. *Brain Struct Funct* 220:3323–3337. doi:10.1007/s00429-014-0859-9
- Lenz M, Vlachos A. 2016. Releasing the Cortical Brake by Non-Invasive Electromagnetic Stimulation?
 rTMS Induces LTD of GABAergic Neurotransmission. *Front Neural Circuits* 10:96.
 doi:10.3389/fncir.2016.00096
- Li C, Lu J, Wu C, Duan S, Poo M. 2004. Bidirectional modification of presynaptic neuronal excitability
 accompanying spike timing-dependent synaptic plasticity. *Neuron* 41:257–268.
 doi:10.1016/s0896-6273(03)00847-x
- Lombardo J, Sun J, Harrington MA. 2018. Rapid activity-dependent modulation of the intrinsic
 excitability through up-regulation of KCNQ/Kv7 channel function in neonatal spinal motoneurons.
 PloS One 13:e0193948. doi:10.1371/journal.pone.0193948
- Mahon S, Charpier S. 2012. Bidirectional plasticity of intrinsic excitability controls sensory inputs
 efficiency in layer 5 barrel cortex neurons in vivo. J Neurosci 32:11377–11389.
 doi:10.1523/JNEUROSCI.0415-12.2012
- Mahon S, Deniau J-M, Charpier S. 2004. Corticostriatal plasticity: life after the depression. *Trends Neurosci* 27:460–467. doi:10.1016/j.tins.2004.06.010
- 593 Makowiecki K, Harvey AR, Sherrard RM, Rodger J. 2014. Low-intensity repetitive transcranial 594 magnetic stimulation improves abnormal visual cortical circuit topography and upregulates BDNF

595	in mice. J Neurosci 34:10780–10792. doi:10.1523/JNEUROSCI.0723-14.2014				
596	Manns ID, Sakmann B, Brecht M. 2004. Sub- and suprathreshold receptive field properties of pyramidal				
597	neurones in layers 5A and 5B of rat somatosensory barrel cortex. J Physiol 556:601-622.				
598	doi:10.1113/jphysiol.2003.053132				
599	Mix A, Hoppenrath K, Funke K. 2015. Reduction in cortical parvalbumin expression due to intermittent				
600	theta-burst stimulation correlates with maturation of the perineuronal nets in young rats. Dev				
601	Neurobiol. 75:1-11. doi:10.1002/dneu.22205				
602	Müller-Dahlhaus F, Vlachos A. 2013. Unraveling the cellular and molecular mechanisms of repetitive				
603	magnetic stimulation. Front Mol Neurosci 6:50. doi:10.3389/fnmol.2013.00050				
604	Naudé J, Paz JT, Berry H, Delord B. 2012. A theory of rate coding control by intrinsic plasticity effects.				
605	PLoS Comput Biol 8:e1002349. doi:10.1371/journal.pcbi.1002349				
606	Niemegeers CJ, Schellekens KH, Van Bever WF, Janssen PA. 1976. Sufentanil, a very potent and				
607	extremely safe intravenous morphine-like compound in mice, rats and dogs. Arzneimittelforschung				
608	26 :1551–1556.				
609	Oostenveld R, Fries P, Maris E, Schoffelen J-M. 2011. FieldTrip: Open source software for advanced				
610	analysis of MEG, EEG, and invasive electrophysiological data. Comput Intell Neurosci				
611	2011 :156869. doi:10.1155/2011/156869				
612	Pashut T, Magidov D, Ben-Porat H, Wolfus S, Friedman A, Perel E, Lavidor M, Bar-Gad I, Yeshurun				
613	Y, Korngreen A. 2014. Patch-clamp recordings of rat neurons from acute brain slices of the				
614	somatosensory cortex during magnetic stimulation. Front Cell Neurosci 8:145.				
615	doi:10.3389/fncel.2014.00145				
616	Pasley BN, Allen EA, Freeman RD. 2009. State-dependent variability of neuronal responses to				
617	transcranial magnetic stimulation of the visual cortex. Neuron 62:291-303.				
618	doi:10.1016/j.neuron.2009.03.012				
619	Paxinos G, Watson C. 1986. The Rat Brain in Stereotaxic Coordinates: Hard Cover Edition. Elsevier.				
620	Radman T, Ramos RL, Brumberg JC, Bikson M. 2009. Role of cortical cell type and morphology in				
621	subthreshold and suprathreshold uniform electric field stimulation in vitro. Brain Stimulat 2:215-				
622	228, 228.e1–3. doi:10.1016/j.brs.2009.03.007				
623	Rodger J, Mo C, Wilks T, Dunlop SA, Sherrard RM. 2012. Transcranial pulsed magnetic field				
624	stimulation facilitates reorganization of abnormal neural circuits and corrects behavioral deficits				
625	without disrupting normal connectivity. FASEB J 26:1593-1606. doi:10.1096/fj.11-194878				
626	Rohan ML, Yamamoto RT, Ravichandran CT, Cayetano KR, Morales OG, Olson DP, Vitaliano G, Paul				
627	SM, Cohen BM. 2014. Rapid mood-elevating effects of low field magnetic stimulation in				
628	depression. Biol Psychiatry 76:186–193. doi:10.1016/j.biopsych.2013.10.024				
629	Rossini PM, Burke D, Chen R, Cohen LG, Daskalakis Z, Di Iorio R, Di Lazzaro V, Ferreri F, Fitzgerald				
630	PB, George MS, Hallett M, Lefaucheur JP, Langguth B, Matsumoto H, Miniussi C, Nitsche MA,				
631	Pascual-Leone A, Paulus W, Rossi S, Rothwell JC, Siebner HR, Ugawa Y, Walsh V, Ziemann U.				
632	2015. Non-invasive electrical and magnetic stimulation of the brain, spinal cord, roots and				
633	peripheral nerves: Basic principles and procedures for routine clinical and research application: An				
634	updated report from an I.F.C.N. Committee. Clin Neurophysiol. 126:1071-1107.				
635	doi:10.1016/j.clinph.2015.02.001				

