Stereotaxic access to brain areas underneath the superior sagittal sinus (SSS) is notoriously challenging. As a major drainage vessel, covering the whole extension of the sagittal fissure, the SSS impedes direct bilateral access to underlying regions for recording and stimulation probes, drug-delivery cannulas, and injection devices. We now describe a new method for transection and retraction of the SSS in rats, that allows the accurate placement of microinjection devices, or chronic electrode probes, while avoiding hemorrhage and the ensuing deleterious consequences for local structures, animal health, and behavior. To demonstrate the feasibility of this approach we evaluated its consequences acutely during surgery, and thereafter during surgical survival, recovery, behavioral testing, as well as postmortem analysis of histologic impact in the related brain structures of male rats. This method provides a new approach enabling direct access for manipulation and recording of activity in brain areas previously obstructed by the SSS.

Key words: superior sagittal sinus; surgery
Significance Statement

The superior sagittal sinus (SSS) constitutes a physical obstacle for the vertical bilateral stereotaxic access to cortical and subcortical midline structures, often leading to extensive bleeding, damage to the underlying brain tissue and animal death. To overcome these frequent practical difficulties, we have developed and validated an innovative surgical method to transect and retract a small portion of the SSS, exposing the sagittal fissure with negligible vascular or neural damage. Transection of the SSS provides a new approach for manipulation and recording of activity in directly inaccessible midline brain regions, preserving cognitive, physiological, and anatomic integrity.

Introduction

Recording from and manipulating populations of neurons in target brain regions is fundamental for understanding neural function (Cui et al., 2013; Jun et al., 2017; Liang et al., 2017; Kampasi et al., 2018; Pisanello et al., 2018; Steinmetz et al., 2018; Luo et al., 2020). However, bilateral stereotaxic access to cortical and subcortical midline structures has been notoriously difficult because of the obstruction of vertical approach paths by the overlying superior sagittal sinus (SSS). Surgical manipulation of the SSS or insertion of probes in close proximity often leads to ruptures and extensive bleeding, consequently damaging the underlying brain tissue with critical implications for neural function. Bleeding from injuries to the SSS frequently causes accumulation of blood under the meninges, increasing intracranial pressure (Jussen et al., 2017) and inflammation (Quan et al., 2015), and causing ischemic neuronal damage to the underlying cortex (Chen et al., 1991). This damage often extends beyond the underlying cortex, affecting subcortical structures, such as the hippocampus (Jussen et al., 2017). Moreover, the consequences of acute subdural hematomas are frequently more harmful than expected from simple increases in intracranial pressure, with evidence suggesting that the presence of blood constituents aggravates histologic and functional damage (Sasaki and Dunn, 2001; Baechli et al., 2010; Jussen et al., 2017).

It is thus critical to avoid damage and bleeding from the SSS to preserve tissue viability. One technique to avoid or minimize such damage is to target deep midline structures at an angle from a lateral coordinate, instead of directly from above (i.e., vertically). This method has the advantage of being relatively simple and allowing potentially damaging probes or instruments, like electrodes or sharp pipettes, to be kept away from the SSS. However, angled approaches are problematic for reaching deeper structures, such as small midline hypothalamic nuclei, where minor deviations from the intended path lead to gross mistargeting. Additionally, in the case of particular targets (e.g., hypothalamic nuclei), an angled approach is suboptimal for in vivo electrophysiological recordings (Jun et al., 2017; Liang et al., 2017), as depth adjustments and probe designs lead electrodes to cross target structures through a plane of reduced thickness (i.e., orthogonal to the major axis of the region), thus minimizing the number of recorded units per penetration. A second, commonly employed approach, is to dislocate the sinus laterally to expose the target mediolateral coordinate (Wirtshafter et al., 1979). This approach has recently been used to implant optical recording devices for prefrontal cortex and medial entorhinal cortex imaging in head-fixed animals (Low et al., 2014). However, bulky tetrode drives and sensitive silicone probes preclude the simultaneous mobilization of the SSS, and the lowering of probes in the confined space of a craniotomy without increased risks of damage to either. Furthermore, laterally dislocating the SSS only exposes one hemisphere and simultaneous bilateral access is still impeded. Importantly, mobilization per se can lead to injury and bleeding (personal observations), and implanting unilateral probes in close contact to the SSS can cause ruptures at a later point in time if probes need to be lowered postsurgically to sample a new pool of units (e.g., silicone probes).

