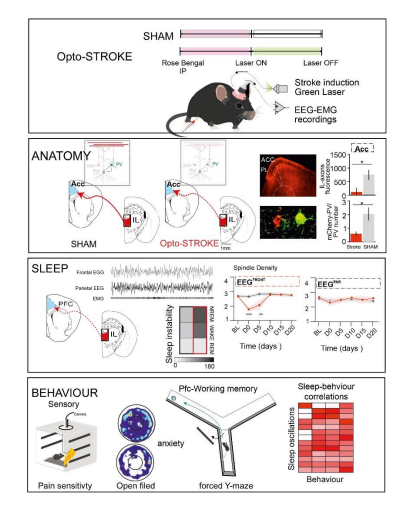
1 2	<i>Title:</i> Optical mini-stroke of thalamic networks impairs sleep stability, topography and cognition.
3	
4	Lenzi I ² ., Borsa M ² ., Czekus C ¹ ., Rusterholz T ¹ , Bassetti C. L., ^{1,2} , Gutierrez Herrera C ^{1,3*} .
5 6	¹ Center for Experimental Neurology, Department of Neurology, Inselspital University Hospital Bern, University of Bern, Bern, Switzerland.
7 8	² Sleep-Wake-Epilepsy Center, Department of Neurology, Inselspital University Hospital Bern, University of Bern, Bern, Switzerland.
9 10	³ Department of Biomedical Research (DBMR), Inselspital University Hospital Bern, University of Bern, Bern, Switzerland.
11	
12	
13	
14	
15	
16	
17	
18	
19	
20	
21	* Correspondence to:
22 23 24 25 26 27 28 29 30 31	Carolina Gutierrez Herrera, PhD, PD. Center for Experimental Neurology, Department of Neurology Inselspital University Hospital, University of Bern Freiburgstrasse 18 3010 Bern, Switzerland Tel: +41 (0) 31 632 55 93 carolina.gutierrez@dbmr.unibe.ch
32 33 34 35	
36	Abstract: 194 words
37 38	Main text: 11293 words Figures: 5
39	Suppl. figures 7

40 Abstract

Modelling stroke in animals remains a challenge for translational research, especially for the 41 infraction of small subcortical arteries. Using combined fibre optics and photothrombosis 42 technologies, we developed a novel model of optically-induced infarcts (Opto-STROKE). 43 44 Combining our model with electrophysiological recordings in freely-behaving mice, we studied 45 early and late consequent patho-physiological changes in the dynamics of sleep-wake circuits 46 and cognitive performance. Here, focusing on inducing Opto-STROKE lesions in the intralaminar thalamus (IL), which in humans cause severe impairments of arousal, cognition, 47 48 and affective symptoms, our model recapitulated important deficits on sleep disorders presented in humans including arousal instability, concurrent to an augmented slow-wave 49 activity and a reduction gamma power bands during wakefulness. Moreover, during NREM 50 sleep, spindle density was decreased and topographically shifted to frontal cortices when 51 compared to control animals. Remarkably, gamma power and spindle density were correlated 52 with decreased pain threshold and impaired prefrontal cortex- dependent working memory in 53 Opto-STROKE mice relative to controls. Collectively, our combined method influences both 54 anatomical and functional outcomes of the classical stroke procedures and offers new insights 55 on the fundamental role of the media thalamus as a hub for the regulation of both sleep-wake 56

57 architecture and cognition.



58 Introduction

Stroke is a devastating brain disorder leading cause of morbidity and mortality worldwide that 59 60 involves tissue damage and extensive structural modifications that are associated with severe 61 physiological and behavioural consequences. The severity of clinical outcomes partly depends 62 on the localization of the thrombus and the vascular territories involved. Although ischemic 63 stroke management has advanced along with the management of pathological deficits, this 64 disorder still remains a leading cause of long-term serious consequences in motor-, sensory-, sleep- and cognitive-related functions in patients which life quality never fully recovers. This is 65 particularly true in thalamic infarcts involving the territory of the paramedian arteries (1). 66 67 Paramedian strokes represent a unique category bearing a variety of outcomes, including altered conscious states, counting coma (2), hypersomnia with alterations in non-rapid eye 68 movement (NREM) sleep architecture (3, 4); marked cognitive disturbances involving 69 reference memory recall, attention, enhanced distractibility and confusion (5-7); dysfunctional 70 emotional regulation, memory loss and perseveration (2); sensory processing disturbances, 71 such as in pain sensitivity. While sleep-related dysfunctions often improve over time, however 72 73 memory deficits and changes in behaviours tend to persist (2, 3). These numerous clinical symptoms suggest the insurgence of disconnection to subcortical structures. Despite major 74 advances on acute stroke management and neuro rehabilitation, most of disabilities remain 75 76 untreatable.

77 Paramedian strokes result from lesions of the tuberothalamic artery or, in rare cases, of the paramedian artery, irrigating the thalamic reticular nucleus (TRN), intralaminar thalamus (IL) 78 79 which comprises the parafascicular nucleus (PF), medio-dorsal thalamus (MD) and central median thalamus (CMT), ventral anterior nucleus of the thalamus (VA), rostral part of the 80 81 ventrolateral nucleus of the thalamus (VL), mamillothalamic tract, ventral amygdalofugal pathway, the ventral part of the internal medullary lamina, and the anterior medial (AM), ventral 82 (AV) and dorsal (AD) thalamic nuclei (8, 9). Together as a functional class (2), these nuclei 83 subserve arousal and various cognitive functions. Specifically, they are involved in the 84 modulation of wake-sleep patterns, sleep oscillations, sensory processing, attention, goal-85 oriented behaviours and associative memory through their substantial connectivity to the 86 cortex (thalamo-cortical), to other thalamic nuclei (intra-thalamic), as well as to other 87 subcortical regions (10-16). According to their broad system of efferent and afferent 88 89 projections, the IL have been recognized as higher order thalamic nuclei, involved in higher 90 associative cognitive functions. Previous works have shown that MD and CMT neurons extensively project to parvalbumin expressing interneurons (PV+) in the anterior cingulate 91 92 cortex (ACC) (17-20), which are responsible for feedforward inhibition ultimately modulating cortical excitability and brain states (21–23). Moreover, IL -prefrontal cortex (PFC) circuits have 93 94 been shown to play a role in memory and attention-related tasks (24-26). Thus, understanding

the relationship between the topography of sleep oscillatory activities and behavioural 95 outcomes after stroke lesions is indispensable. Experimental stroke models have greatly 96 advanced our understanding of development and consequences of stroke lesions. However, 97 98 infractions of small subcortical arteries - one of the three major causes of human ischemic 99 stroke (48) – still remains a challenge, as the absence of non-anesthetized freely-behaving 100 models thus limiting the comparability of pre-clinical models to clinical cases of stroke (48, 49). 101 These latter factors are particularly relevant in presence of vigilance disturbances, such as the 102 ones caused by paramedian stroke. Here, we developed a novel model for optically induced 103 mini-photothrombotic ischemia targeting subcortical areas (Opto-STROKE) that does not 104 require anaesthesia and allows longitudinal (up to 4 weeks) multi-site electrophysiological 105 recordings of sleep and circuit activity in freely behaving mice. By means of this novel technique, we studied the link between the topography of IL ischemic lesions and related 106 changes in postsynaptic targets, sleep/wake and behaviour. Our findings showed that IL Opto-107 108 STROKE increases the fragmentation of sleep and wakefulness associated with sustained changes in slow wave activity and gamma oscillations in wakefulness when compared to 109 110 SHAM control animals. Fronto-parietal expression of spindles was affected in IL-opto-STROKE animals in the acute phase after stroke. Interestingly, we found a negative correlation 111 between sensory-related responses (pain) at day 20 after stroke and the power of gamma at 112 the day of stroke induction, as well as between impairments in PFC-dependent memory 113 114 retrieval in the subacute phase and spindle density on the acute phase post-stroke. In 115 summary, our data implicates the IL as a fundamental hub for the regulation of both sleepwake architecture and cognition, providing further insights on the relationship between sleep 116 117 and cognition. Furthermore, this work contributes to the understanding of the temporal progression of thalamic strokes' symptoms and plastic changes at the lesion site and within its 118 post-synaptic targets. Overall, we propose the Opto-STROKE model as novel selective 119 methodological approach to further dissect structural and functional consequences of local 120 121 mini-strokes in subcortical structures and to explore crucial time windows for interventions.

122 Methods

123 Animals

(https://www.janvier-labs.com/en/fiche produit/%20c57bl-6jrj 124 C57BL/6JRj male mice 125 mouse/10-20 weeks old, 23-30 g). Animals were kept in groups of 2-5 per individually ventilated cages (IVC) under controlled conditions (regular circadian cycle of 12:12 hours light: 126 dark; lights on at 8:00 AM (ZT0), constant temperature 22-24°C and humidity 30%-50%). 127 Animals included for sleep and behavioural experimentations were habituated to a light-dark 128 129 cycle of 12 h (lights on at 08:00, ZT0) for 5 days prior the surgical procedures. Following viral transduction and chronic electroencephalogram (EEG) and electromyogram (EMG) electrodes 130 131 implantation, mice were all housed individually and let recover for 7-10 days in custom-

designed polycarbonate cages (300 mm x 170 mm). Then, animals were tethered and allowed 132 to progressively adapt to the EEG/EMG and optic cables in their home cage for an additional 133 7-10 days to then remain plugged for the duration of the experiment. Animals were randomly 134 135 assigned to two experimental groups: SHAM (control) and Opto-STROKE (experimental). Both 136 groups underwent same injection, instrumentation and sleep and behavioural monitoring protocols. However, only stroke animals underwent the stroke induction protocol. All animals 137 were treated according to Swiss animal care laws, and experimental procedures were 138 approved by local authorities (Veterinary Office, Canton of Bern, Switzerland; license numbers 139 140 BE 41/17 and BE 118/2020).