- Rotem A, Moses E. 2008. Magnetic stimulation of one-dimensional neuronal cultures. *Biophys J* 94:5065–5078. doi:10.1529/biophysj.107.125708
- Schramm AE, Carton-Leclercq A, Diallo S, Navarro V, Chavez M, Mahon S, Charpier S. 2020.
 Identifying neuronal correlates of dying and resuscitation in a model of reversible brain anoxia.
 Prog Neurobiol 185:101733. doi:10.1016/j.pneurobio.2019.101733
- 641 Sherrard RM, Morellini N, Jourdan N, El-Esawi M, Arthaut L-D, Niessner C, Rouyer F, Klarsfeld A,
 642 Doulazmi M, Witczak J, d'Harlingue A, Mariani J, Mclure I, Martino CF, Ahmad M. 2018. Low 643 intensity electromagnetic fields induce human cryptochrome to modulate intracellular reactive

644 oxygen species. *PLoS Biol* **16**:e2006229. doi:10.1371/journal.pbio.2006229

- Silvanto J, Pascual-Leone A. 2008. State-dependency of transcranial magnetic stimulation. *Brain Topogr* 21:1–10. doi:10.1007/s10548-008-0067-0
- 647 Silver RA. 2010. Neuronal arithmetic. Nat Rev Neurosci 11:474–489. doi:10.1038/nrn2864
- Stengel C, Vernet M, Amengual JL, Valero-Cabré A. 2021. Causal modulation of right
 hemisphere fronto-parietal phase synchrony with Transcranial Magnetic Stimulation during a
 conscious visual detection task. *Sci Rep* 11:3807. doi:10.1038/s41598-020-79812-y
- Steriade M. 2004. Neocortical cell classes are flexible entities. *Nat Rev Neurosci* 5:121–134.
 doi:10.1038/nrn1325
- Tang AD, Hong I, Boddington LJ, Garrett AR, Etherington S, Reynolds JNJ, Rodger J. 2016. Low-653 intensity repetitive magnetic stimulation lowers action potential threshold and increases spike 654 layer 335:64-71. firing in 5 pyramidal neurons in vitro. Neuroscience 655 doi:10.1016/j.neuroscience.2016.08.030 656
- Tremblay R, Lee S, Rudy B. 2016. GABAergic Interneurons in the Neocortex: From Cellular Properties
 to Circuits. *Neuron* 91:260–292. doi:10.1016/j.neuron.2016.06.033
- Vlachos A, Muller-Dahlhaus F, Rosskopp J, Lenz M, Ziemann U, Deller T. 2012. Repetitive Magnetic
 Stimulation Induces Functional and Structural Plasticity of Excitatory Postsynapses in Mouse
 Organotypic Hippocampal Slice Cultures. *J Neurosci*. doi:10.1523/JNEUROSCI.0409-12.2012
- Wilent WB, Contreras D. 2004. Synaptic responses to whisker deflections in rat barrel cortex as a
 function of cortical layer and stimulus intensity. J Neurosci 24:3985–3998.
 doi:10.1523/JNEUROSCI.5782-03.2004
- Williams MS, Altwegg-Boussac T, Chavez M, Lecas S, Mahon S, Charpier S. 2016. Integrative
 properties and transfer function of cortical neurons initiating absence seizures in a rat genetic
 model. *J Physiol* 594:6733–6751. doi:10.1113/JP272162
- Wilson CJ, Kawaguchi Y. 1996. The origins of two-state spontaneous membrane potential fluctuations
 of neostriatal spiny neurons. *J Neurosci* 16:2397–2410.
- Kang J. 2005. The mechanisms and functions of activity-dependent long-term potentiation of
 intrinsic excitability. *Rev Neurosci* 16:311–323. doi:10.1515/revneuro.2005.16.4.311
- Zhu JJ, Connors BW. 1999. Intrinsic firing patterns and whisker-evoked synaptic responses of neurons
 in the rat barrel cortex. *J Neurophysiol* 81:1171–1183. doi:10.1152/jn.1999.81.3.1171
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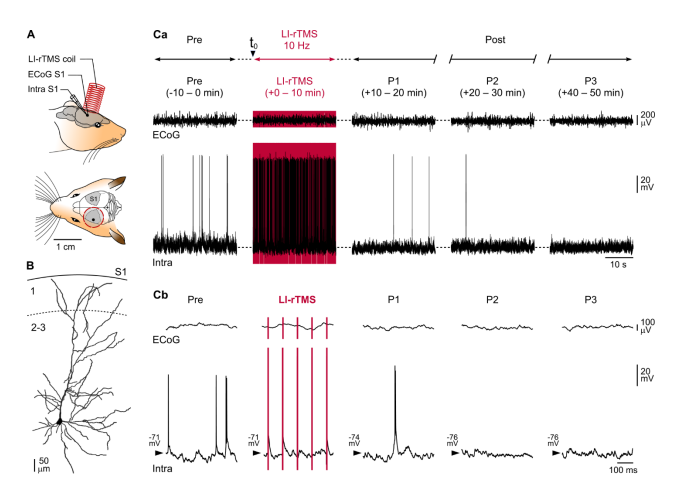
688 Author contributions:

- 689 Conceptualization: SM, RS, SC, AL, AVC
- 690 Methodology: MB, SM, SC, RS, AL, AVC
- 691 Investigation: MB
- 692 Data analysis: MB, PB, CS, SM, RS
- 693 Supervision: SM, RS, SC, AL, AVC
- 694 Writing original draft: SM, SC, RS, AL
- 695 Writing review & editing: SM, SC, RS, AL, MB, PB, AVC, CS
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- 698

- 699 Data and materials availability: All data are available in the main text or the supplementary materials.
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704 Figures and Legends

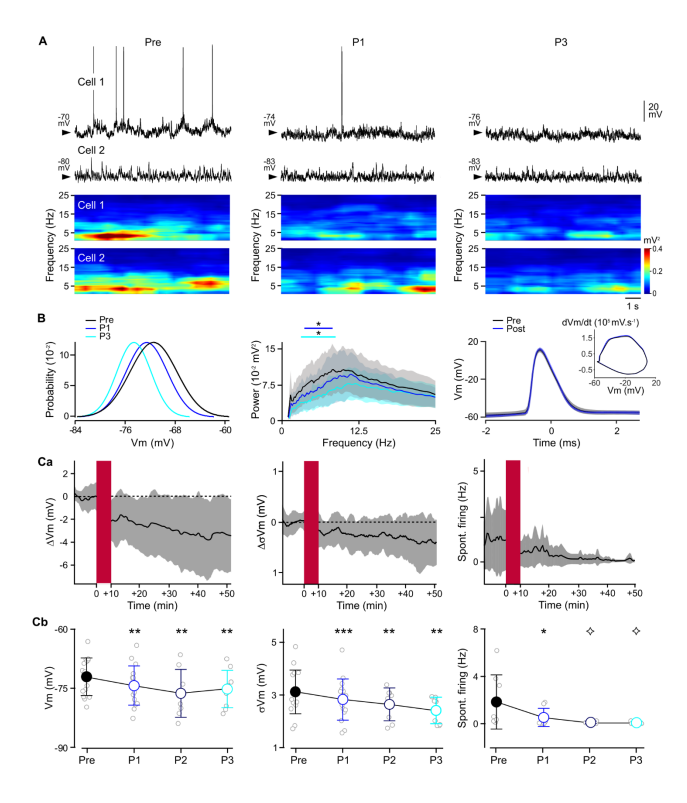




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707 Fig. 1. Experimental paradigm and definition of recording periods.

(A) The experimental set-up illustrating the positions of the LI-rTMS coil and the ECoG (ECoG S1) and 708 intracellular (Intra S1) recording electrodes. LI-rTMS (10 Hz for 10 minutes) was delivered through an 709 8-mm coil placed \sim 2 mm above the brain surface. (B) Reconstruction of a layer 2/3 neurobiotin-filled 710 pyramidal neuron (Ca) Simultaneous ECoG and intracellular recordings from a layer 2/3 pyramidal 711 neuron, before (Pre), during LI-rTMS, and at different times (+10-20 min, P1; +20-30 min, P2; +40-50 712 min, P3) after the onset of LI-rTMS (Post). (Cb) Expanded view (500 ms) of the recordings shown in 713 (Ca). The LI-rTMS artifacts are shown in red. Here and in the following figures, the Vm is indicated at 714 715 the left of intracellular recordings.