Direct, unobstructed, vertical access solves many of the challenges inherent to bilateral stereotaxic targeting of deep midline brain regions. Using simplified approach paths would greatly improve the positioning of the recording and stimulation devices, and minimize surgical complications resulting from SSS damage. Here, we report a two-step surgical procedure to transect and retract the SSS, enabling direct bilateral access to the underlying cortex and the longitudinal fissure. First, the SSS is
Materials and Methods

Subjects
A total of 28 male rats (27 Sprague Dawley and one Long–Evans; *Rattus norvegicus*) were used for all procedures. Animals were two to three months old, 350 and 480 g body weight (Charles River Laboratories) at the time of the surgeries. One animal died during the surgery.

Rats were housed in groups of four to five per ventilated double-decker cage and kept under constant experimental conditions: 14/10 h light/dark cycle, 21 ± 0.5°C, 60 ± 10% relative humidity, food and water *ad libitum*.

All the procedures were approved by the Portuguese National Authority for Animal Health (DGAV), as well as by the institute’s animal well-being office.

Surgical procedures

**Transection of SSS**

Materials and step-by-step surgical procedure are described in Detailed Protocol ([https://imm.medicina.ulisboa.pt/intranet/links/view_file/TAS66TLXI/2633](https://imm.medicina.ulisboa.pt/intranet/links/view_file/TAS66TLXI/2633)).

In control animals, without SSS transection, all surgical steps were equal to the experimental group, except the ligation and sectioning of the sinus. In the sham surgery, the suture was threaded under the sinus as described above and removed shortly after, without ligation.

**Microinjection of virus and tracer**

After SSS transection was performed, animals were injected either with a viral construct (AAV8-hSyn-hM4D(Gi)-mCherry) or an anterograde neural tracer [biotinylated dextran amine (BDA)–Texas Red conjugated], using a microinjection control system attached to the stereotaxic frame and a glass micropipette.

The micropipette was filled with mineral oil and placed in the microinjector, after which the virus or tracer were aspirated to fill the tip of the pipette.

Using bregma as a reference, the pipette was carefully positioned over the microinjection coordinates, slowly lowered until reaching the targets [medial preoptic area (MPOA): −0.60 mm AP, +/−0.3 mm ML, −8.4 mm DV; suprachiasmatic nucleus (SCN): −0.60 mm AP, +/−0.3 mm ML, −9.2 mm DV] and left in place for 5 min. The virus/tracer was then delivered (0.5 μl in MPOA or 0.1 μl in SCN), at a rate of 100 nl/min. Micropipette was slowly retracted after waiting 10 min, to avoid contamination of overlaying structures.

Craniotomy was then covered with a single drop of 1.5% agarose at body temperature (~37°C). Skin incision was sutured and Ringer’s lactate was administered for hydration. For postoperative recovery, the animals were placed in a heated cage during the first 24 h. During this period, *ad libitum* food, nutritional gel, and water were provided.

**Hyperdrive implant**

After exposing the skull and identifying stereotaxic landmarks, seven anchor screws were fixed to the skull. An additional screw was placed over the cerebellum to serve as ground. The craniotomy (AP: 1 to −3, ML: 2 to −2) was carefully opened and the SSS transection performed as described above. Mineral oil was then applied to the surface of the brain, and the hyperdrive was carefully lowered and firmly fixed to the skull and to the anchor screws with self-curing dental acrylic (Kerr Tab 2000).

The hyperdrive (adapted from Liang et al., 2017) carried 15 independent octrodes each consisting of eight 12.7-μm nichrome wires (Sandvik). The distance between otrrodes was ~166 μm. A single rectangular octrode bundle (1.46 × 1.27 mm, including the surrounding epoxy used to fix the polyamide tubes) was implanted to bilaterally target the hypothalamus at AP: −0.33 to −1.08, ML: 0.5 to −0.5 and DV: −9.2.

Once the dental acrylic hardened, the octrodes were manually lowered to the desired depth and glued in place. We attached a small “ball” of super glue at the desired octrode length (above the upper end of the polyamide tubes) to serve as a bumber, effectively allowing octrodes to be manually lowered to their target depths. This lowering movement was smooth and unimpeded and we did not observe blood exiting the polyamide tubes. Octrodes were fixed throughout the recording experiment.

The surgical wound was closed, Ringer’s lactate (5 ml) and carprofen (5 mg/kg) were administered subcutaneously, and the animal was left to recover over the period of one week. To prevent infections, the animal received minocycline diluted in the drinking water (100 mg/l) throughout the experiment.

The postmortem extraction of the probe was performed by first ventrally exposing the brain and gently freeing it from the surrounding skull. Octrodes were found to be straight and the hyperdrive was retrieved after careful removal of bone and dental acrylic remains. Although aiming at simultaneous bilateral recordings, postmortem histologic verification revealed unilateral targeting and final coordinates more posterior than expected, most likely because of unlevelled drive and octrode insertion during the surgery.