141

142 Viral targeting

Viral injections were performed in animals of 10 to 16 weeks old as previously described (39, 143 50). Briefly, animals were anesthetized with isoflurane (4.0% induction, 1.5% maintenance). 144 145 Body temperature was constantly monitored and kept at physiological range using a rectal thermo-probe and feedback-controlled heating system. Animals were fixed in a digital 146 stereotaxic frame, analgesia was administered subcutaneously (meloxicam, 5 mg/kg), and 147 lidocaine 2 mg/kg infused subcutaneously at the incision site. All animals received an 148 intracranial injection of 200 nl AAV2-CaMKII-mCherry viral vectors (50 nl / min infusion rate) 149 through 28 G stainless steel cannula (Bilaney), connected by a tubing to a 10 ml Hamilton 150 syringe in an infusion pump (model 1200, Harvard Apparatus). Injections were performed to 151 target the Intralaminar thalamus (IL, AP: -1.6 mm; ML: -0.75 mm; DV: -3.9 mm, 10°). Animals 152 were given seven days to recover before instrumentation surgery. 153

154

155 Instrumentation

156 Animals at 12-20 weeks of age were chronically implanted with a unilateral optic fibre (diameter 200 µm, part number = FT200UMT, ThorLabs) unilaterally above the IL (IL, AP: -1.6 mm; ML: 157 -0.75 mm; DV: -3.8 mm, 10°) and bilaterally over the reticular thalamic nucleus (TRN, AP 158 -0.8 mm, ML +1.7 mm, DV -3.5 mm) and the anterior dorsal thalamus (AD, AP -0.86 mm, ML 159 +0.75 mm, DV -2.71 mm) together with an EEG / EMG connector (Straight Male PCB Header: 160 cat. # 852-10-100-10-001101, Preci-Dip) as previously reported (39). Animals were 161 instrumented with two screws over the frontal cortices (AP: 2 mm; ML: +/- 2 mm), two over the 162 posterior cortices (AP: -3 mm; ML: +/-2 mm) and one the cerebellum as ground (Stainless steel 163 screws diameter: 1,9 x 3,18 mm, Paul Korth GmbH & Co. KG) were in planted. In addition, two 164 bare-ended EMG wires (Cat. # W3MUF 8/30-4046 55, 3 wire international (3WI)) were sutured 165 to the neck muscles to record postural tone (Suppl. Figure 1A). 166

- 167
- 168

169 Thalamic targeted strokes

Opto-STROKE induction was performed in animals at 13 to 23 weeks of age were 170 intraperitoneally (IP) injected with 0.10 ml photosensitive dye Rose Bengal (Sigma Aldrich, cat 171 No 632-69-9) diluted to 10 mg/mL in NaCI. Within 5-10 minutes after injection, 10 mW of 532-172 173 nm laser (LRS-0532-GFM-00100-03, Laserglow Technologies) was delivered through the optic fibre coupled to the laser into the AD, TRN or IL. For the SHAM procedure, animals received 174 175 an IP injection of Rose Bengal but light was delivered. For anatomical investigation of at day 0, 5 and 10 of infarct induction, mice underwent the same procedure, but the optic fibre was 176 177 removed from the brain 5 minutes post-illumination. All stroke inductions took place between ZT 4-5. Please note that the rage in the animals age was with the intent to model population 178 179 variability and thus reduce discrepancies with the human disease.

180

181 Data acquisition

Sleep recordings: EEG and EMG signals were amplified (× 1,000) using a multichannel differential amplifier (model 3500, AM System) and digitized at 512 Hz (NIDAQ 6363, National Instruments) using a sleep recording software (Sleep Score, View Point). A 48 hours' baseline of spontaneous sleep-wake behaviour were recorded for all animals. Following, SHAM and Opto-STROKE animals were recorded in the infarct's acute phase, or rather during stroke induction and for the following 12 hours (day 0), and semi-chronic phase at days five, ten, fifteen and twenty post-stroke, to follow the progression of sleep behaviours and oscillations.

Behavioural tests: Anxiety-, motor-, cognitive-related outcomes were assessed in sleep recordings-free days. Video recordings of the performances were analysed with Ethovision software (Noldus), allowing the tracking of the mice movements within the whole area and predetermined fractions of the arenas, such centre in the open filed test. For the pain sensitivity testing sessions, videos were recorded via an USB-camera included in the fear conditioning system (Fear Conditioning 2.1, Ugo Basile).

195

196 Sleep staging

As previously described (39, 50) electrophysiological data were manually scored and analysed 197 using EEGlab (39). Exact transitions of three vigilance states were identified based on 198 EEG/EMG frequency and amplitude. Wakefulness was determined by low-amplitude EEG and 199 high amplitude EMG signals. Whereas NREM sleep was scored in period of high-amplitude 200 EEG signals, rich in low-frequency oscillations (0.5-4 and 10-16) concomitant to reduced EMG 201 tone. REM sleep was characterized by low amplitude, high EEG theta (5-9 Hz) power and 202 isoelectric EMG with intermittent muscle twitches. Microarousals were scored and defined as 203 204 cortical fast rhythm and EMG bursts of at least 1 s. Transitional states were defined as periods between Wake-NREM-Wake featured by the presence of low EMG power and EEGs including
both slow and theta oscillations. Minimal transitional period length was defined as 5 sec.

207 Slow waves and spindles detection

208 Electrophysiological analysis was completed using custom written MATLAB scripts. Slow 209 waves and spindles were detected via customised MATLAB scripts. For slow waves, adapted 210 SWA-MATLAB detection toolbox (39, 53) was used to detect the negative envelope across the four EEG channels, filtered between 0.5 and 4 Hz, and consecutive zero-crossings were 211 212 detected. The threshold in the amplitude of the detected event eliminates the potential 213 individual differences on distances between electrodes and electrode depth that would affect 214 the record amplitude. In a second pass, the activity over all four channels was examined for 215 each SW detected on the negative envelope to obtain individual channel data. For spindle detection, a wavelet power was estimated between 10 and 16 Hz. Then wavelet functions 216 217 classification was performed using the ratio between average power of spindle segments and spindle-free segments. The wavelet energy time series was smoothed using the 200 ms Hann 218 window, and a threshold equal to 3 SD (SD: standard deviation) was applied above the mean 219 220 to detect potential spindle events (54).

221

222 Behavioural tests

All animals underwent three behavioural tests aimed to characterize the motor-, anxiety-, and cognitive-related differences between SHAM and Opto-STROKE animals see Supplemental Figure 1B-D. All tests were run at distinct times and days between sleep recordings. All measures were conducted at least 12 h apart from the last sleep recording in the light phase. To avoid a rewarding odour-based bias to mice choices, after each behavioural session, testing setups were carefully cleaned with 70% ethanol.

Open field test (OFT). To quantify changes in exploratory tendency, motor integrity and anxiety,
 mice were placed one by one, in a round-shaped arena (60 cm in diameter) and let free to
 explore for 5 min (Supplemental Fig 1B). The OFT was repeated two times, once before stroke
 induction (OFT pre-stroke), and another time after (OFT post-stroke) (Suppl. Figure 1A).
 Measured outcomes were total time spent in the centre (s), total distance moved (cm) and
 mean velocity (cm/s) during the 5 min session.

Pain sensitivity test. Differences in sensitivity to foot shocks were assessed via a novel protocol
using triplets of 2 s shocks of increasing intensity (0,05 mA - 0,6 mA) in 0.05 mA increments.
Shocks with same intensity were interspersed with a 10 s interval (inter-triplet), while a 20 s
time was used as intra-triplet interval. Pain responses to shocks were first defined, then
categorized in 3 different pain thresholds (PT): absence of response, or "no response" (PT0);

240 "backward movements" (PT1); "escape run", "jump" and "immobility" (PT2) (modified from
241 (51)). Scoring of the behaviours was done off-line.

242 Y-Maze alternation test (YM). This test aimed to measure PFC-related cognitive performance 243 (52). For habituation, mice were handled for one hour daily for one week prior to the beginning 244 of the experiment. 3 days before starting of the behavioural test's procedures animals were food restricted and monitored to maintain ~ 80% of basal body weight (Suppl. Figure 7). For 245 246 the reward habituation, animals were provided with 0.1 ml of Sweetened condensed milk diluted 1:10 in water, that was later used as reward during training and testing in the YM. 247 248 Animals were habituated to the Y maze arena (Figure 5H and Suppl. Figure 1C, top) on the 249 second day of the food restriction period, where they were allowed to explore the maze for 10 250 min at ZT 3 and ZT 9. The food dispenser positioned in the three-arm ends were baited with 251 food reward (0.1 ml of the diluted sweetened condensed milk). Then animals were trained for two sessions (T1 and T2) to alternate between left and right arms ("goal" arms) to find the 252 reward (See Suppl. Figure 7D for learning curves). During this training sessions, goal arms 253 were alternatively closed and baited to train the mice to run from a starting arm in the open 254 255 goal arm and get the reward with a time limit of 1 min. Testing session consisted of 10 trials, during which animals were expected to remember the alternating strategy learned in the 256 training sessions in order to get the reward. In the first sample trial (trial 0), one of the goal 257 arms was closed, and mice were forced to get the reward in the opposite arm. In the next 258 consecutive trials, mice were not prompt anymore with the door, and needed to alternate 259 260 between right and left arm to find the reward. For a single trial, mice were given a maximum of five runs (or consecutive errors) interleaved by 30 s delay to get the reward. In the case that 261 five consecutive errors occurred, animals were re-directed to the rewarding arm by closing the 262 not-goal arm (see trial 0) to then proceed with a new trial. Performances were video recorded, 263 264 and behaviour scored off line. Errors were ranked from E1 to E5 to evaluate choice perseveration (Suppl. Figure 1C, bottom). Finally, latency to the reward was measured to 265 disclose possible motor deficiency in stroke animals. 266

267

268 Immunohistochemical staining

Animals were sacrificed at the day of stroke induction (day 0) and post-stroke day 5, 10 and 269 270 21 with 150 mg/kg or 0.5 - 1 ml/kg pentobarbital intraperitoneal injection (Esconarkon ad us. vet., Streuli Pharma) and transcardially perfused with 1x phosphate-buffered saline (PBS) 271 followed by 4% paraformaldehyde. Brains were postfixed for 24 hours, cryoprotected in 30% 272 273 sucrose (48 h at 4°C), frozen in 2-methyl-butane on dry ice and cut into 40 µm sections. To 274 measure the stroke volumes, every third slice was mounted onto a glass slide (the other two sets of sections were used for immunostaining), dried at room temperature, rehydrated, and 275 276 processed for Nissl staining. Briefly, sections were immersed in Cresyl violet (Klüver Barrera,

Bio-Optica), washed in distilled water and dehydrated in graded alcohols, cleared in xylene
(Sigma Millipore), and mounted (Eukitt mounting medium, Bio-Optica) on gelatin-coated
microscope slides.

The other two remaining free-floating brain sections were washed in 0.1% PBS with 0.1 % 280 Triton A-10 (PBS-T) and incubated in blocking solution (1 h at room temperature; PBS-T with 281 10% donkey serum, Sigma Life Science). Then, sections were incubated with primary antibody 282 283 to: mCherry (rat anti-mCherry, 1:1000, # M11217), microglia (ionized calcium binding adaptor molecule 1 (lba1) (rabbit anti-lba1 1:1000 Wako 019-19741), reactive astrocytes (mouse anti-284 285 glial fibrillary acidic protein GFAP, 1:800, Catalog # 13-0300), parvalbumin (rabbit antiparvalbumin, 1:600, Ab11427 RRID:AB 298032) and NeuN (mouse anti-neuronal nuclei 286 287 (1:500; cat. # ab104224; Abcam). Following repeated washes in PBS-T, sections were incubated with secondary antibodies (Anti-mouse Alexa Fluor 488; cat. # ab150113; Abcam; 288 Goat anti-rat Alexa Fluor 555 (Cat. # ab150166); Goat anti-rabbit Alexa Fluor 488 (Cat. # 289 ab150077); and Goat anti-mouse Alexa Fluor 647 (Cat. # ab150115) in PBS-T (2 h at room 290 temperature). Finally, slices were washed in PBS 1X, mounted on microscope slides, and 291 292 covered.