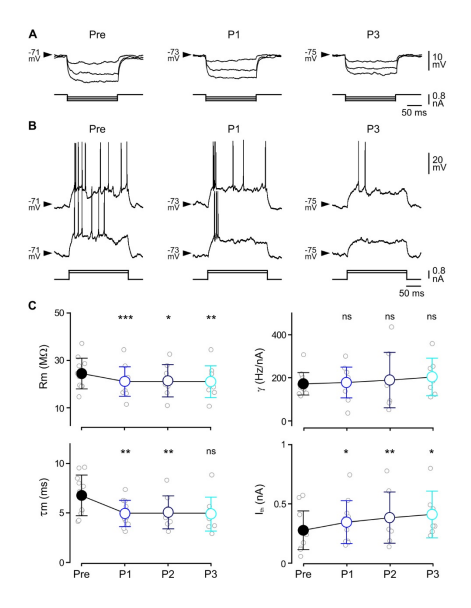




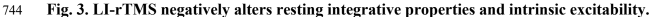
719 Fig. 2. LI-rTMS reduces cortical neuron spontaneous activity.

(A) Spontaneous intracellular activity recorded from active (cell 1) and silent (cell 2) S1 neurons before
 LI-rTMS (Pre) and during the first (P1) and third (P3) post-stimulation periods. The associated time-

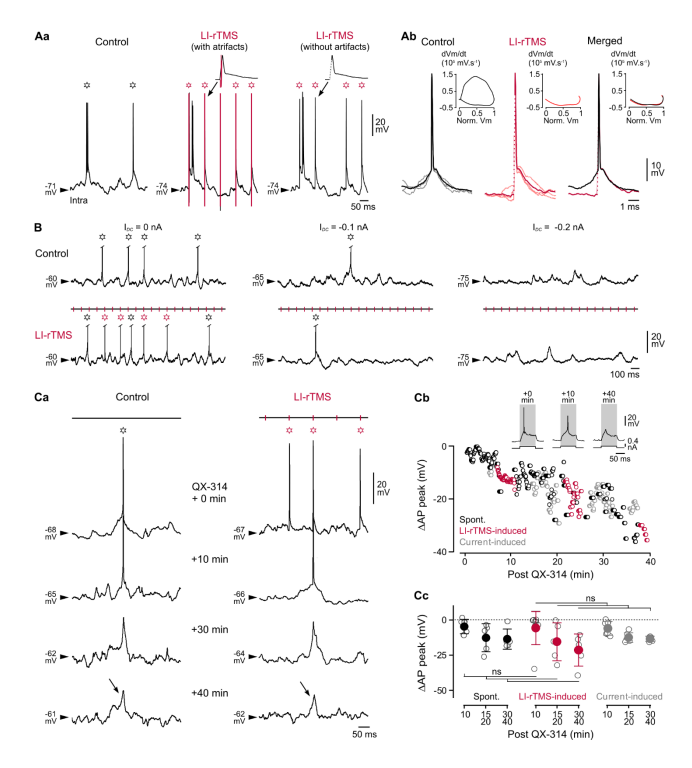
frequency maps, after removal and interpolation of APs, are temporally-aligned below the records. (B) 722 Left: probability densities of Vm values computed from cell 2 during the different periods (60 sec of 723 724 recording, bin size 1 mV). Middle: mean (solid lines) \pm SD (shaded areas) frequency power spectra computed from intracellular subthreshold signals in the pre- and post-stimulation periods (n = 16725 neurons) showing a progressive decrease in the power of the 5-10 Hz frequency band after LI-rTMS 726 (Wilcoxon signed rank test with Benjamini-Hochberg's post hoc correction, comparing P1 and P3 with 727 Pre). Right: superimposed examples of average ± SD (shaded areas) of spontaneous APs and 728 corresponding phase-plots (inset) during Pre (black trace, n = 138) and Post (blue trace, n = 28) periods. 729 (C) Time-dependent changes in spontaneous activity after 10 minutes LI-rTMS. (Ca) Time course of the 730 mean change (black line) \pm SD (gray shaded area) in Vm (after subtraction of the mean Pre value), in the 731 amplitude of Vm fluctuation (σ Vm; after subtraction of the mean Pre value), and in the spontaneous 732 firing frequency of active neurons (Spont, Firing), calculated over successive 30-second periods before 733 and after LI-rTMS application (red rectangle). (Cb) Comparison of Vm, σ Vm and spontaneous firing 734 rate values (mean \pm SD) during baseline (Pre) and at the three post-LI-rTMS times (P1, P2, P3) 735 736 [RMANOVA and Bonferroni post-hoc comparisons; Vm: n = 16, 16, 8 and 9, $F_{0.39, 3.89} = 12.35$, P =0.034; σ Vm: n = 16, 16, 8 and 9, $F_{1,36, 13.55} = 11.69$, P = 0.0025; spontaneous firing: n = 8, 8, 3 and 3, 737 $F_{1.64,6.03} = 9.74$, P = 0.015]. Here and in the following figures, the grav circles in the pooled-data graphs 738 indicate values for individual neurons. In (Ca) and (Cb), analysis of ongoing firing was restricted to 739 neurons which were spontaneously active during baseline. *P < 0.05, ** P < 0.01, *** P < 0.001, ns: 740 non-significant; \Diamond = too few data for post-hoc comparison. 741