**Behavioral testing**

To assess potential effects of the sinus sectioning on animals’ locomotor and memory performances, all animals were tested before the surgery, as well as four and eight weeks postsurgery. Rats were handled for 5 d before behavioral tests. After basal behavioral assessment,
animals were randomized between control and experimental groups. All behavioral tests were performed in a sound attenuated room, during the light phase of the circadian cycle, between 9 A.M. [zeitgeber time (ZT)2] and 7 P.M. (ZT12).

**Open-field test**

Locomotor and exploratory behavior were measured as routinely performed in the lab. The test consisted in free exploration of an arena (40 × 40 × 40 cm) for 5 min. Activity was recorded and analyzed using a video-tracking software (Smart 2.5, PanLab) to assess total distance traveled, average speed, resting time and permanence time in peripheral versus central portions of the arena.

**Y-maze test**

Short-term reference memory was assessed in a spontaneous novelty-based spatial preference Y-maze test. The test was performed as a two-trial recognition test in a Y-shaped maze with three arms (each with 35 × 10 × 20 cm), angled at 120° and with opaque walls. Different visual cues were placed at the end of each arm. Allocation of arms was counterbalanced. During the habituation phase (learning trial) rats were placed at the end of the “start” arm, allowed to explore two arms of the maze for 8 min and returned to the home cage. Access to the third arm (“novel” arm) was blocked by an opaque door. After 1 h, the door of the novel arm was removed and animals were placed again in the start arm to freely explore the maze for 5 min (test trial). Rat tracings were recorded using SMART video-tracking software. Preference for the novel arm is considered a measure of short-term reference memory.

**SHIRPA**

SHIRPA is a three-stage protocol for the comprehensive analysis of behavioral phenotype of mice and rats (Rogers et al., 1997).

In order to assess possible adverse side effects of the SSS transection on general behavior phenotype that could be indicative of extensive neural damage, we resorted to the first stage of SHIRPA protocol. This is an adaptation of the Irwin procedure (Irwin, 1968) that consists of standard methods to provide a behavioral and functional profile by observational assessment. The screening of each animal begins by observing undisturbed behavior, looking for manifestation of bizarre or stereotypical behaviors, defects in gait, posture, motor control, coordination, and breathing rate, as well as alterations in coat appearance, piloerection, palpebral closure, lacrimation, and tremor. The observation of spontaneous behavior is followed by a sequence of manipulations, with tail suspension and use of the grid across the width of the cage, to assess positional passivity, limb grasping, irritability, aggression, and visual perception.

Evaluated parameters are scored to provide qualitative assessment of functional phenotype with gross measures of sensory, spinocerebellar, neuropsychiatric, autonomic, and motor functions.

**Histology**

**Pathology**

A total of 10 weeks after SSS transection, rats were euthanized with isoflurane overdose and immediately decapitated. Whole heads were immersion fixed in 10% neutral buffered formalin, followed by 72 h in a decalcifying solution (Surgipath Decalcifier Leica), and then routinely processed for paraffin embedding, sectioned at 4 μm (15–20 sections distancing 100 μm), and stained with hematoxylin and eosin. Lesions were classified by a veterinary pathologist blind to the experimental groups, and scored according to the severity: absent, minimal, mild, moderate, marked, and severe.

To quantify the histologic damage, lesioned area was measured in five serial sections per animal (across 1.5 mm) centered in the zone where the transection was performed and normalized to the whole-brain area.

All slides were digitally scanned using Hamamatsu NanoZoomerSQ, and representative images were obtained using NDP.view2 software (Hamamatsu).

**Anatomical tracing**

Five weeks (viral vector) or 11 d (BDA) postinjection, animals were euthanized as above and transcardially perfused with ~300 ml of PBS followed by ~500 ml of 10% neutral buffered formalin. Brains were gently dissected from the skull to avoid damage to the optic chiasm and SCN, kept in formalin for 24 h at room temperature, then placed in 15% sucrose in PBS until sinking, followed by 30% sucrose in PBS. After sinking, the brains were embedded in gelatin and frozen. Frozen brains were sectioned coronally in 50-μm slices using a cryostat (LEICA, CM3050 S). Fixed brain slices were de-gelatinized in PBS at 37°C for 10 min, mounted in Fluoromount, coverslipped, and left to dry in a dark place. The virus/tracer expression was confirmed using an Axio Observer widefield fluorescence microscope (Zeiss) equipped with an Axiocam 506 mono CCD (Zeiss).

**Octrode tracks**

Four weeks after the hyperdrive implant, the animal was euthanized and perfused. The brain was processed as above, and slices were incubated for 10 min with Hoechst (Hoechst 33342, Thermo Scientific, 12-μg/ml final concentration), washed for 15 min with PBS, mounted in Fluoromount, cover-slipped and left to dry in a dark place. Octrode locations were confirmed using an Axio Observer widefield fluorescence microscope (Zeiss) equipped with an Axiocam 506 mono CCD (Zeiss).