293

294 Anatomical quantification

The extent of the Opto-STROKE lesion was evaluated by quantifying stroke edges delineated per section using ImageJ software (<u>https://imagej.nih.gov/ij/</u>). As described before (39), the lesioned area was measured in each brain slice and multiplied by the distance between sections to define the infarct volume.

299 Fluorescent signals from the immunohistological staining were performed by drawing regions 300 of interest (ROI) normalized by the background fluorescence level (same size ROI) per 301 individual section in ImageJ. A subset of brain sections was used to quantify the changes in 302 neuronal projections to parvalbumin-expressing neurons in post-synaptic targets of the IL. For synaptic contact quantification, a squared area of about 250 um² was used to delimit a fixed 303 region in the ACC. Within the delimited zone, IL synaptic contacts (puncta) on to PV⁺ neurons 304 were manually counted. After quantification, our measures were normalized by the number of 305 PV+ neurons counted in the chosen area. Three to four sections per animal were analysed. 306

307

308 Microscopy

Images for anatomical analysis were acquired using a Nikon Eclipse Ti-E fluorescence
 microscope (M.I.C. facility - Mu40, CH-3008 Bern). For stroke volume quantification, images
 were acquired with a using an Olympus light microscope (Widmer Laboratory - Inselspital CH 3010 Bern) and magnified with 10x or 20x objectives. For the fluorescence

immunohistochemical quantification, 10x and/or 20x magnification was used, and 60x for the
 quantification of IL-mCherry⁺ ACC-PV⁺ neurons.

315

316 Statistical analysis

Differences in outcome parameters between SHAM and Opto-STROKE groups at each day of 317 318 stroke progression were analysed using two-way ANOVA with multiple comparisons using 319 LSM model (Prism 6 GraphPad; https://www.graphpad.com/scientific-software/prism/) and Post hoc Bonferroni testing for multiple comparisons between experimental groups and time 320 321 points after stroke induction or otherwise as stated in the text and figure legends. All data are 322 presented as mean \pm SEM, and levels of statistical significance were set at threshold P < 0.05. 323 For the analysis of the behaviour, paired and unpaired Student's t test, or two-way ANOVA were used for the analysis between groups or within groups using Bonferroni post-hoc multiple 324 comparison test. Animals that did not perform behavioural tests or lost the EEG/ EMG signals 325 326 during longitudinal measurements were excluded from the analysis. Experiments were not conducted in a blinded fashion. Data were scored independently by two experimenters. At 327 least 4 cohorts of animals were used for statistical analysis. 328

329

330 Results

331

332 Histological characterization of optical mini-stroke (Opto-STROKE) in thalamic regions.

333 To overcome the limitations of current photothrombotic stroke models (49) and allow the study of subcortical strokes in awake behaving mice, we targeted different regions irrigated by the 334 335 tuberothalamic artery to restrict the ischemic infarct to three discrete reticular and intralaminar (IL) functional thalamic class: the reticular thalamic nucleus (TRN), and the anterior dorsal 336 thalamus (AD) and the IL (via the central median-thalamus (CMT)) (see methods). 337 Quantification of the volume of ischemic infarcts across different stroke locations were 338 consistent in size and restricted to the targeted nucleus and, to a lesser extent, to some of the 339 adjacent nuclei (AD = 0.51 ± 0.06 mm³; TRN= 0.896 ± 0.109 mm³; IL 0.53 ± 0.23 mm³) (Figure 340 1A-F and Suppl. Figure 2 for TRN and AD and Suppl. Figure 3 for IL stroke). Labelling of GFAP 341 and Iba-1 were used to quantify reactive astrocytes and microglia activation as markers of 342 inflammatory pathological processes. We found a higher level of gliosis and microglia 343 activation in Opto-STROKE animals as compared to SHAM at the lesion site (Suppl. Figure 344 2C and E). Quantification of the fractions of IL nuclei affected by the stroke revealed that the 345 central median thalamic (CMT); dorsomedial thalamus (DMT); intermediodorsal (IMD); 346 paracentral (PC); rhomboid (RH); and reuniens (RE) were affected in different proportions 347

348 (CMT = 40.94 ± 15.70 ; DMT = 40.41 ± 8.97 ; IMD = 19.23 ± 7.37 ; PC = 19.83 ± 20.95 ; RH = 349 9.81 ± 4.68 ; RE = 13.26 ± 9.62) (Figure 1F).

Next, we characterized the structural changes of IL neuronal projections following Opto-350 351 STROKE using an anterograde tracing strategy (Figure 1G-H). To label IL neurons and their 352 projections, AAV2-CamKII-mCherry viral vector was stereotactically injected in the IL area for stable expression of reporter fluorescent mCherry protein. Major IL neuron terminals were 353 354 found in the anterior cingulate (ACC), prelimbic region (PL) of the frontal cortex, the anterior insular cortex (AI), zona incerta (ZI), caudo-putamen (CP), nucleus accumbens (ACB) and the 355 356 basolateral amygdala (BLA) (Figure 1G). Interestingly, we found a significant decrease in mCherry fluorescence level in ACC, in Opto-STROKE compared to SHAM animals (ACC: 357 358 SHAM = 843.89 ± 360.28 ; Opto-STROKE = -66.48 ± 419.70 , unpaired *t-test*) (Figure 1H). To track IL circuit-specific synaptic changes, relevant for sleep modulation and cognition, we 359 quantified IL synaptic contacts onto PV+ cell in the ACC after immunolabelling of GABAergic 360 PV⁺ interneurons. Quantification of mCherry labelled post-synaptic contacts onto 361 immunoreactive PV+ interneurons revealed a dramatic reduction in the number of IL puncta in 362 Opto-STROKE as compared to SHAM animals (IL puncta: Opto-STROKE = 0.60 ± 0.31 ; 363 SHAM = 2.05 ± 0.84 ; unpaired *t-test*) (Figure 1I and J). These results show: (1) specificity of 364 the Opto-STROKE model in targeting restricted subcortical areas; (2) changes in the IL-target 365 regions projection patterns following Opto-STROKE affecting particularly frontal cortices (PL 366 and ACC); and (3) IL postsynaptic contacts on PV+ interneurons in the ACC were decreased 367 368 post-Opto-STROKE.

369

Intralaminar thalamic Opto-STROKE leads to rapid changes in arousal stability at the lesiononset.

372 Previous clinical and fundamental investigations on the implication of thalamic lesions on regards to arousal, led to contrasting results with either loss (3, 4, 55) or no changes in 373 374 arousability (56, 57). Here, we took advantage of the optical fibre technology for better temporal 375 and spatial infarct resolution to investigate the functional consequences of IL Opto-STROKE. A remarkable feature of the Opto-STROKE model is the possibility to track changes from the 376 377 onset (during) of stroke induction (5 min of laser on- Figure 3A) until later phases (here, up to 378 20 days) in un-anesthetised freely behaving animals. Animals were chronically implanted with EEG/EMG electrodes for characterization of their sleep-wake cycles (see methods and Figure 379 2A). Interestingly, we found that IL Opto-STROKE animals presented a decrease in the latency 380 to the first stable NREM sleep (more than 50s long) immediately after stroke induction as 381 compared to SHAM animals (SHAM = 1.07 ± 7.67 sec; Opto-STROKE = -13.55 ± 11.52 sec; 382 unpaired *t-test*) (Figure 2B). Moreover, during the subsequent 24 hrs, sleep-wake macro-383 architecture was fragmented indicated by the increase of 56 +/ 14 % on the number of wake-384

NREM, NREM-wake transitions in OPTO-stroke mice relative to SHAMs (Figure 2C and Suppl. Figure 4A). Overall, analysis of the sleep architecture revealed a significant decrease in NREM sleep (SHAM = 0.57 ± 15.63 %; Opto-STROKE = -21.48 ± 17.66 %; unpaired *t-test*) (Figure 2D) and an increase in wake (SHAM = 6.94 ± 21.87 %; Opto-STROKE = 19.00 ± 19.00 %; unpaired *t-test*) as compared to SHAM animals on the subsequent 24 hours after Opto-STROKE.

To further characterized sleep/wake quality, oscillatory dynamics of cortical activity were 391 investigated after Opto-STROKE induction. Time frequency analysis of the power 5 min before, 392 during and after Opto-STROKE showed a shift towards lower frequencies with a significant 393 394 augmentation in the power of the delta band (δ) (0.5-4.5Hz) following stroke in comparison to SHAM animals (SHAM = 0.10 ± 0.03 ; Opto-STROKE = 0.16 ± 0.03 ; unpaired *t-test*) (Figure 395 2E-F and Suppl. Figure 4B). This echoed a change in δ^2 component (2.75-3.75 Hz) - a medio-396 dorsal thalamic indicator of sleep homeostasis (58) - restricted the frontal EEG cortices 397 (EEG^{FRONT}) (δ2^{FRONT} SHAM = -16.54 ± 33.38 %; Opto-STROKE = 18.96 ± 52 %; unpaired *t*-398 *test*) (Figure 2G), whereas no significant difference was found for the δ 1 component (0.75-1.5) 399 (Suppl. Figure 4B). The differences in $\delta 1$ and $\delta 2$ modulation after IL lesions were found to be 400 persistent over the 20 days of sleep monitoring (Suppl. Figure 4). 401

402 IL Opto-STROKE promotes arousal instability and reduces sleep efficiency.

403 To characterize the semi-chronic effects of IL Opto-STROKE lesions on sleep-wake architecture and sleep oscillations, recordings were followed from day 0-20 from both 404 experimental groups. Analysis was performed in ZT4-8 (light phase) and ZT16-20 (dark 405 phase). Results showed that Opto-STROKE animals exhibited a general and marked acute 406 fragmentation of NREM sleep and wakefulness in both the light and dark periods (number of 407 NREM bouts % change from baseline light phase: SHAM = -6.17 ± 21.76; Opto-STROKE = 408 409 17.58 ± 16.65 ; NREM bouts dark: SHAM = 7.129 \pm 13.395; Opto-STROKE = 42.471 \pm 23.304; wake light phase (%) SHAM = -12.266 ± 18.695 %; Opto-STROKE = 8.858 ± 18.695 %; wake 410 dark phase: SHAM = 1.96 ± 24.57 ; Opto-STROKE = 32.63 ± 24.95 ; unpaired *t-test*) (Figure 411 412 3A-D, and Suppl. Figure 5A), which tended to renormalize over time. Although, percentages 413 of wake and NREM sleep were unchanged during the light period, lower amounts of wakefulness and increase in NREM sleep were present in Opto-STROKE mice during the dark 414 period. REM sleep parameters were unaffected (Suppl. Figure 5 B and C). 415

Further, a long-lasting increase on sleep instability was observed during the dark period following Opto-STROKE (NREM/wake bouts dark phase: SHAM = 0.67 ± 0.09 ; Opto-STROKE = 1.15 ± 0.043; *two-way ANOVA*) (Figure 3E). In contrast, during the light phase, Opto-STROKE animals showed small, but significant, increase in sleep instability at day 0 (NREM/wake bouts SHAM = 0.93 ± 0.075 ; Opto-STROKE = 1.18 ± 0.041; *two-way ANOVA*) 421 (Suppl. Figure 3C). Unstable sleep was accompanied by a high number of wake-NREM-wake 422 transitions (Number of transitions dark phase: SHAM = 7.25 ± 9.27 ; Opto-STROKE = $30.45 \pm$ 423 13.60; light phase: SHAM = 12.63 ± 9.88 ; Opto-STROKE: 27.73 ± 14.34 ; unpaired *t-test*) 424 (Figure 3F, Suppl. Fig 5D). Changes in wake and NREM sleep architecture progressively 425 returned to baseline levels. However, sleep instability remained high over time.