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(A) Average (n = 6-30 trials) voltage responses (upper traces) of an S1 pyramidal neuron to 745 hyperpolarizing current steps of increasing intensity (-0.2 to -0.6 nA, lower traces) before (Pre) and after 746 LI-rTMS (P1, P3). (B) Current-evoked firing responses of an S1 neuron to depolarizing current pulses 747 of increasing intensity during Pre, P1 and P3. (C) Population data showing the temporal evolution of 748 mean (\pm SD) values of Rm, τ m, current threshold (I_{th}) and neuronal gain (γ) values. [RMANOVA and 749 Bonferroni post-hoc comparisons; Rm: n = 11, 11, 8 and 9, $F_{0.14, 1.19} = 10.54, P = 0.08; \tau m: n = 11, 11, 8$ 750 and 9, $F_{1.61, 13.44} = 8.24$, P = 0.006; I_{th} : n = 12, 11, 8 and 7, $F_{0.72, 5.55} = 16.91$, P = 0.01; γ : n = 12, 11, 8 and 751 7, $F_{0.14, 1.11} = 0.66$, ns]. *P < 0.05, ** P < 0.01, *** P < 0.001, ns: non-significant. 752

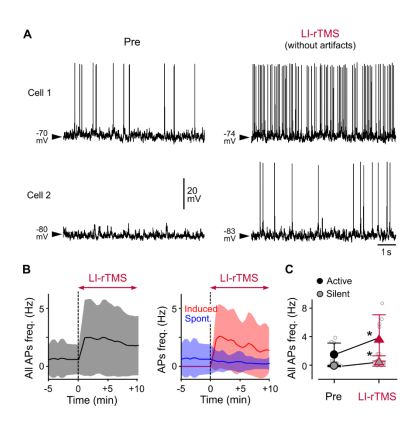


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755 Fig. 4. Evidence for LI-rTMS-induced APs.

(A) Comparison of spontaneous and LI-rTMS-induced APs. (Aa) Intracellular activity of an S1 pyramidal neuron in control and during LI-rTMS, before (with artifacts, vertical red lines) and after (without artifacts) removal of stimulation artifacts. Here and in the following panels, open stars mark the occurrence of spontaneous (black) and putative LI-rTMS-evoked (red) APs. A typical example of a putative LI-rTMS-evoked AP, with and without artifact, is enlarged above the records. Note that the 761 depolarizing phase of the LI-rTMS-evoked AP is hidden within the artifact. (Ab) Superimposition of 3 spontaneous (black) or LI-rTMS-induced (red) APs (light-colored traces) with their corresponding 762 763 average waveform (deeper-colored traces). The AP amplitude is normalized and LI-rTMS artifacts are indicated by dotted red lines. Note the similarity in the repolarization phase of spontaneous and LI-rTMS-764 induced APs when average waveforms are merged (right). The insets show the corresponding AP phase-765 plots (with only the repolarization phase illustrated for LI-rTMS induced APs). (B) Spontaneous and LI-766 rTMS-induced APs are blocked by membrane hyperpolarization. Cortical intracellular activity in absence 767 of (Control, top traces) and during LI-rTMS (bottom traces) at rest (left panel) and during membrane 768 hyperpolarization induced by intracellular injection of negative DC-current (middle and right panels). 769 Red vertical markers above the traces indicate the position of the removed artifacts. (C) AP blockade by 770 OX-314 in control and during LI-rTMS. (Ca) Examples of intracellular records obtained in absence of 771 (left) and during (right) LI-rTMS at the indicated times after impalement with a microelectrode 772 773 containing 100 mM QX-314. Times after impalement are indicated between each set of traces. Note the progressive decrease in amplitude of both spontaneous and LI-rTMS-triggered APs, which start to 774 resemble sharp synaptic depolarizations at 30–40 min after onset of QX-314 diffusion (arrows). (Cb) 775 Changes in peak potential value of individual spontaneous (black), LI-rTMS-induced (red) and current-776 induced (gray) APs in an S1 neuron, as a function of time after onset of QX-314 diffusion. The peak 777 value of the first AP in each group, measured just after cell impalement, was taken as reference and 778 artificially set to zero mV. Insets show representative current-evoked responses at the indicated times 779 after cell impalement. (Cc) Pooled data comparing the peak potential decrease of spontaneous (black), 780 LI-rTMS-induced (red) and current-induced (gray) APs at different times after QX-314 diffusion onset. 781 [Mixed Effects ANOVA; Spont: n = 7, 5, 5 neurons; LI-rTMS-induced: n = 8, 5, 5 neurons; Current-782 induced: n = 8, 6, 5. Effect of time, $F_{1,21}, 9.72 = 16.64, P = 0.0017$; Effect of stimulation, $F_{1,33}, 10.64 = 20.13$ 783 P = 0.0006). Between groups ANOVA: t0, $F_{2, 24} = 2.34$; t10, $F_{2, 20} = 1.56$; t15-20, $F_{2, 13} = 0.68$; t30-40, 784 $F_{2,12} = 1.81$. At each time-point, all comparisons, ns]. ns: non-significant. 785