**Analyses**

**Analysis of electrophysiological recordings from hypothalamic nuclei**

Analysis of in vivo electrophysiological signals was performed using the Spikeinterface toolbox (Buccino et al., 2020) together with custom written Python and MATLAB (MathWorks) scripts. Raw data were bandpass filtered between 300 and 6000 Hz and spiking activity was extracted based on four SD threshold crossings. Spike sorting was performed using mountainsort4 (Chung et al., 2017).

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Statistical analysis

All statistical analyses were performed with GraphPad Prism software. Values are presented as dot blots with individual values plus bar with mean ± SEM. Statistical comparisons included two-sided unpaired t test, repeated measures one-way or two-way ANOVA, and two-way ANOVA, followed by a Bonferroni’s or Tukey’s multiple comparison post hoc tests, as specified in the figure legends; p < 0.05 was considered to be statistically significant.

Materials

Please see detailed protocol https://imm.medicinaulisboa.pt/intranet/links/view_file/TAS66TLXI/2633

Results

Transsection of the SSS, overview

We developed a two-step method for safely transecting the SSS with minimal damage to the surrounding vasculature or brain tissue (detailed protocol and materials available at https://imm.medicina.ulisboa.pt/intranet/links/view_file/TAS66TLXI/2633 and minimal increase in surgery time or surgical complexity (~124 min from anesthesia to wound closure in ST animals, as compared with 86 min in sham animals).

Briefly, rats are anaesthetized and securely placed on a stereotaxic frame (Fig. 1A). After exposing the skull, bregma and lambda are identified and used to level the animal’s head (Fig. 1B,C). Using a small drill-bit (Fine Science Tools, catalog #19007-05) the craniotomy window is gradually and carefully opened until a single bone flap is loosened (Fig. 1D). The flap is lifted on one side with fine tweezers and with the help of a small, blunt instrument (e.g., the small tip of a battery-powered cautery; Fine Science Tools, catalog #18010-00) the dura mater is carefully detached from the underside of the bone piece (blunt dissection; Fig. 1E,F). After exposure, the dura mater is punctured with a small needle or sharp tweezers and gradually opened from the lateral borders of the craniotomy toward the SSS (Fig. 1G). Next, the SSS is lifted by the loose dural flaps, and with the help of a ligation aid (Fine Science Tools, catalog #18062-12) a 1.5 absorbable surgical suture (~10 cm long) is threaded under the sinus (Fig. 1H–J). The suture loop is cut and each of the two remaining threads is gently pulled toward the anterior and posterior borders of the craniotomy (Fig. 1K,L). Here, the SSS is securely ligated and blood flow is interrupted (Fig. 1M). Finally, the SSS is cauterized between the ligations and retracted to maximally expose the underlying cortex and the longitudinal fissure (Fig. 1N). The simultaneous bilateral targeting of midline structures is now possible.

In humans, occlusions of the SSS cause severe clinical symptoms and are often fatal (Ferro et al., 2004). Yet in rodents, reports suggest that compensatory mechanisms might minimize the potential harmful impact of such obstructions (Stolz et al., 2011; Wang et al., 2019), as long as cortical venal blood flow is preserved, SSS blockage alone does not alter cerebral blood flow, nor does it cause infarction and hemorrhage (Nakase et al., 1996; Röttger et al., 2005). To examine possible adverse side effects of SSS blockage, we compared the results of a battery of motor, memory function, and general behavior tests, in a group of SSS-transected rats (ST group) with those of age-matched controls (sham group), submitted to all surgical procedures except transection. All animals were tested at baseline (presurgery), and at four and eight weeks postsurgery (for details, see Materials and Methods). Furthermore, to evaluate possible mechanical or ischemic damage, we performed a histopathological study of representative brains from animals in each group.

Transecting the SSS does not impair general behavior

Injuries to the SSS can lead to submeningeal hemorrhages and ischemic lesion to the underlying cortex (Chen et al., 1991; Jussen et al., 2017). To test for possible adverse side effects of the SSS transection on general behavior, and evaluate putative sensorimotor deficits induced by focal lesions, we applied an adapted version of the SHIRPA protocol (Rogers et al., 1997; Hatcher et al., 2001), consisting of an observational screening based on Irwin procedure (Irwin, 1968), to all animals of both groups.

Four weeks after the surgery, there were no qualitative differences between SSS-transected (ST group) and aged-matched sham-operated animals as all parameters scored normal in both groups, indicating that transecting the SSS did not induce behavioral alterations that could be indicative of gross neural damage.