426 The presence of wake and NREM sleep fragmentation during the animals' active (dark) phase 427 suggested potential further changes in arousal related activity. Indeed, spectral analysis of cortical EEG during wake episodes revealed a unique brain state signature featured by high 428 amplitude 10-13 Hz oscillations that was accompanied by a decrease in the muscle tone, which 429 430 was enhanced in Opto-STROKE relative to SHAM animals (Figure 3H-I). This state was further 431 guantified using double-blind visual scoring to identify EEG with high 10-13Hz oscillation and 432 low EMG power at wake-NREM sleep transitions which represented 6.84 ± 2.43 % of the total amount of vigilance states in Opto-STROKE as compared to 2.62 ± 0.29 % in SHAM animals. 433 Remarkably, the quantity of these transitional state remained high across the light cycles for 434 up to 20 days (Figure 3G, Suppl. Figure 5F and Suppl. Table 1). Moreover, additional analyses 435 revealed an increased power in the 10-13 Hz frequency range in Opto-STROKE animals in 436 comparison to SHAM during the duration of the experiment (Normalized power μV^2 dark phase: 437 SHAM = 0.02 ± 0.02 ; Opto-STROKE = 0.02 ± 0.001 ; two-way ANOVA) (Figure 3J and Suppl. 438 439 Figure 6A).

440 Qualitative analysis of wake period indicated an enhancement of slow wave activity (SWA) across the light cycle (Normalized power (μV^2) dark phase: SHAM = 0.15 ± 0.01; Opto-441 STROKE = 0.2 ± 0.024; light phase: SHAM = 0.15 ± 0.02; Opto-STROKE = 0.20 ± 0.03; two-442 way ANOVA) (Figure 3K). Remarkably, SWA changes were also restricted to the frontal EEG, 443 444 as mentioned above for the acute changes (Suppl. table 2-3 and Suppl. Figure 6B). Furthermore, analysis of the wake spectral power during the dark-active phase showed a 445 decrease in the theta (5-9 Hz) in IL Opto-STROKEs (Normalized power (μ V²) SHAM= 0.09 ± 446 447 0.004: Opto-STROKE = 0.07 \pm 0.01: two-way ANOVA) and gamma (30-60 Hz) power in 448 comparison to SHAM control animals (Normalized power (μ V²) SHAM = 0.04 ± 0.002; Opto-STROKE 0.03 ± 0.004; two-way ANOVA) (Figure 3L and M; Suppl. Figure 6C and D). 449

450 Temporal and topographic renormalization of NREM sleep and oscillatory activities after IL
451 Opto-STROKE.

Paramedian strokes with inclusions of the IL have been reported to present a decrease in NREM sleep spindle density in acute phases, with a subset of the patients presenting long lasting spindle deficits (3, 4). Here, we sought to characterize the spindle oscillatory activities from IL Opto-STROKE animals in the acute and semi-acute phases following Opto-STROKE. First, we ran a spectral analysis of cortical EEG^{FRONT} and EEG^{PAR} signals during NREM for both light and dark phase. At day 0, we found a frontal- parietal dissociation of the sigma band (11-16 Hz) presented a decreased in EEG^{FRONT} power in IL Opto-STROKE animals (Normalized power SHAM D0 = $0.06 \pm 0.01 \mu V^{2}$; Opto-STROKE D0 = $0.01 \pm 0.01 \mu V^{2}$; *twoway ANOVA*).

Further, NREM sleep discrete spindle events were detected as previously reported (54). Remarkably, reduction in the spindle density was found in the frontal but not the parietal EEG derivation at day 0 and 5 (Spindle density (spindles/min): SHAM D0 = 2.65 ± 0.24 ; Opto-STROKE D0 = 1.77 ± 0.24 ; D5 SHAM = 2.83 ± 0.24 ; Opto-STROKE = 2.06 ± 0.62 ; *two-way ANOVA*) and to a minor extent during, the dark phase (Figure 4B).

Expression of spindles has been linked to delta activity in humans, and rodents (54, 59, 60) as well as their temporal relationship with the expression of other oscillatory events (Figure 4A and C). Here, we found that in parallel to disrupted frontal spindle power, Opto-STROKE delta power was enhanced anteriorly (Normalized delta power all states light phase: SHAM = -6.01 $\pm 3.58 \ \mu$ V²; Opto-STROKE = 8.37 $\pm 5.27 \ \mu$ V²; dark phase: SHAM = -8.45 $\pm 4.51 \ \mu$ V²; Opto-STROKE = 18.36 $\pm 11.43 \ \mu$ V²; *two-way ANOVA*) (Figure 4C and D). Changes in delta power spontaneously recovered after day 5 post-stroke.

473 IL Opto-STROKE increases pain sensitivity and impaired PFC-dependent working memory.

The IL thalamus has been implicated in the regulation of stress and pain responses (2, 14, 61– 64). To test this, mice underwent behavioural phenotyping for the assessment of locomotion/anxiety, pain threshold, and cognition-related responses (figure 5A). First, using an open field test (OFT), we observed a decrease in exploration (Figure 5B). However, we found no significant difference in the total distance travelled and the time spend in the centre of the open field arena either between SHAM and Opto-STROKE animals post -STROKE (Figure 5C-D), suggesting no impairments in the motor or stress related responses.

Then, to assess changes in pain perception, we used a pain sensitivity test (PT). A mild electric foot-shock test was employed with increasing shock intensity (0.05 mA-0.60 mA; Figure 5 E). Mice with IL Opto-STROKE presented lower pain threshold to shocks of lowest intensity as compared to SHAM animals (mean triplet pain threshold response level to 0.05 mA: SHAM = 0.38 \pm 0.30; Opto-STROKE = 0.80 \pm 0.18; unpaired *t-test*). Yet, no significant changes were found at higher shock intensities between groups (Fig 5F).

Lastly, to address the consequences of IL Opto-STROKE on learning and working memory, we used the forced alternation task in the Y-maze (YM). Mice were habituated, trained, and tested in the YM at day 12, 13 and 14 post-Opto-STROKE (Figure 5H, Suppl. Figure 1C, and Suppl. figure 7A). We found no significant differences in the number of errors during the training sessions or the latency-to-reward between the two groups (Figure 5I and J Suppl. Figure 7D),

suggesting an absence of deficits in learning or motor performances. However, during testing, 492 the total number of errors was significantly higher in Opto-STROKE as compared to SHAM 493 494 animals (Total errors number: SHAM = 2.69 ± 2.84 ; Opto-STROKE = 7.73 ± 1 ; unpaired *t-test*) 495 (Figure 5K). In fact, IL Opto-STROKE mice showed higher number of errors of type 1-3 with a 496 significant lower percentage of correct trials compared to SHAM animals (Error level effect: P < 0.0001; interaction error level x group P < 0.0001, two-way ANOVA) (Figure 5L, Suppl. Figure 497 498 7C and Suppl. Table 4). Note that verification for potential reward-location preference was quantified. No preference was found in both experimental and control groups. (Suppl. Figure 499 7D). 500

501 To gain insights on the relationship between sleep and behaviour-related performances scores 502 in OPTO-stoke mice, we performed a correlation analysis. Pain threshold was found negatively 503 correlated with gamma power at both day 0 and 10 (Pain threshold-gamma D0: r = -0.983 P =0.003: D10: r = -0.951 P = 0.049, Pearson Correlation) and theta power at D10 (Pain threshold-504 gamma D0: r =-0.935, P = 0.003; D10: r = -0.999, P = 0.005, Pearson correlation) (Figure 5G, 505 506 Suppl. Figure 7E). Noteworthy, the number of errors in the YM was negatively correlated with gamma power (r = -0.809, P = 0.001, Pearson correlation) during the acute phase (D0). 507 508 Furthermore, the long-term impairment in spindles had a negative correlation with working memory performance (r = -0.826, P = 0.006, Pearson Correlation) (Figure 5M, Suppl. Figure 509 510 7F). Collectively, these results are supported by previous studies showing the role of the IL in 511 goal-oriented memory consolidation (21, 24, 62, 65-68). Importantly, our findings on the 512 correlation between sleep and memory performance addresses the importance of alterations 513 on sleep oscillations in the level of behavioural symptoms present at the onset and acute phases of stroke. 514

515 Discussion

Our study introduced a novel model of Opto-STROKE in freely behaving mice, allowing the 516 517 investigation of animals' behaviour evolution from acute to semi-acute stroke phase. Here, we focused on subcortical IL Opto-STROKE lesions and related consequences on arousal and 518 cognition. Our findings demonstrate the validity of such model via: (1) anatomical 519 characterization, showing lesions to be limited to the intralaminar thalamus and stroke-related 520 521 changes in inflammation and IL projection patterns (Figure 1 and Suppl. Figure 2 and 3); (2) 522 acute sleep behaviour analysis revealing fast changes in arousability and stability (Figure 2 523 and 3); (3) over-time analysis of sleep architecture and oscillations showing stable fragmentation and increase in transitional states and progressive recovery (Figure 2 and 3); 524 525 (4) study of cognitive impairments, revealing enhanced pain sensitivity and impaired working memory performance in a PFC-dependent task (Figure 5). 526

Current and widely-used stroke rodents models, including the middle cerebral artery occlusion 527 528 (MCAO) and some photothrombotic ischemia, in large require a deep general anaesthesia and targets widespread areas, limiting the relevance of these translational approaches (44). Our 529 530 photothrombotic Opto-STROKE model uses optical fibres (69) to precisely target sub-cortical 531 brain areas and vascular territories in un-anesthetized freely behaving mice, allowing the 532 monitoring of naturally occurring behaviours from stroke induction to acute and semi-chronic 533 stages (up to 20 days). Importantly, plastic events within acute-stroke time window, and concurrent changes in sleep pattern and oscillations are key for intervention and recovery of 534 535 cognitive, motor, and sensory functions (39, 44, 69), further supporting the relevance of our 536 model.

Interestingly, we found that stroke lesions in the IL induced significant changes in the thalamocortical connectivity, and, in particular, in the IL-to- ACC and - PL circuits including synaptic contacts onto parvalbumin-expressing cells (Figure 1G, H, I, J). It is possible that some of IL projections contact other cell types in these areas, however, previous reports have shown that the majority of them project towards inhibitory interneurons (18, 26). Determining the exact contribution of IL-stroke to the changes in local cortical connectivity awaits further investigation.