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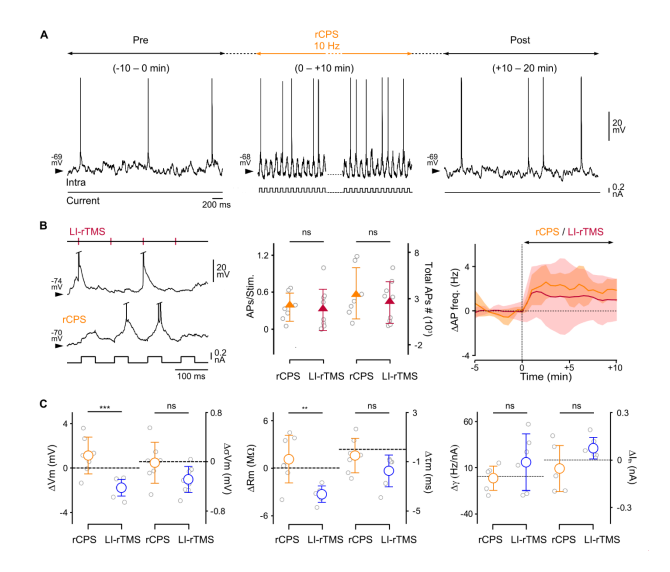


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790 Fig. 5. Changes in firing rate during LI-rTMS.

(A) Intracellular activity of representative active (Cell 1, located in layer 2/3) and silent (Cell 2, located 792 793 in layer 5) cortical neurons recorded before (Pre) and during LI-rTMS after artifact removal. (B) Time course of the mean (\pm SD, n = 16 neurons) changes in global firing frequency (including both spontaneous 794 and LI-rTMS-induced APs; left) and of the specific rate of spontaneous (blue line) and TMS-mediated 795 (red line) APs (right), calculated over 30-second periods before and during LI-rTMS (red arrow). (C) 796 Summary plot comparing the mean (± SD) total AP frequency in Pre and LI-rTMS periods, separating 797 cells that were active (black circles) and silent (gray circles) during baseline. [n = 8 active and 7 silent]798 neurons, Wilcoxon Signed Rank test, *P < 0.05]. 799

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Fig. 6. Repetitive current-evoked firing in a single cortical neuron does not reproduce LI-rTMS
 effects

(A) Cortical intracellular activity before, during and after application of repetitive current pulse 805 stimulation (rCPS). The rCPS protocol consisted of injecting a series of depolarizing current pulses (0.2 806 nA, 40 ms duration) at 10 Hz for 10 min (lower trace). (B) Comparison of neuronal firing during rCPS 807 and LI-rTMS protocols. Left: Short periods of intracellular activity during LI-rTMS (top trace) and rCPS 808 (bottom trace) paradigms. Onset times of removed LI-rTMS artifacts are indicated by red vertical 809 markers and the timing of depolarizing current pulses is shown below the bottom trace. Middle: Pooled 810 data comparing the mean (± SD) number of APs/stim. evoked by LI-rTMS or current pulses, and the 811 total number of APs (evoked and spontaneous) measured over the whole stimulation period in response 812 to rCPS (n = 7 neurons, orange triangles) or LI-rTMS (n = 9 neurons, red triangles). Right: Time course 813 of the mean change (solid lines, n = 7 neurons) \pm SD (shaded areas) in AP frequency (after subtraction 814 of the mean Pre value) calculated over successive 30-second periods, before and during rCPS (orange) 815

or LI-rTMS (red) protocols. Dotted line indicates the mean Pre value, artificially set to zero. (C) 816 Summary data comparing the changes in Vm, σ Vm, Rm, τ m, γ and Ith values between pre- and post-817 stimulation periods after application of rCPS (orange circles) or LI-rTMS (blue circles) protocols. 818 Dashed lines indicate the mean pre-stimulation value (Pre), artificially set to zero [Vm: $n = 7, 8; \sigma$ Vm: 819 $n = 7, 9, \text{Rm}: n = 7, 6; \gamma: n = 7, 6; \text{Ith}: n = 7, 6; \text{ unpaired Student t-test; and } \tau\text{m}: n = 7, 6; \text{Mann Whitney}$ 820 Rank Sum test; ** P < 0.01, *** P < 0.001, ns: non-significant]. 821 822 823 824 **Supplementary Materials** 825 826

