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Hardware Article

A low-cost device for cryoanesthesia of neonatal rodents

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ABSTRACT

Studying the development of neural circuits in rodent models requires surgical access to the neonatal brain. Since commercially available stereotaxic and anesthetic equipment is designed for use in adults, reliable targeting of brain structures in such young animals can be challenging. Hypothermic cooling (cryoanesthesia) has been used as a preferred anesthesia approach in neonates. This commonly involves submerging neonates in ice, an approach that is poorly controllable. We have developed an affordable, simple to construct device – CryoPup – that allows for fast and robust cryoanesthesia of rodent pups. CryoPup consists of a microcontroller controlling a Peltier element and a heat exchanger. It is capable of both cooling and heating, thereby also functioning as a heating pad during recovery. Importantly, it has been designed for size compatibility with common stereotaxic frames. We validate CryoPup in neonatal mice, demonstrating that it allows for rapid, reliable and safe cryoanesthesia and subsequent recovery. This open-source device will facilitate future studies into the development of neural circuits in the postnatal brain.

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Specifications table.

Hardware name	CryoPup
Subject area	Neuroscience
Hardware type	Hypothermic anesthetic device
Open source license	GNU General Public License 3.0 (GPL 3.0)
Cost of hardware	Materials (162 GBP) + PCB (45 GBP) + CNC (96 GBP) = 303 GBP per unit (without shipping costs)
Source file repository	https://doi.org/10.17605/OSF.IO/MXGZ8 https://github.com/FrancisCrickInstitute/Cryopup

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Hardware in context

Systems neuroscience research heavily relies on visualizing and manipulating defined neural circuits in the brain. In rodent models, this is commonly achieved via precise microinjection of chemicals, implantation of cannulas, electrodes and optic fibers, or delivery of viral vectors into the brain using stereotaxic surgery [1,2]. For this purpose, animals are secured into a stereotaxic frame while anesthetics are delivered via intraperitoneal injection or inhalation through a mouth-piece [3].

While this procedure is routinely used in adults, it is suboptimal for neonates, which are increasingly used to study how neural circuits develop postnatally [4–8]: inhalation anesthetics can negatively and lastingly affect the brain of neonatal rodents, in particular areas associated with memory [9–12]. Instead, cryoanesthesia, i.e. anesthesia by deep hypothermia, has been commonly used in such young animals. This is typically carried out by submerging neonates in crushed ice or using simple cooling pads [7,8,13–15]. These approaches, however, do not provide accurate or consistent temperature control, and questions about the safety and appropriateness of hypothermic anesthesia for neonates have thus been raised [16]. A system capable of adjustable temperature control for hypothermic surgeries in adult rodents was recently published, but it is incompatible with stereotaxic surgeries due to its dimensions [17].

We have designed a device (CryoPup) to overcome these limitations. Based on a bidirectionally controlled Peltier element, CryoPup is capable of rapid cooling and heating (temperature range 0 °C to + 42 °C) at ambient temperature (21 °C), thus also acting as a heating pad (see section 7 for safe temperatures for cryoanesthesia and recovery). Due to its compact size, it is fully compatible with modern stereotaxic equipment. We have also designed a 3D printed head holder which facilitates the positioning of neonates during stereotaxic surgeries, thus increasing targeting success. Finally, we show that CryoPup allows for rapid, reliable and safe cryoanesthesia in neonatal mice.

This device thus represents a cost-effective and versatile solution for cryoanesthesia in neonatal rodents.

Hardware description

CryoPup combines a thermoelectric module (Peltier element) which rapidly cools to temperatures suitable for cryoanesthesia of neonatal rodents, with 3D printed components to form a surgery bed. The device consists of a thermoelectric element which pumps heat from a low to a high temperature side and a CPU water cooling system which removes heat from the high temperature side. Peltier-based cooling provides accurate temperature control for cryoanesthesia in a closed loop with a thermocouple confined between the bed top plate and the Peltier element. Temperature is set via an intuitive control panel and the device is compatible with standard stereotaxic frames without further modifications. A 3D printed head holder ensures cranial stability during stereotaxic surgeries. Importantly, CryoPup can seamlessly switch from cooling to heating, and thus acts as a heating pad during recovery, or during the use of other forms of anesthesia in adults. This device can in principle also be equipped with a temperature probe to monitor external or internal (e.g. rectal) temperature.

Main components

The device is composed of a main printed circuit board (PCB), the heat removing system and the bed which encloses a thermoelectric element (Fig. 1a-b).