543 Previous studies have indicated the role of the IL thalamic neurons in regulating arousal states 544 and maintaining sleep stability (22, 61). Consistent with this, our results showed that IL Opto-545 STROKE is immediately followed by a decreased latency of the NREM sleep onset and 546 increased sleep fragmentation, similar to clinical reports (3, 37, 70) (Figure 2B and C). This is accompanied by an overall increase in delta during the acute-stroke phase, with marked 547 increase in the δ^2 component in stroke animals (Figure 2G). These results are congruent with 548 549 previous studies showing increased in δ2 following central median thalamic cell optogenetic 550 inhibition (58), which mimicked post-sleep deprivation effects. Therefore, our results indicate a post-stroke sleep homeostatic need, which could be due to a change of sleep-dependent 551 plasticity in the ischemic circuitry or circuit-related rearrangements, or both. Further 552 553 confirmation of the former includes higher SWA during wakefulness, typically considered as a 554 sign of increased sleep pressure (58). Yet this may also result from the high sleep and wake fragmentation in opto-stroke animals (Figure 3C and D). Overall, these results indicate a high 555 556 level of sleep pressure and decreased ability to sustain arousal in IL stroke animals acutely, with a tendency to renormalize after 10 days post-stroke induction. This renormalization may 557 558 harmonize contrasting results from other works on thalamic lesions and EEG recordings at 559 semi-chronic stages, where no significant changes in the sleep or spectral power were 560 observed (56, 57). Notably, our results provide evidence on the role of the IL in controlling the 561 expression of slow oscillatory events in a topographic specific manner, which leads to their archetypal activity in the frontal cortex in healthy conditions, and to changes in such activity 562 563 post-IL connectivity reorganization following Opto-STROKE.

A key feature of IL Opto-STROKE was an increased in transitional states, characterized by a 10-13 Hz frequency band (*alpha-like* activity) concomitant to a gradual reduction of the EMG power (Figure 3I)(Figure 3G and Suppl. Figure 5F). Interestingly, activity in the alpha-band has been hypothesized to reflect cortical activation, or more precisely, cortical excitation (71–73). For instance, during anesthetized states, the so-called "alpha-anteriorization" leads to a migration of alpha oscillations from the posterior cortex to the frontal cortex, particularly over the prefrontal cortex, in monkeys and humans (74–76).

- 571 At the cellular level, Lőrincz ML et al. (77) demonstrated that a subtype of excitatory thalamocortical neurons fire in burst at alpha frequency, driving inhibitory interneurons. Related to the 572 573 IL Opto-STROKE, it may be that the reduction in IL excitatory connectivity to ACC PV+ cells 574 (Figure 1J) is the underlying mechanism for the featured high 10-13 Hz power and the parallel 575 increase in transitional states. Nonetheless, our results suggest a stroke-related reduction in cortical regulation and weakening of pyramidal neurons inhibition due to lack of IL input to PV+ 576 577 interneurons (78–80), possibly leading to a higher cortical excitation. Notwithstanding early 578 research (86) suggested a link between alpha-band activity to spindle activity and to thalamo-579 cortico-thalamic re-entrant loops (71), it is thought that alpha-band oscillations and spindle 580 oscillations have a strikingly different physiological basis(81). Our results suggested that IL may be important for modulating the alpha band (72, 82). 581
- 582 Concomitant to changes in SWAs topography and alpha power, we found an acute decrease 583 in sigma power and spindles density in IL Opto-STROKE animals (Figure 4). These results are consistent with human studies where reduction of spindles was observed after stroke (3, 4). 584 Here, we reported evidence in favour of IL regulation of spindles and delta expression across 585 the frontal cortex (54, 59, 83-85). Remarkably, we found a fronto-parietal dissociation of 586 587 oscillatory activities where changes spindle density prominently affected the frontal cortices (Figure 4B-C) as reported for other thalamic nuclei such as the reticular thalamic nucleus (86-588 90). Future studies should concentrate on the understanding of how plastic changes occurring 589 590 after stroke are related to changes in neuronal activity of distinct thalamo-cortical networks and 591 their oscillatory activities from stroke progression's acute to semi-chronic phases.
- Previous studies in animals and humans have implicated the medial thalamus in memory and 592 cognition (21, 25, 26, 33, 62, 67, 68, 91–93). Interestingly, in humans other brain disorders -593 594 characterized by strong deficits in cognition - have been often associated with altered PFC 595 excitation including spindles density and changes in the structure and/or activity of the medial 596 thalamus (19, 94–96). These investigations are further supported by the high density of medial 597 thalamic projections to frontal cortical regions, while other thalamic nuclei preferentially regulate activity of more parietal cortices (18, 22, 25, 26, 61, 97). Here, we found a negative 598 599 correlation between frontal spindles density and working memory performance that highlight

the circuit specificity of our Opto-STROKE model and further supports the notion that these 600 two phenomena might be functionally related and dependent on IL-PFC projections integrity 601 602 (Figure 5M). Additional support comes from the found sensory-related deficits induced by IL 603 Opto-STROKE, consisting in lower pain sensitivity threshold (Figure 5E-F), and impairments 604 in recalling sequence of actions necessary to obtain reward and perseveration in a working memory task performed in the YM (Figure 5H-L). Our results further confirmed the importance 605 606 of the IL thalamus in regulating both salient sensory stimuli processing and cognitive performance, possibly due to reduced central median-mediated PV+ inhibition onto pyramidal 607 608 neurons in the PFC (17, 21, 23, 25, 26, 66, 91, 97-99).

609

610 Sleep- and arousal-related distinct oscillations, as slow waves, spindles, and gamma rhythms, has been shown to be beneficial in the recovery from traumatic brain injuries (39, 41, 69, 100-611 103). Moreover, previous studies have shown that sleep-related neural rhythms regulate 612 613 synaptic plasticity and memory consolidation, while their enhancement is beneficial for both neurological and psychiatric improvements (41, 104-106). In this context, our un-614 anaesthetised mini-stroke model will be of particular interest since it enables the anatomical 615 and functional dissection of specific brain areas in variable and complex pathological condition 616 such as stroke. Thus, this study provides important insights about sleep- and arousal-617 dependent circuit activities and its relation to sensory and cognitive processes. Ultimately, it 618 may open new ways of future circuit- and/or region- specific therapeutic strategies and 619 620 improved personalized treatment.

621

622 AUTHOR CONTRIBUTIONS

Author contributions: C.G.H. and I.L. conception and design of research; I.L and M.B.
performed experiments; T.R. wrote and adapted Matlab custom scripts for data analysis, I.L.,
M.B. and C.K. analysed data; C.G.H. and I.L. interpreted results of experiments; I.L. and
C.G.H. prepared figures; I.L and C.G.H drafted manuscript; C.G.H, I.L., M.B. and C.K edited
and revised manuscript; C.G.H. approved final version of manuscript.

628

629 Acknowledgements

We thank the Zentrum for Experimental Neurology for hosting our research and providing with technical support. To Prof Widmer for the use of the fluorescence microscope and the imaging facility of the University of Bern (Mu40). This work and I.L. were supported by University of Bern Interfaculty Research Cooperation "Decoding Sleep" (Gutierrez Herrera, C.); C.K and T.R were supported by the Inselspital University Hospital (Gutierrez Herrera, C), the University of Bern. M.B. work was part of the Biomedical Sciences Master program at the University of Fribourg-University of Bern.

637 **References**

638

- 639 1. Bogousslavsky J, Regli F, Uske A. Thalamic infarcts: Clinical syndromes, etiology, and
 640 prognosis. *Neurology* 1988;38(6):837–837.
- 2. Schmahmann JD. Vascular Syndromes of the Thalamus. *Stroke* 2003;34(9):2264–2278.
- 642 3. Hermann DM et al. Evolution of Neurological, Neuropsychological and Sleep-Wake
 643 Disturbances After Paramedian Thalamic Stroke. *Stroke* 2008;39(1):62--68.
- 4. Bassetti C, Marhis J, Gugger M, Lovblad KO, Hess CW. Hypersomnia following
 paramedian thalamic stroke: A report of 12 patients. *Ann Neurol* 1996;39(4):471--480.
- 5. Danet L et al. Medial thalamic stroke and its impact on familiarity and recollection. *Elife* 2017;6:e28141.
- 648 6. Witte LD et al. Cognitive, affective and behavioural disturbances following vascular 649 thalamic lesions: A review. *Cortex* 2011;47(3):273--319.
- 7. Carrera E, Bogousslavsky J. The thalamus and behavior. *Neurology* 2006;66(12):1817-1823.
- 652 8. Foix, Hillemand. Les artères de l'axe encéphalique jusqu'au diencéphale 653 inclusivement.1925;32.
- 9. H. D. Recherches anatomiques sur la circulation de l'encéphale. *Arch Physiol Norm Pathol*1874;(2):919–957.
- 10. Cover KK et al. Activation of the Rostral Intralaminar Thalamus Drives Reinforcement
 through Striatal Dopamine Release. *Cell Reports* 2019;26(6):1389-1398.e3.
- Merf YDVD, Jolles J, Witter MP, Uylings HBM. Contributions of Thalamic Nuclei to
 Declarative Memory Functioning. *Cortex* 2003;39(4–5):1047–1062.
- 12. Burk JA, Mair RG. Effects of intralaminar thalamic lesions on sensory attention and motor
 intention in the rat: a comparison with lesions involving frontal cortex and hippocampus. *Behav Brain Res* 2001;123(1):49–63.
- 13. Brunzell DH, Kim JJ. Fear Conditioning to Tone, but Not to Context, Is Attenuated by
 Lesions of the Insular Cortex and Posterior Extension of the Intralaminar Complex in Rats. *Behav Neurosci* 2001;115(2):365–375.
- 14. Sun Y et al. Involvement of the Ventrolateral Periaqueductal Gray Matter-Central Medial
 Thalamic Nucleus-Basolateral Amygdala Pathway in Neuropathic Pain Regulation of Rats.
 Front Neuroanat 2020;14:32.
- Minamimoto T, Kimura M. Participation of the Thalamic CM-Pf Complex in Attentional
 Orienting. *J Neurophysiol* 2002;87(6):3090–3101.
- 16. Bordi F, LeDoux JE. Response properties of single units in areas of rat auditory thalamus
- that project to the amygdala. *Exp Brain Res* 1994;98(2):275–286.