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Supplementary Materials for

In vivo low-intensity magnetic pulses durably alter neocortical neuronal excitability and spontaneous activity

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Figs. S1 to S3 Table S1

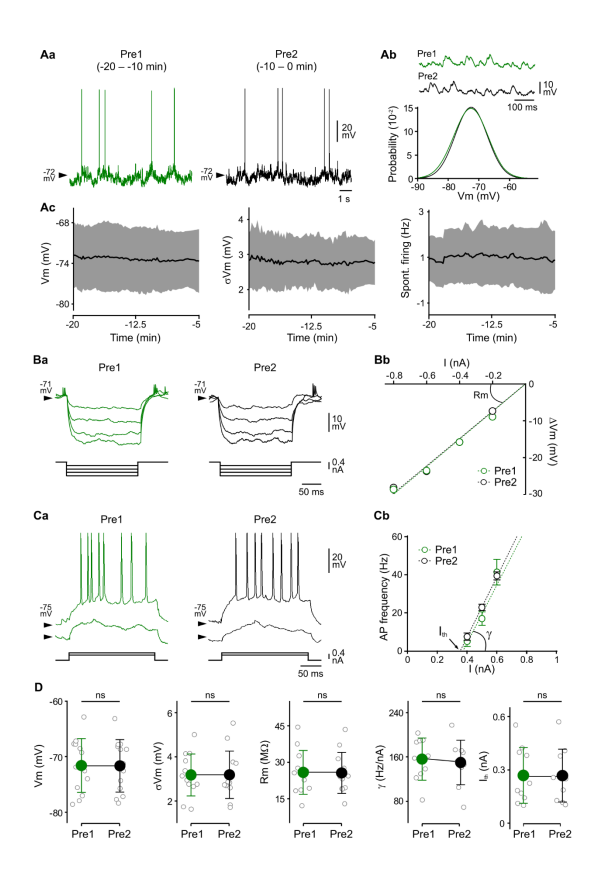


Fig. S1. Stability of resting integrative properties and intrinsic excitability of cortical neurons over baseline periods.

(Aa) Spontaneous intracellular activity of an S1 neuron recorded for two successive periods of 10 minutes: between 20-10 min (green trace, Pre1) and 10-0 min (black trace, Pre2) before the onset of the 10 min 10 Hz LI-rTMS protocol. Neuronal activity was characterized by a barrage of smallamplitude, broadband frequency, synaptic fluctuations (see also Fig. 2, A and B), confirming that sufentanil sedation induced cortical dynamics comparable to those observed during the waking state (Bruno and Sakmann, 2006; Constantinople and Bruno, 2011; Altwegg-Boussac et al., 2014). This background synaptic activity induced firing in 7 out of 13 neurons. Spontaneous firing rate of the active neurons was 2.9 ± 3.5 Hz during the first and 2.8 ± 3.5 Hz during the second prestimulation period (paired Student t-test, P = 0.29). Vm values are indicated at the left of intracellular records. (Ab) The probability densities of Vm values (60 s of recording, bin size 1 mV) demonstrate stability in the amplitude of synaptic activity over time. (Pre1 Vm = -71.6 ± 4.9 mV, n = 13 neurons versus Pre2 Vm, -71.7 ± 4.7 mV, n = 13 neurons; paired Student t-test, P =0.92). (Ac) Time course of the mean values (solid lines, n = 8 neurons) \pm SD (grav shaded areas) of Vm, σ Vm and spontaneous firing frequency computed over successive 30-second epochs during the 15 minutes of recording preceding LI-rTMS onset. Pearson's correlations indicate a lack of significant changes over time. (B) Stability of membrane input resistance. (Ba) Average (n = 10-17 trials) voltage responses (top traces) to negative current pulses of increasing intensity (bottom traces) recorded from an S1 pyramidal neuron during the two baseline periods (green traces, Pre1 and black traces, Pre2). The corresponding V-I relationships shown in (Bb) were best fitted by linear regressions (dashed lines, both $r^2 > 0.99$), indicating a lack of substantial membrane rectification in the hyperpolarizing direction. (C) Constancy of cell intrinsic excitability over the pre-stimulation period. (Ca) Examples of current-induced firing responses to depolarizing current steps of increasing intensity recorded during the two successive baseline periods (green traces, Pre1; black traces, Pre2). (Cb) F-I relationships and corresponding linear fits for the neuron illustrated in (Ca). Each data point is the mean (\pm SD) firing rate calculated from 10–18 successive trials. (D) Population data comparing Vm, σ Vm, Rm, γ and I_{th} values during the two successive baseline periods. Gray circles correspond to values of individual neurons and green and black symbols show the mean \pm SD. [Vm: n = 13; σ Vm: n = 13; Rm: n = 12; γ : n = 10; Ith: n = 10; paired Student t-test; *P < 0.05, ***P < 0.001, ns: non-significant].