Transecting the SSS does not impair motor behavior

Motor cortical lesions in humans have devastating effects (Heller et al., 1987; Jorgensen et al., 1995). However, rodents seem to recover remarkably well, with little to no lasting impairments in general motor behavior or in dexterous skill execution (Kawai et al., 2015). After confirming that the general behavioral phenotype of ST and sham animals was identical, we asked whether there were alterations in locomotion or exploratory behavior that could be indicative of injury restricted to adjoining primary and secondary motor areas (Neafsey et al., 1986).

We recorded exploratory behavior in both groups, during a 5-min open-field test (Fig. 2A), and quantified various locomotor parameters. We found that transecting the SSS did not alter the total individually traveled distance (repeated measures two-way ANOVA followed by Bonferroni’s multiple comparisons post hoc test, no significant effect of procedure; \( F_{(1,14)} = 0.211, p = 0.662 \); Fig. 2C), nor the total resting time (repeated measures two-way ANOVA followed by Bonferroni’s multiple comparisons post hoc test, no significant effect of procedure; \( F_{(1,14)} = 0.189, p = 0.670 \); Fig. 2D).

Furthermore, exploratory behavior was also unaffected by the transection of SSS, as the permanence in the open-field’s subregions was comparable between groups.
**Figure 1.** Detailed photographs of the surgical procedure. **A**, Induce anesthesia and verify the absence of reflexes. After careful fixation to the stereotaxic frame, scrub the surgical site with antiseptic to avoid contamination. **B**, Make an incision along the sagittal fissure. Expose the skull by scraping periosteum off the bone. **C**, Identify the stereotaxic landmarks (Bregma and Lambda) using the sutures on the skull surface. Validate horizontality of the skull. **D**, Mark the outline of the craniotomy on the skull surface. **E**, Lift the single bone piece using a double hand technique. Use forceps on one hand to slowly lift the bone flap and a cauterizer with the smallest tip attached on the other hand to scrape the dura of the bone as it is gradually lifted. **F**, Open each durotomy at the lateral limit of the craniotomy, next to the bone, and extend it carefully along the anterior and posterior boundaries of the craniotomy until the border of the sagittal sinus. **G**, Slowly lift and fold the bilateral dura flaps over the sinus. **H**, Elevate the attached sinus and thread a suture beneath it, using a vascular ligation instrument. **I**, Use one hand to lift the sinus and the other to slide the tip of the instrument under the sinus, carefully avoiding cerebral veins. Once the instrument’s tip is visible on the opposite side, lower the sinus and carefully thread the suture under it without removing the instrument, stabilizing it very carefully. **J**, Once a 2-3cm loop of suture is threaded, carefully retract the instrument while firmly holding the suture in place at the opposite side of the sinus. **K, L**, Cut the loop in the middle leaving the suture in place, with two strands of thread perpendicular to the sinus. Gently pull one strand towards the anterior limit of the craniotomy and the second towards the posterior limit. **M**, Double ligate the one at the posterior border, followed shortly by the anterior one. **N**, Gently lifting the sinus with one hand, cauterize the mid-point between sutures and extend the severed borders of the sinus posteriorly and anteriorly, exposing the longitudinal fissure. **O**, Cover the whole extension of the
at the time points tested [four weeks after surgery: two-way ANOVA followed by Tukey’s multiple comparisons post hoc test, no interaction ($F_{(2,42)} = 2.002, p = 0.148$); eight weeks after surgery: two-way ANOVA followed by Tukey’s multiple comparisons post hoc test, no interaction ($F_{(2,42)} = 0.295, p = 0.746$); Extended Data Fig. 2-1C].

In essence, ST rats exhibit normal locomotor and exploratory behavior, comparable to sham-operated animals in all behavioral measures acquired (Fig. 2). These results were stable over time, thus excluding the development of lasting deficits resulting from transecting the SSS.

**Transecting the SSS does not impair short-term spatial memory**

Having found that locomotor function was fundamentally preserved, we asked whether memory-dependent behaviors might have been affected by transecting the SSS. Underlying the SSS is the medial mesocortex (anterior cingulate, midcingulate, and retrosplenial), which communicates bi-directionally with the hippocampus and is proposed to be critical for spatial memory and memory-guided behaviors (Ferreira-Fernandes et al., 2019). Considering the proximity of the transected SSS to the anterior cingulate cortex, we assessed the consequences of possible damage to these structures using the Y-maze test for short-term spatial reference memory (Fig. 3A).