- 17. Delevich K, Tucciarone J, Huang ZJ, Li B. The Mediodorsal Thalamus Drives
- Feedforward Inhibition in the Anterior Cingulate Cortex via Parvalbumin Interneurons. J
 Neurosci 2015;35(14):5743--5753.
- 18. Mukherjee A et al. Variation of connectivity across exemplar sensory and associative
 thalamocortical loops in the mouse. *Elife* 2020;9:e62554.
- 19. Kaskie RE, Graziano B, Ferrarelli F. Topographic deficits in sleep spindle density and
 duration point to frontal thalamo-cortical dysfunctions in first-episode psychosis. *J Psychiatr Res* 2019;113:39–44.
- 20. Spellman T, Svei M, Kaminsky J, Manzano-Nieves G, Liston C. Prefrontal deep
 projection neurons enable cognitive flexibility via persistent feedback monitoring. *Cell*2021;184(10):2750-2766.e17.
- 684 21. Bolkan SS et al. Thalamic projections sustain prefrontal activity during working memory 685 maintenance. *Nat Neurosci* 2017;20(7):987--996.
- 686 22. Gent TC, Bandarabadi M, Herrera ACC, Adamantidis AR. Thalamic dual control of sleep 687 and wakefulness. *Nat Neurosci* 2018;21(7):974--984.
- 23. Vertes RP, Hoover WB, Rodriguez JJ. Projections of the central medial nucleus of the
 thalamus in the rat: Node in cortical, striatal and limbic forebrain circuitry. *Neuroscience*2012;219:120--136.
- 691 24. Mandelbaum G et al. Distinct Cortical-Thalamic-Striatal Circuits through the 692 Parafascicular Nucleus. *Neuron* 2019;102(3):636-652.e7.
- 25. Namboodiri VMK et al. Relative salience signaling within a thalamo-orbitofrontal circuit
 governs learning rate. *Biorxiv* 2021;2020.04.28.066878.
- 26. Schmitt LI et al. Thalamic amplification of cortical connectivity sustains attentional control.
 Nature 2017;545(7653):219–223.
- 697 27. Murphy TH, Corbett D. Plasticity during stroke recovery: from synapse to behaviour. *Nat* 698 *Rev Neurosci* 2009;10(12):861–872.
- 28. Biernaskie J, Chernenko G, Corbett D. Efficacy of Rehabilitative Experience Declines
 with Time after Focal Ischemic Brain Injury. *J Neurosci* 2004;24(5):1245–1254.
- 29. Qin L et al. An Adaptive Role for BDNF Val66Met Polymorphism in Motor Recovery in
 Chronic Stroke. *J Neurosci* 2014;34(7):2493–2502.
- 30. Jones TA, Adkins DL. Motor System Reorganization After Stroke: Stimulating and
 Training Toward Perfection. *Physiology* 2015;30(5):358–370.
- 31. Bergmann TO, Born J. Phase-Amplitude Coupling: A General Mechanism for Memory
 Processing and Synaptic Plasticity?. *Neuron* 2018;97(1):10–13.
- 32. Wei Y, Krishnan GP, Bazhenov M. Synaptic Mechanisms of Memory Consolidation
 during Sleep Slow Oscillations. *J Neurosci* 2016;36(15):4231–4247.
- 33. Chen Z, Wimmer RD, Wilson MA, Halassa MM. Thalamic Circuit Mechanisms Link
 Sensory Processing in Sleep and Attention. *Front Neural Circuit* 2016;9:83.

- 34. Tononi G, Cirelli C. Sleep and the Price of Plasticity: From Synaptic and Cellular
- Homeostasis to Memory Consolidation and Integration. *Neuron* 2014;81(1):12–34.
- 35. Astori S, Wimmer RD, Lüthi A. Manipulating sleep spindles expanding views on sleep,
 memory, and disease. *Trends Neurosci* 2013;36(12):738–748.
- 36. Soddu A, Bassetti CL. A good sleep for a fresh mind in patients with acute traumatic
 brain injury. *Neurology* 2017;88(3):226–227.
- 37. Hermann DM, Bassetti CL. Sleep-related breathing and sleep-wake disturbances in
 ischemic stroke. *Neurology* 2009;73(16):1313–1322.
- 38. Cam E, Gao B, Imbach L, Hodor A, Bassetti CL. Sleep deprivation before stroke is
 neuroprotective: A pre-ischemic conditioning related to sleep rebound. *Exp Neurol* 2013;
 247:673–679.
- 39. Facchin L et al. Slow waves promote sleep-dependent plasticity and functional recovery
 after stroke. *J Neurosci* 2020;40(45): JN--RM-0373-20.

40. Zhang Y, Quiñones GM, Ferrarelli F. Sleep spindle and slow wave abnormalities in
schizophrenia and other psychotic disorders: Recent findings and future directions. *Schizophr Res* 2020; 221:29--36.

41. Ladenbauer J et al. Promoting Sleep Oscillations and Their Functional Coupling by
 Transcranial Stimulation Enhances Memory Consolidation in Mild Cognitive Impairment. J
 Neurosci 2017;37(30):7111--7124.

- 42. Prehn-Kristensen A et al. Transcranial Oscillatory Direct Current Stimulation During
- 731 Sleep Improves Declarative Memory Consolidation in Children With Attention-
- deficit/hyperactivity Disorder to a Level Comparable to Healthy Controls. *Brain Stimul* 2014;7(6):793--799.
- 43. Leng Y et al. Sleep duration and risk of fatal and nonfatal stroke. *Neurology* 2015;84(11):1072--1079.
- 44. Duss SB et al. The role of sleep in recovery following ischemic stroke: A review of human
 and animal data. *Neurobiology Sleep Circadian Rhythm* 2017;2:94--105.
- 45. laccarino HF et al. Gamma frequency entrainment attenuates amyloid load and modifies
 microglia. *Nature* 2016;540(7632):230–235.
- 46. Halje P et al. Oscillations in cortico-basal ganglia circuits: implications for Parkinson's
 disease and other neurologic and psychiatric conditions. *J Neurophysiol* 2019;122(1):203–
 231.
- 47. Ocariz M del MS de, Nader JA, Santos JA, Bautista M. Thalamic Vascular Lesions: Risk
 Factors and Clinical Course for Infarcts and Hemorrhages. *Stroke* 1996;27(9):1530--1536.
- 48. Jickling GC, Sharp FR. Improving the translation of animal ischemic stroke studies to
 humans. *Metab Brain Dis* 2015;30(2):461–467.
- 49. Carmichael ST. Rodent models of focal stroke: Size, mechanism, and purpose. *Neurorx*2005;2(3):396–409.

- 50. Herrera ACC et al. Hypothalamic feedforward inhibition of thalamocortical network
 controls arousal and consciousness. *Nat Neurosci* 2016;19(2):290--298.
- 51. Dedic N et al. Chronic CRH depletion from GABAergic, long-range projection neurons in
 the extended amygdala reduces dopamine release and increases anxiety. *Nat Neurosci*2018;21(6):803–807.
- 52. Zhang C-L et al. Protein Kinase A Deregulation in the Medial Prefrontal Cortex Impairs
 Working Memory in Murine Oligophrenin-1 Deficiency. *J Neurosci* 2017;37(46):11114–
 11126.
- 53. Mensen A, Riedner B, Tononi G. Optimizing detection and analysis of slow waves in
 sleep EEG. *J Neurosci Meth* 2016;274:1–12.
- 54. Bandarabadi M et al. A role for spindles in the onset of rapid eye movement sleep. *Nat Commun* 2020;11(1):5247.
- 55. Lin G et al. Paramedian Thalamic Ischemic Infarction: A Retrospective Clinical
 Observation. *Eur Neurol* 2017;77(3–4):197–200.
- 56. Hindman J et al. Thalamic strokes that severely impair arousal extend into the brainstem.
 Ann Neurol 2018;84(6):926--930.
- 57. Fuller P, Sherman D, Pedersen NP, Saper CB, Lu J. Reassessment of the structural
 basis of the ascending arousal system. *J Comp Neurol* 2011;519(18):3817–3817.
- 58. Hubbard J et al. Rapid fast-delta decay following prolonged wakefulness marks a phase
 of wake-inertia in NREM sleep. *Nat Commun* 2020;11(1):3130.
- 59. Klinzing JG et al. Spindle activity phase-locked to sleep slow oscillations. *Neuroimage*2016;134:607–616.
- 60. Latchoumane C-FV, Ngo H-VV, Born J, Shin H-S. Thalamic Spindles Promote Memory
 Formation during Sleep through Triple Phase-Locking of Cortical, Thalamic, and
 Hippocampal Rhythms. *Neuron* 2017;95(2):424-435.e6.
- 61. Mátyás F et al. A highly collateralized thalamic cell type with arousal-predicting activity
 serves as a key hub for graded state transitions in the forebrain. *Nat Neurosci*2018;21(11):1551--1562.
- 62. Barsy B et al. Associative and plastic thalamic signaling to the lateral amygdala controls
 fear behavior. *Nat Neurosci* 2020;23(5):625–637.
- 63. Liebermann D et al. Subjective cognitive-affective status following thalamic stroke. J
 Neurol 2013;260(2):386–396.
- 64. Ferro JM, Caeiro L, Figueira ML. Neuropsychiatric sequelae of stroke. *Nat Rev Neurol*2016;12(5):269–280.

783	65. Prasad JA, MaACCregor EM, Chudasama Y. Lesions of the thalamic reuniens cause
784	impulsive but not compulsive responses. Brain Struct Funct 2013;218(1):85–96.

66. Beck A-K et al. Neuronal activation in the human centromedian-parafascicular complex
 predicts cortical responses to behaviorally significant auditory events. *Neuroimage*

787 2020;211:116583.

67. Keyes PC et al. Orchestrating Opiate-Associated Memories in Thalamic Circuits. *Neuron* 2020;107(6):1113-1123.e4.

68. Saund J, Dautan D, Rostron C, Urcelay GP, Gerdjikov TV. Thalamic inputs to
dorsomedial striatum are involved in inhibitory control: evidence from the five-choice serial
reaction time task in rats. *Psychopharmacology* 2017;234(16):2399–2407.

69. Mohajerani MH, Aminoltejari K, Murphy TH. Targeted mini-strokes produce changes in
 interhemispheric sensory signal processing that are indicative of disinhibition within minutes.
 Proc National Acad Sci 2011;108(22):E183--E191.

796 70. Luigetti M et al. Bilateral thalamic stroke transiently reduces arousals and NREM sleep 797 instability. *J Neurol Sci* 2011;300(1–2):151--154.

71. Purpura DP. Physiological basis of the alpha rhythm P. Andersen and S.A. Andersson.
(Appleton-Century-Crofts, New York, 1968, 235 p., \$12.00). *Electroen Clin Neuro*1969;27(2):222–223.

72. Hughes SW, Crunelli V. Thalamic Mechanisms of EEG Alpha Rhythms and Their
Pathological Implications. *Neurosci* 2005;11(4):357–372.

73. Sauseng P, Klimesch W, Gerloff C, Hummel FC. Spontaneous locally restricted EEG
alpha activity determines cortical excitability in the motor cortex. *Neuropsychologia*2009;47(1):284–288.

74. Tinker JH, Sharbrough FW, Michenfelder JD. Anterior Shift of the Dominant EEG Rhythm
 during Anesthesia in the Java Monkey. *Anesthesiology* 1977;46(4):252–259.

75. Bastos AM, Loonis R, Kornblith S, Lundqvist M, Miller EK. Laminar recordings in frontal
 cortex suggest distinct layers for maintenance and control of working memory. *Proc National Acad Sci* 2018;115(5):1117–1122.