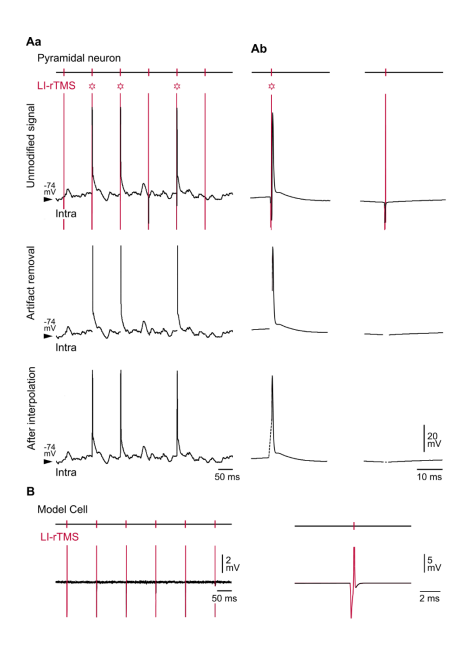


Fig. S2. Procedure for removing LI-rTMS artifacts.

(A) Off-line cutting and interpolation of electrical artifacts. (Aa) Short periods of intracellular activity recorded from a layer 2/3 pyramidal neuron during LI-rTMS, before removal of stimulation artifacts (vertical red lines, Unmodified signal), after their removal (Artifact removal), and after their replacement by linear interpolation (After interpolation). Red stars mark the onset of LI-rTMS-induced AP occurrences. Vm values are indicated at the left of the records. (Ab) Left, Representative example of an averaged (n = 2800) LI-rTMS-induced AP before and after removal of the LI-rTMS pulse stimulation artifact. Right, average cell responses (n = 3200) to magnetic pulses in absence of evoked APs. (B) Current-clamp recording during LI-rTMS illustrating the

absence of evoked electrical responses in the aftermath of stimulation artifacts (in red) when a CLAMP-1U model cell is simulating the recording of a pyramidal neuron. An expanded view of a single stimulation is shown at right.

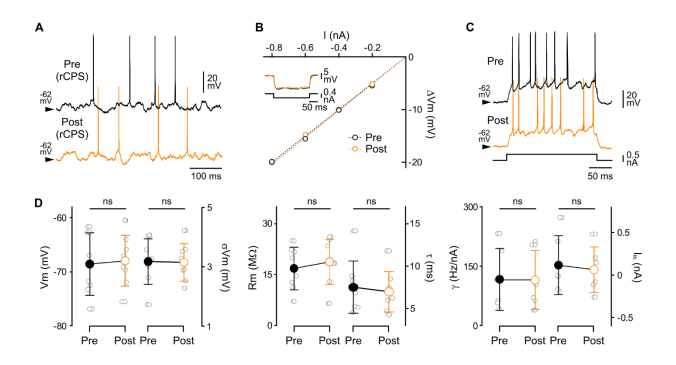


Fig. S3. Stability of neuronal resting integrative properties and intrinsic excitability after rCPS.

(A) Spontaneous intracellular activity recorded from an S1 neuron before (black trace, Pre) and 15 min after (orange trace, Post) the onset of the rCPS protocol. Vm values are indicated at the left of intracellular records. (B) *V*–*I* relationships computed from the neuron illustrated in (A) before and following the application of the rCPS protocol. The average voltage responses (20–30 trials) to negative current pulses of -0.4 nA (200 ms duration) during pre- and post-stimulation periods are superimposed in the inset. *V*–*I* relationships were best fitted by linear regressions (dashed colored lines, $r^2 > 0.99$). (C) Examples of current-evoked firing responses in baseline (Pre) and following rCPS (Post). (D) Population data comparing Vm, σ Vm, Rm, τ m, γ and I_{th} values during the pre- and post-rCPS periods. Gray circles correspond to values of individual neurons and black and orange symbols show the mean ± SD. [Vm: n = 7; σ Vm: n = 7; Rm: n = 6; τ m: n = 6; γ : n = 6; Ith: n = 6; paired Student t-test, ns: non-significant].

	Threshold (mV)	Amplitude (mV)	Duration at mi- amplitude (ms)	Peak amplitude (mV)
Pre (-10 – 0 min.)	-52.2 ± 1.1	59.7 ± 4.6	0.66 ± 0.2	7.5 ± 4.9
P1 (+10 – +20 min.)	-52.5 ± 2.2	59.2 ± 5.0	0.67 ± 0.2	6.4 ± 4.9

Table S1: Properties of spontaneous APs are similar before and after LI-rTMS.

Values are mean \pm SD. No significant difference was found between the pre- and post-stimulation (P1) period (n = 8 neurons; paired Student t-test; P > 0.16 for each parameter).