ST animals showed a preference for the novel arm similar to sham-operated animals at four and eight weeks after surgery (two-way ANOVA followed by Tukey’s multiple comparisons post hoc test, significant novelty effect; $F_{(1,56)} = 179.9, p < 0.0001$; Fig. 3B). No significant differences were found between the performances of ST and Sham animals after the procedure (Fig. 3). Two-way ANOVA followed by Tukey’s multiple comparisons post hoc test indicates no significant effect of SSS transection ($F_{(3,56)} = 0, p > 0.9999$) nor interaction ($F_{(3,56)} = 1.534, p = 0.2158$).

Furthermore, within subject analysis showed no significant differences in the performance levels of ST animals and Sham animals at the time points tested [four weeks after surgery: two-way ANOVA followed by Tukey’s multiple comparisons post hoc test, no interaction ($F_{(2,42)} = 2.002, p = 0.148$); eight weeks after surgery: two-way ANOVA followed by Tukey’s multiple comparisons post hoc test, no interaction ($F_{(2,42)} = 0.295, p = 0.746$); Extended Data Fig. 2-1C].
across time points (repeated measures one-way ANOVA, $F_{(1.517,10.62)} = 0.192, p = 0.769$; Extended Data Fig. 3-1D).

To exclude possible effects of locomotor activity on exploratory behavior, confounding the latter, we counted and used the number of transitions between arms as an indirect measure of general locomotor activity. Once again, no differences were found between groups at each time point (repeated measures two-way ANOVA followed by Bonferroni’s multiple comparisons post hoc test, no significant effect of procedure ($F_{(1,14)} = 1.202, p = 0.291$) nor interaction ($F_{(2,28)} = 1.819, p = 0.181$); Extended Data Fig. 3-1C).

Overall, transection of the SSS had no transient or long-lasting impact on short-term spatial reference memory.

**Brain tissue viability is preserved after SSS transection**

After confirming that general behavior, motor, and memory functions were fundamentally preserved, we...
performed histopathological analysis of brain samples to test tissue integrity and viability.

A total of 10 weeks after surgery, the skull and meninges in the craniotomy area showed typical features of normal wound healing in both groups (Fig. 4). There was moderate scar tissue formed in the bone and overlying skin, mainly composed of fibroblasts, neovascularization, and macrophages, with occasional hemosiderin accumulation (Fig. 4B,C). In a few cases, this extended through the superficial brain parenchyma in both ST and sham-operated animals.

Histologic damage was assessed in three animals from each group by measuring the lesion area in five serial sections per animal (across 1.5 mm), centered in the zone where the transection was performed, and normalizing it to the whole-brain area. Percentage damage was lower than 0.5% in both groups and not significantly different (sham $0.245 \pm 0.144$, $n=3$; ST $0.249 \pm 0.119$, $n=3$, two-sided unpaired t test, $t_{(4)} = 0.0206$, $p = 0.985$). Furthermore, no correlation was found between histologic damage and behavioral performance ($p > 0.05$; Extended Data Fig. 4-1A).

Since any surgery involving dura mater opening can potentially lead to CSF leakage (Kawai et al., 2014), we inspected ventricular dimensions and confirmed the absence of sizable alterations (Extended Data Fig. 4-1B). At the cellular level, no major lesions were

Figure 4. Transection of SSS does not affect brain tissue viability. Histopathological analysis indicates no extensive neural damage or focal lesion 10 weeks after SSS transection. ST animals are comparable to sham-operated animals in all features analyzed. A, Low magnification of brain, skull and meninges stained with hematoxylin and eosin. Scale bar: 5000 μm. B, Brain and skull magnification. Features of normal surgical wound healing were found within the skull and meninges in the craniotomy area of both groups. Note the scar tissue (scored as moderate). No alterations were found in the cerebral parenchyma. Scale bar: 1000 μm. C, Magnification of scar tissue (scored as moderate), composed mainly of fibroblasts, neovascularization, and macrophages (black arrows) with occasional hemosiderin accumulation (blue arrows). Scale bar: 250 μm. D, High magnification of cerebral parenchyma. No alterations were observed in the neurons and glial cells. Scale bar: 100 μm. Histologic damage is not correlated with behavioral performance and did not involve ventricular alterations (Extended Data Fig. 4-1).
found in the gray and white matter, and no alterations were observed in neurons or glial cells (Fig. 4D).

Histopathological analysis denotes no relevant alterations in ST animals that could be indicative of significant neural damage, or focal lesions. Therefore, SSS transection procedure does not affect viability and integrity of underlying brain tissue.

SSS transection enables microinjections and simultaneous tetrode recordings in deep hypothalamic nuclei

Having established that ST animals have normal motor and memory functions, and that tissue health is preserved, we next validated our method by injecting rats with viral vectors and neural tracers, and implanting octrode drives in deep hypothalamic areas. We used octrodes because of their increased rigidity. However, only four channels (i.e., tetrodes) per octrode were effectively used for recordings.