Electronics

The PCB hosts a microcontroller (Espressif ESP32) which controls the Peltier element and heat exchanger (CPU water cooling loop). Two Maxim integrated thermocouple amplifiers and digital converters (MAX31855KASA + T) acquire data from two Type-K temperature sensors: one in the bed and one external sensor which can be used to monitor experimental animals' temperature. Two power MOSFETs (IRFH8318TRPBF) control the power to the All-In-One liquid cooler (Corsair Hydro H60) and the thermoelectric element (TEC1-12709). A rotary encoder is used to set the desired bed temperature. The desired temperature as well as the two thermocouple temperatures are continuously displayed on a 0.96-inch OLED display (Fig. 5b).

Firmware

CryoPup is capable of both cooling and heating. Rapid cooling is achieved by the thermoelectric element pumping heat over a short distance and by the highly effective heat removal by the closed loop liquid cooler (Fig. 1c). Bed temperature is adjusted combining two bang-bang control algorithms. When cooling (i.e. desired bed temperature is below a predefined ambient temperature) the microcontroller modulates power to the Peltier element using a bang-bang control while the heat removing system is constantly ON. In contrast, if a higher-than-ambient desired temperature is selected, the heat removal system shuts down and a bang-bang control with a small pulse width modulation (PWM) duty-cycle value is initiated. With the thermoelectric element operating in its inefficient regime, heat gradually accumulates on the bed. The oscillations