- 76. Cimenser A et al. Tracking brain states under general anesthesia by using global
 coherence analysis. *Proc National Acad Sci* 2011;108(21):8832–8837.
- 77. Lőrincz ML, Kékesi KA, Juhász G, Crunelli V, Hughes SW. Temporal Framing of
 Thalamic Relay-Mode Firing by Phasic Inhibition during the Alpha Rhythm. *Neuron*2009;63(5):683–696.
- 78. Bogart LJ, O'Donnell P. Multiple long-range inputs evoke NMDA currents in prefrontal
 cortex fast-spiking interneurons. *Neuropsychopharmacol* 2018;43(10):1–8.
- 79. Fan Z, Hu H. Medial Prefrontal Cortex Excitation/Inhibition Balance and Schizophrenialike Behaviors Regulated by Thalamic Inputs to Interneurons. *Biol Psychiat* 2018;83(8):630–
 631.
- 821 80. Ferguson BR, Gao W-J. Thalamic Control of Cognition and Social Behavior Via
- 822 Regulation of Gamma-Aminobutyric Acidergic Signaling and Excitation/Inhibition Balance in
- the Medial Prefrontal Cortex. *Biol Psychiat* 2018;83(8):657–669.

- 81. Steriade M, Gloor P, Llinás RR, Silva FHL da, Mesulam M-M. Basic mechanisms of 824 825 cerebral rhythmic activities. *Electroen Clin Neuro* 1990;76(6):481–508.
- 826 82. Saalmann YB, Pinsk MA, Wang L, Li X, Kastner S. The Pulvinar Regulates Information Transmission Between Cortical Areas Based on Attention Demands. Science 827 828 2012;337(6095):753-756.
- 829 83. Geva-Sagiv M, Nir Y. Local Sleep Oscillations: Implications for Memory Consolidation. Front Neurosci-switz 2019;13:813. 830
- 84. Yordanova J, Kirov R, Verleger R, Kolev V. Dynamic coupling between slow waves and 831 sleep spindles during slow wave sleep in humans is modulated by functional pre-sleep 832 833 activation. Sci Rep-uk 2017;7(1):14496.
- 85. Demanuele C et al. Coordination of Slow Waves With Sleep Spindles Predicts Sleep-834 Dependent Memory Consolidation in Schizophrenia. Sleep 2016;40(1). 835 doi:10.1093/sleep/zsw013
- 836
- 86. Fernandez LMJ, Lüthi A. Sleep Spindles: Mechanisms and Functions. Physiol Rev 837 838 2020;100(2):805-868.
- 87. Anderer P et al. Low-resolution brain electromagnetic tomography revealed 839
- 840 simultaneously active frontal and parietal sleep spindle sources in the human cortex. Neuroscience 2001;103(3):581-592. 841
- 842 88. Cox R, Schapiro AC, Manoach DS, Stickgold R. Individual differences in frequency and 843 topography of slow and fast sleep spindles. *Biorxiv* 2017;113373.
- 844 89. Nir Y et al. Regional Slow Waves and Spindles in Human Sleep. Neuron 2011;70(1):153-169. 845
- 90. Niethard N, Ngo H-VV, Ehrlich I, Born J. Cortical circuit activity underlying sleep slow 846 847 oscillations and spindles. Proc National Acad Sci 2018;115(39):201805517.
- 848 91. Mitchell AS, Chakraborty S. What does the mediodorsal thalamus do?. Frontiers Syst Neurosci 2013;7:37. 849
- 92. Deng J et al. The Parabrachial Nucleus Directly Channels Spinal Nociceptive Signals to 850 851 the Intralaminar Thalamic Nuclei, but Not the Amygdala. Neuron 2020;107(5):909-923.e6.
- 93. Rikhye RV, Gilra A, Halassa MM. Thalamic regulation of switching between cortical 852 representations enables cognitive flexibility. Nat Neurosci 2018;21(12):1753–1763. 853
- 94. Schmitt LI, Halassa MM. Interrogating the mouse thalamus to correct human 854 neurodevelopmental disorders. Mol Psychiatr 2017;22(2):183-191. 855
- 856 95. Buchmann A et al. Reduced mediodorsal thalamic volume and prefrontal cortical spindle activity in schizophrenia. Neuroimage 2014;102:540-547. 857
- 96. Ferrarelli F, Tononi G. Reduced sleep spindle activity point to a TRN-MD thalamus-PFC 858 circuit dysfunction in schizophrenia. Schizophr Res 2017;180:36-43. 859

97. Delevich K, Jaaro-Peled H, Penzo M, Sawa A, Li B. Parvalbumin interneuron dysfunction
in a thalamo-prefrontal cortical circuit in Disc1 locus impairment mice. *Eneuro*2020;7(2):ENEURO.0496-19.2020.

98. Meda KS et al. Microcircuit Mechanisms through which Mediodorsal Thalamic Input to
Anterior Cingulate Cortex Exacerbates Pain-Related Aversion. *Neuron* 2019;102(5):944959.e3.

- 866 99. Rikhye RV, Wimmer RD, Halassa MM. Toward an Integrative Theory of Thalamic
 867 Function. *Annu Rev Neurosci* 2018;41(1):1–21.
- Malekmohammadi M, AuYong N, Ricks-Oddie J, Bordelon Y, Pouratian N. Pallidal deep
 brain stimulation modulates excessive cortical high β phase amplitude coupling in Parkinson
 disease. *Brain Stimul* 2018;11(3):607–617.
- 101. Martorell AJ et al. Multi-sensory Gamma Stimulation Ameliorates Alzheimer's Associated Pathology and Improves Cognition. *Cell* 2019;177(2):256-271.e22.
- 102. Fröhlich F, Lustenberger C. Neuromodulation of sleep rhythms in schizophrenia:
- Towards the rational design of non-invasive brain stimulation. *Schizophr Res* 2020;221:71– 875 80.
- 103. Balbi M et al. Gamma frequency activation of inhibitory neurons in the acute phase after
 stroke attenuates vascular and behavioral dysfunction. *Cell Reports* 2021;34(5):108696.
- 104. Chauvette S, Seigneur J, Timofeev I. Sleep Oscillations in the Thalamocortical System
 Induce Long-Term Neuronal Plasticity. *Neuron* 2012;75(6):1105–1113.
- 105. Binder S et al. Transcranial Slow Oscillation Stimulation During Sleep Enhances
 Memory Consolidation in Rats. *Brain Stimul* 2014;7(4):508–515.
- 106. Benington JH, Frank MG. Cellular and molecular connections between sleep and
 synaptic plasticity. *Prog Neurobiol* 2003;69(2):71–101.
- 107. Konsman J-P. The mouse brain in stereotaxic coordinates Second Edition (Deluxe) By
 Paxinos G. and Franklin, K.B.J., Academic Press, New York, 2001, ISBN 0-12-547637-X. *Psychoneuroendocrino* 2003;28(6):827–828.
- 887
- 888
- 889
- 890
- 891
- 892
- 893
- 894

895 Figures

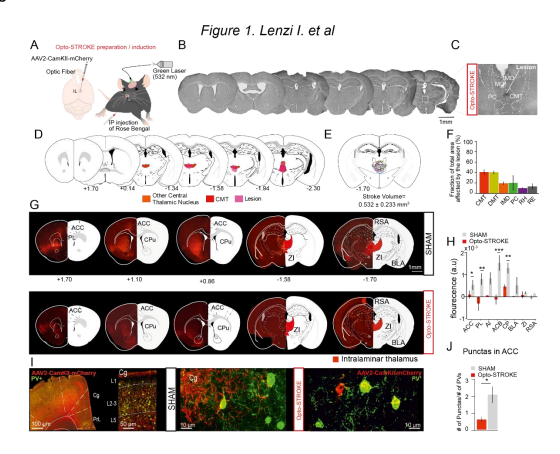


Figure 1. Characterization of Opto-STROKE lesions in the intralaminar thalamus (IL). (A) 896 897 Schematic of animal model of targeted Opto-STROKE in the IL. (B) Representative AP distribution of IL lesion (Cresyl violet). (C) Close-up of IL thalamic nuclei showing the lesion 898 site. (D) Schematic representation of anatomical atlas sections to (C) and coordinates from 899 bregma (107). (E) Overlap of all obtained IL stroke lesions at coordinate -1.75 from bregma 900 and mean lesion volume. (F) Bar graphs of the fraction (%) of IL thalamic nuclei affected by 901 the lesion in Opto-STROKE animals (n = 9). Data are mean \pm SEM. (G) Representative AP 902 903 distribution of sections from SHAM (top) and Opto-STROKE (bottom) injected with AAV2-CamKII-mCherry. (H) Bar graph of the normalized mCherry fluorescence intensity in the target 904 905 regions of IL in SHAM and Opto-STROKE animals (mCherry intensity in area/background fluorescence; SHAM (n = 3) vs Opto-STROKE (n = 4), (I) From left, in order: representative 906 907 image of IL projections (mCherry, red) to prefrontal cortices (Anterior cingulate cortex (ACC) and prelimbic (PL)) and parvalbumin positive neurons (PV+, green); close-up of ACC cortex 908 showing intense IL projections to ACC cortical lavers; close-ups images from SHAM (left) and 909 Opto-STROKE (right) animals showing IL-PV+ synaptic contacts. (J) Bar graph showing the 910 quantification of the number of IL punctas on PV+ interneurons (Number of punctas/number of 911 PV+ interneurons; SHAM (n = 4) vs Opto-STROKE (n = 3), two-way ANOVA with Bonferroni 912 913 post hoc test. Data are mean \pm SEM. Two-way ANOVA with Bonferroni post hoc test; *P < 0.05, ***P* < 0.002, ****P* <0.0002). 914

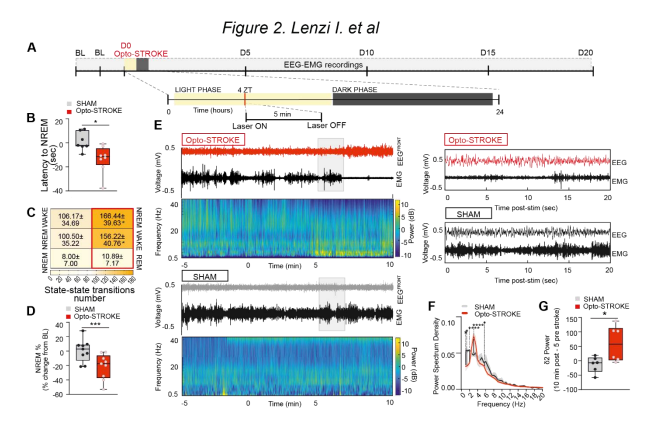


Figure 2. Changes in arousability immediately after IL Opto-STROKE induction. (A) Experimental timeline for acute EEG-EMG recordings (D0). (B) Min-Max box-plots of the summary data of the NREM episode (min. 5 sec) onset latency (SHAM (n = 6) vs Opto-STROKE (n = 6), *P < 0.05, **P < 0.002, ***P < 0.002), unpaired *t-test*. (C). Heat maps of WAKE-NREM-WAKE transitions' level (Colour map light-dark yellow as increasing transition number, SHAM (n = 6) vs Opto-STROKE (n = 9), unpaired *t-test* *P < 0.05, **P < 0.002, **P < 0.002, **P < 0.005, **P < 0.002, **P < 0.005, **P < 0.002, (E) EEG-EMG traces and heat map of frequency analysis over 15 min (5 min prestroke induction; 5 min lase ON; 5 min post-stroke induction). (F) Power spectrum density of the frequency range 0-20 Hz within the time frame 5 min pre-stroke – to – 10 min post-stroke (SHAM (n = 6) vs Opto-STROKE (n = 6), unpaired *t-test*, *P < 0.005, **P < 0.002, ***P < 0.0002. (G) Min-max box-plots showing delta 2 (δ 2) power between time 10 min post and 5 min pre-stroke (SHAM (n = 6) vs Opto-STROKE (n = 6), unpaired *t-test*. *P < 0.05, **P < 0.005, **P < 0.002, ***P < 0.002. All data represents mean ± SEM.