After transecting the SSS, animals were stereotaxically injected with either a viral vector (AAV8-hSyn-hM4D(Gi)-mCherry, n = 2) in the MPOA or an anterograde neural tracer (BDA–Texas Red labeled, n = 7) in the SCN (Fig. 5A).

Success ratio was defined as the ability to vertically reach our target area without extensive bleeding, damage to underlying brain tissue or causing animal death. Although precise targeting of deep hypothalamic nuclei is dependent on several other variables (i.e., correct determination of coordinates, appropriate head leveling and fixation), this method particularly improves the accuracy of the injections in deep structures by allowing correct determination of the dorsoventral coordinate “zero,” on the brain surface (DV = 0).

Accordingly, five weeks (viral vector) or 11 d (BDA) postinjection, fluorescence microscopy imaging shows extensive viral expression or tracer uptake in the targeted regions (Fig. 5B) as well as anterograde spreading to hypothalamic nuclei (Fig. 5C) in all the nine ST and injected animals. In three animals, the needle tract is visible within the nucleus (Extended Data Fig. 5-1, #1, #3, #4). The remaining animals have staining within
the nucleus consistent with correct targeting, indicating accurate stereotaxic delivery of compounds in the MPOA and SCN. All injections performed using the same virus and tracer showed similar patterns (Extended Data Fig. 5-1).

As transecting the SSS allowed accurate targeting of deep hypothalamic nuclei with minimal increase in surgery time or surgical complexity, we then tested our method for the recording of neural activity in deep midline brain areas.

After SSS transection, we implanted two animals with octrodes. The first animal served as a targeting test and was implanted with a bundle of octrodes dipped in DiI stain not connected to any recording system. One day after the surgery, electrolytic lesions were performed under anesthesia and the animal was killed. Postmortem analysis revealed accurate bilateral targeting of SCN (Extended Data Fig. 6-1A).

The second animal was chronically implanted with a functional hyperdrive (Liang et al., 2017; Fig. 6A) carrying 15 independent octrodes (Fig. 6B) to simultaneously record neural activity (Local Field Potentials (LFP) and single units) from anterior and intermediate hypothalamic nuclei in a freely moving animal. Octrodes were manually lowered to the desired depth and glued in place, remaining fixed throughout the recording experiment.

Using this method, we were able to successfully perform simultaneous single unit and LFP recordings from

**Figure 6.** SSS Transection enables multisite recording of deep midline brain areas. A, Representation of the SLIQ hyperdrive used to chronically record neural activity across multiple regions in freely moving animals. B, Schematic representation of the experimental approach (left). An array of 15 octrodes was chronically implanted to simultaneously record neural activity from different hypothalamic nuclei (right). C, Photograph of ventral brain surface, confirming octrode targeting of deep hypothalamic areas surrounding optical chiasm. D, Schematic of tetrode recording of neural activity from SCN, ReCh, and Arc (left). Two seconds of example LFP recorded simultaneously from SCN, ReCh, and Arc (right). Anatomical location of octrode tips was confirmed by histologic analysis (Extended Data Fig. 6-1B).
SCN, retrochiasmatic area (ReCh), and arcuate nucleus (Arc; Fig. 6D) as confirmed by histologic verification of octrode tracks (Fig. 6C; Extended Data Fig. 6-1B). Although aiming at simultaneous bilateral recordings, postmortem histologic verification revealed unilateral targeting and final coordinates more posterior than expected, most likely because of unleveled drive and octrode insertion during the surgery.

Units were well isolated (Fig. 7C) and the recordings were stable across the duration of a session (Fig. 7A,B). Units B and C started firing when the animal started sleeping or stopped moving inside the sleep-box (Fig. 7A). The number of units varied between recording days. Eight days after the surgery the number of units was relatively low and increased in the second recording session. From this point onwards, we observed a continuous decline in the number of units, as expected in chronic targeting with fixed probes.

Overall, our results show that transecting the SSS is a viable and safe method to gain direct, vertical access to deep brain structures, as well as superficial ones underlying the SSS. Rats with the transected sinus (ST group) show no behavioral alterations and tissue integrity is minimally disrupted, allowing for accurate recording of neural activity without deleterious effects of extensive brain damage or postsurgical hemorrhage.

**Discussion**

We developed a surgical method that enables direct access to brain regions underlying the SSS. Key steps to


this strategy are the ligation and transection of the SSS, enabling retraction of this major venous vessel, exposing the sagittal fissure. We found that, in animals with transected SSSs, the interruption of normal blood flow and surgery-associated mechanical damage to cortical structures in the vicinity of the transected SSS, had no measurable negative impacts. We performed a comprehensive battery of postsurgical testing and found no evidence of compromises to general behavior, short-term memory or motor function. Moreover, histologic examination confirmed that SSS transection did not increase brain damage or inflammation when compared with sham operated animals, indicating preserved tissue viability. Our data show that transected animals are comparable to sham animals in all parameters measured. This might be surprising since the SSS is a vessel of large dimensions, and in fact, in humans, its interruption can have devastating consequences (Ferro et al., 2004). In rodents, it has been suggested that consequences of occlusions to the SSS can be minimized by compensatory mechanisms (Stolz et al., 2011; Wang et al., 2019). SSS blockage alone does not alter cerebral blood flow, nor does it cause infraction and hemorrhage (Nakase et al., 1996; Röttger et al., 2005), when care is taken to minimally impact cortical venal blood flow, as is the case in our methodology. Importantly, this evidence suggests that particular care needs to be taken to preserve the surrounding vasculature to prevent neural damage. Although we cannot rule out acute effects of SSS transection on brain function, all behavior measures, including the SHIRPA (performed at four weeks), which better examines possible neurologic deficits, did not suggest the presence of significant neural damage. Moreover, at 10 weeks after surgery we can only detect typical features of normal wound healing in both groups, with preserved cortical morphology and no alterations in neurons or glial cells. Therefore, it seems unlikely that relevant acute damage, beyond the normal postsurgical condition, is impacting function later.

After establishing the feasibility of this method to expose the sagittal fissure and confirming that animal health and behavior were preserved, we tested its usefulness in two different neural targeting problems. We started by injecting deep midline hypothalamic nuclei with different neural tracers in a group of animals. Our results showed accurate delivery of tracers in the targeted areas with a minimal time cost for resecting the SSS (~2 h from anesthesia to wound closure) and 100% success ratio. Next, we tested its advantages in a particularly challenging implantation surgery: hyperdrive (Liang et al., 2017) targeting deep midline hypothalamic nuclei. Successful implantation allowed us to record single unit activity and LFP simultaneously across target regions, extending the usefulness of our method to complex implantation surgeries targeting midline structures.

Given adequate training and surgical care, we found the procedure to add few risks to surgeries, as demonstrated by the extremely low mortality rate. From a total of 28 rats only one animal died during surgery and none during the recovery period. This demonstrates the feasibility of this approach especially in implantation techniques for in vivo recordings and manipulations of neural activity in freely behaving animals, in which survival is crucial. One factor directly influencing surgical success is the existence of interindividual differences in brain vasculature. Large bridging or cortical veins located in the transected region of the SSS might warrant cancelling the procedure if slight readjustment of the target coordinates is not feasible. The use of either very young or old animals might also constitute possible complications. Particularly in older animals, adhesion of the dura mater to the skull tends to be higher (personal observations), making the extraction of the bone flap to increase the risk of rupturing the SSS in this initial step. However, we believe that with adequate testing and optimization, this method can be adapted to animals of different ages. Also, since our surgical site was relatively anterior and we did not test the procedure at more posterior coordinates where the SSS is thicker, we cannot claim feasibility across the whole SSS extension. Nonetheless, at least one study employing a surgical procedure to expose the pineal gland, located under the confluence of the SSS and the transverse sinuses, reported no surgical complications and a comparably high success rate in the removal of a more posterior section of SSS (Magalin et al., 2009). This strongly suggests that the procedure might be applicable in other sections of the SSS.

Since there is preservation of the dorsal venous system (SSS, transverse sinuses, rostral and caudal rhinal veins) between rats and mice (Xiong et al., 2017) it seems feasible that this method should be applicable to mice, with minimal adaptations. This method seems to be particularly useful for circumstances requiring wide midline unimpeded access. Implanting larger footprint implants at such coordinates requires SSS to be chronically kept away from the midline after implantation (Wirtshafter et al., 1979). Additionally, most drive designs preclude assembling bundles with lateral-to-midline converging angles for bilateral targeting. In these circumstances, dislocating the SSS or attempting angled approaches might not be viable options, could cause extensive bleeding or reduce targeting accuracy. Our method circumvents these difficulties, while simplifying targeting and increasing success rates. We believe that the added benefit of transecting the SSS largely outweighs the increased surgical complexity and duration (~38 min).

Using the procedures we now report, we have demonstrated precise delivery of neural tracers, as well as chronic recordings of neural activity in the midline hypothalamic nuclei during unrestrained behavior. Bilateral access to deep brain regions, particularly using highly sensitive devices/probes with a comparably large footprint, was previously not possible. At a minimal time cost, this method provides direct access for manipulation and recording of activity in brain areas previously obstructed by the SSS.

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