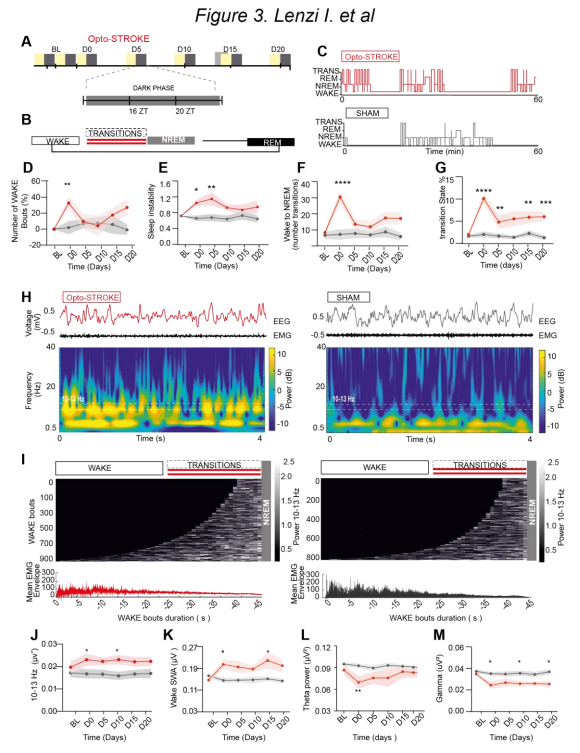


Figure 3. IL Opto-STROKE induces changes in sleep stability and efficiency. (A) Timeline showing sleep recording days and of 12 hours recording during light-dark cycle. (Hours analysed: light: 4-8 ZT; dark: 16-20 ZT). (**B**) Top to bottom: scheme showing natural transitioning between sleep states, with red connecting lines highlighting the WAKE-NREM-WAKE transitional periods enhanced in Opto-STROKE animals. (**C**) Hypnograms showing the increase in manually scored transitional states in Opto-STROKE animals (red) in comparison to SHAM (grey). (**D-G**) From left to right: wake number of bouts (SHAM (n = 9) vs Opto-STROKE (n = 10)) as change from baseline (in %), sleep instability (NREM bouts/WAKE bouts) (SHAM (n = 10) vs Opto-STROKE (n = 10)), transitional states number (SHAM (n = 8) vs Opto-STROKE (n = 11)) and % (SHAM (n = 8) vs Opto-STROKE (n = 11)) over time progression during the dark phase. (**H**) EEG-EMG traces and heatmaps of time-frequency analysis showing increased power in the frequency band between 10-13 Hz in

Opto-STROKE animals during transitional states. (I) Stacked wake episodes at transition to NREM ordered from the shortest (5 sec) to longest (50 sec) in Opto-STROKE (left) and SHAM (right) animals, with respective mean EMG envelop (bottom), showing increase in 10-13 Hz at NREM transition and gradual decrease in EMG power (10-13 Hz power calculated normalizing each animal power within 10-13 Hz to the overall power in overlapping bins of 2 sec). (J-M) From left to right: line plots showing over-time progression of wake 10-13 Hz activity (SHAM (n = 9) vs Opto-STROKE (n = 8)), slow wave activity (SWA) (SHAM (n = 8) vs Opto-STROKE (n = 10)) and gamma power (SHAM (n = 8) vs Opto-STROKE (n = 10)) and gamma power (SHAM (n = 8) vs Opto-STROKE (n = 10)) during the dark active phase. Statistical test. *Two-way ANOVA* with Bonferroni post hoc test was used as statistical test, *P < 0.05, **P < 0.002, ***P < 0.002 were consider significant. Data is represented as mean ± SEM.

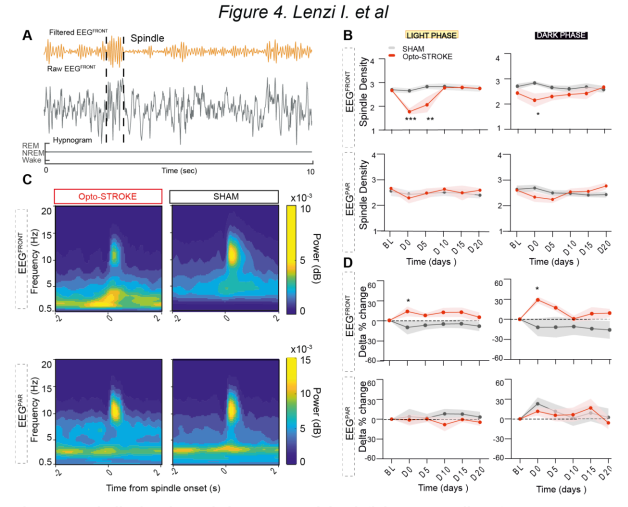


Figure 4. Spindle density and slow wave activity deficits renormalize after D10 post IL Opto-STORKE. (**A**) Representative traces of EEG signal filtered (top) raw (bottom) during NREM sleep for spindle detection. (**B**) Top: line plots showing EEG^{FRONT} sigma power and spindles density during dark phase (SHAM (n = 6) vs Opto-STROKE (n = 7), Bottom: line plots showing EEG^{PAR} sigma power and spindle density during dark phase (SHAM (n = 6) vs Opto-STROKE (n = 7). (**C**) Representative spectrograms showing coupling between spindles and slow wave activity in Opto-STROKE and SHAM animals (Left and right, respectively, upper panel). Bottom: Representative traces showing locking between delta and spindle activity. (**D**) Top: line plots showing increased EEG^{FRONT} delta power (% change from baseline) during both light and dark phase; bottom: line plots showing no change in EEG^{PAR} delta power in light and dark phase in Opto-STROKE animals (SHAM (n = 8) vs Opto-STROKE (n = 9). Two-way ANOVA with Bonferroni post hoc test. Data are mean ± SEM. *P < 0.05, **P < 0.002, ***P < 0.002).

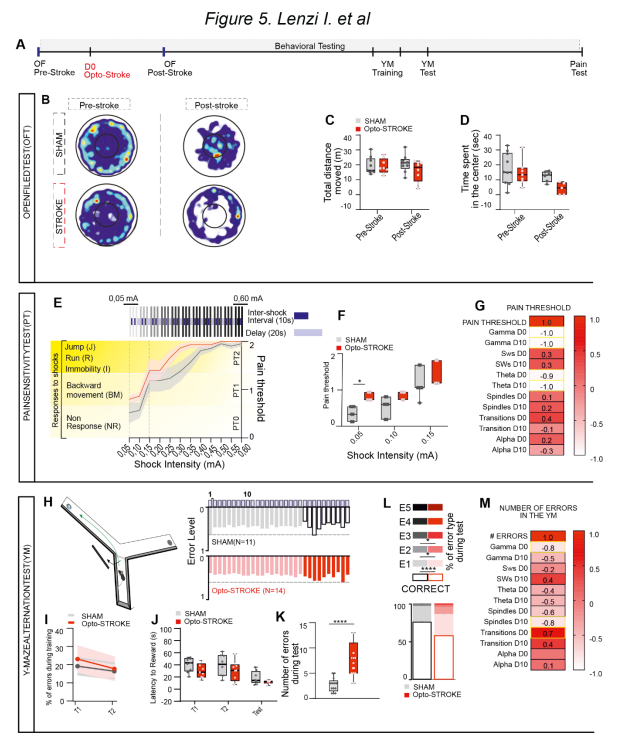


Figure 5. Stress-related responses, pain sensitivity and PFC-dependent working 915 916 memory in IL Opto-STROKE. (A) Timeline of behavioural experimental procedures. Animals were tested before and post- Opto-STROKE induction (D0) in an open field (OFT) arena, and 917 in the forced alteration task in the Y-maze (YM) and pain threshold test (PT) post-Opto-918 STROKE. (B) Representative heatmap showing motor activity of SHAM (upper row) and Opto-919 STROKE (lower row) mice at time pre- (*left*) and post- (*right*) Opto-STROKE. (C-D) Min-max 920 box-plots showing total distance moved (C) and time spent in the centre (D) by SHAM (n = 10) 921 and Opto-STROKE (n = 7) animals at time points pre- and post-Opto-STROKE induction. 922 923 unpaired *t-test*. (E) Schematic representation (top) of the pain sensitivity test. Triplets of shocks (inter-shock interval: 10 secs; delay between triplets: 20 secs) with increasing intensity (0.05 924 925 mA-0.60 mA) were delivered to establish pain threshold in SHAM and stoke animals. Pain 926 threshold to foot shocks was calculated via indexing behavioural responses (Non-response 927 (NR): 0 (PT0); Backward Movements (BM): 1 (PT1); Jump (J), Escape Run (ER) and

Immobility (3) (PT2)) and calculating average response within a triplet of shocks with same 928 929 intensity. Bottom: line graph showing pain threshold in SHAM (n = 7) and Opto-STROKE (n = 7) 5) miceand (I) Boxplot showing difference between groups in level of pain response to shocks 930 of lower intensity (0.05 - 0.15 mA), unpaired *t-test*. (G) Heatmap of the correlation between 931 pain threshold at 0.05 mA and sleep parameters (Pearson correlation, scale bar-1 < r < +1 932 (white= min value; red= max value). (H) Representation of YM set-up (left) and experiment 933 934 structure with bar graph showing distribution of errors in SHAM (upper bar graph) and Opto-STROKE (lower bar graph) animals over sessions' trials. Both training sessions (T1 and T2) 935 and test session were composed by 10 trials interleaved by 30 s intervals. (I) Line graph 936 937 showing percentage of errors accomplished during T1 and T2 by SHAM (n = 13) and Opto-STROKE (n = 15) mice. Two-way ANOVA. (J) Box-plots showing SHAM (n = 13) and Opto-938 STROKE (n = 15) animals' latency to the reward during T1. T2 and test, unpaired *t*-test, (**K**) 939 940 Box-plots showing number of errors made by SHAM (n = 13) and Opto-STROKE (n = 15) 941 animals during testing session. unpaired *t*-test. (L) Stacked bar graphs showing level of error types in testing session (Error 1: E1; error 2: E1; error 3: E3; error 4: E4; error 5: E5) as a 942 943 fraction of cumulative performance (total trials = 100%), with left legend indicating type of errors and significance. Two-way ANOVA, followed by Bonferroni post hoc test. (M) Heatmap of the 944 945 correlation between errors number in the YM and sleep parameters (*Pearson correlation*, scale bar-1 < r < +1 (white= min value; red= max value). Data are mean \pm SEM. *P < 0.05, **P < 946 0.002, ****P* <0.0002. 947