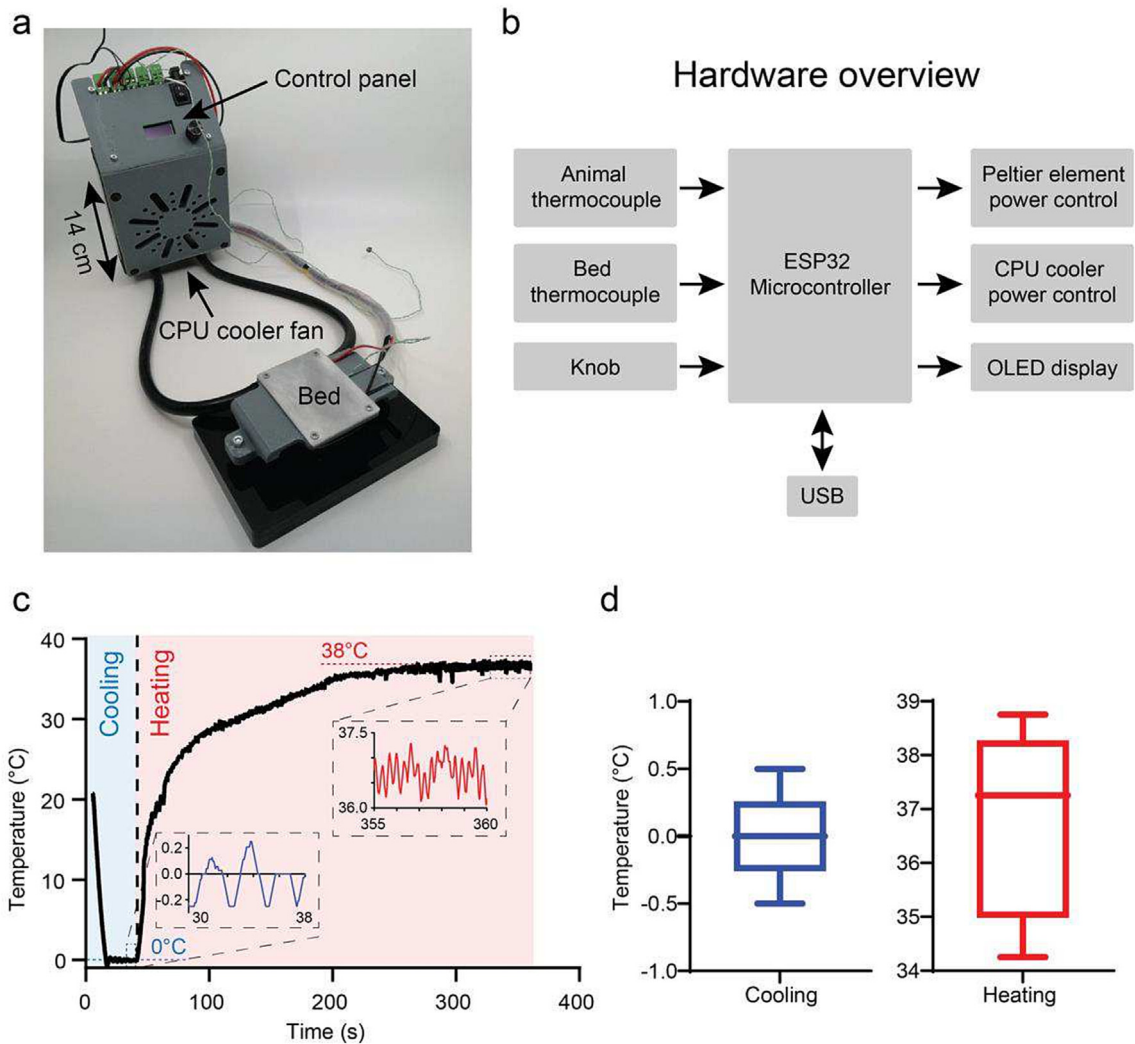


Fig. 1. CryoPup device (a) and schematic control diagram (b). Device performance in a cooling (target temp. 0 °C) and heating (target temp. 38 °C) cycle (c). Temperature variation around the setpoints (d). Data (850 data points each) are from the trial shown in (c). Box-and-whisker plots show median, interquartile range and range. Note that temperatures of 0 °C and 38 °C are not safe for use in live animals.

(ringing) around the set temperature are bigger when warming (± 1.5 °C) than cooling (± 0.5 °C) – thus precise temperature control is ensured during cryoanesthesia (Fig. 1d).

Design files

The design files provide all necessary content to manufacture, assemble and use CryoPup.

Control board: Design and manufacturing files for the PCB as well as Pick and Place (PnP) and Bill of Materials (BOM) files for turn-key PCB manufacturing and assembly.

Peltier holder: Design and manufacturing files for 3D printing the bed frame.

Bed plate: Design and manufacturing files needed for machining the aluminium bed plate.

Electronics holder: Design and manufacturing files for 3D printing the control panel frame which is attached to the heat exchanger and holds the PCB.

Base: Design and manufacturing files for 3D printing or laser cutting the bed base.

Base spacer: Design and manufacturing files for 3D printing or laser cutting the base spacer for the bed.

Head holder: Design and manufacturing files for 3D printing the neonate stereotaxic head holder.

Bill of materials

The components required to build CryoPup, and their costs, are listed in Table 2.

Build instructions

This section describes the build instructions for the PCB, bed base and electronics holder.

Required tools

1. FDM 3D printer (a Zortrax M200 with $200 \times 200 \times 180$ mm building volume was used here)
2. Phillips screwdriver
3. Hex key set
4. Pliers
5. Side cutters / Cable strippers
6. Soldering iron and solder

Manufacturing the custom-designed PCB

The following manufacturing and assembly files for the PCB are available on the OSF repository (link): Gerber files which contain the computer-aided manufacturing files, the bill of materials (BOM) that contains the part number associated with each component and the pick and place file (PnP) which specifies the coordinates on where each component must be placed by the pick and place machine. Following the simple steps outlined below, the board can be reproduced by the manufacturer (JLCPCB) without the need for ordering or soldering components.

1. Download the files from the OSF repository: link.
2. Create customer account on JLCPCB and sign in.
3. Go to JLCPCB instant quote and add Gerber file.
4. Select default settings and preferred PCB color.
5. Under PCB settings, select SMT assembly and indicate quantity (2 or 5).
6. Click next and add the BOM and PnP files.
7. Click next and review if any parts are unavailable. If yes, order from a different supplier (e.g. Mouser, Farnell, Digikey ...).
8. Click next to see the position of each part on the PCB; save to cart and complete payment.

A more detailed document describing the procedure with images can be found on the repository manufacturing files folder (link).

Procuring the remaining components

A comprehensive Bill of Materials can be found in section 4.

Manufacturing of bed base and electronics holder

The four components to be 3D printed can be found in the manufacturing files/mechanical parts folder. The base and the base spacer parts can also be 3D printed but were laser cut from 10 mm acrylic sheets here. M4 and M3 threaded inserts are installed in the base and the Peltier holder parts (Fig. 2a).

Manufacturing the bed

The bed is manufactured from aluminium to ensure good temperature surface homogeneity. This was outsourced to a third-party manufacturer (Hubs), using standard settings (aluminium 6082, no additional treatments). The step files in the Manufacturing files/Mechanical parts folder can be directly uploaded to the instant quote webpage.

Bed assembly

After manufacturing all bed components (Fig. 2a), thermal compound is applied to the CPU cooler surface. Thermal compound improves the heat removal efficiency of the CPU cooler (Fig. 2b). The thermoelectric element is installed on top of the heat sink compound with the writing (hot side) facing the CPU cooler. By convention when the red cable of a thermoelectric element is connected to source positive and the black cable to negative, the ceramic plate on which the cables are soldered on is the hot side. Testing before installation is recommended.

The 3D printed Peltier holder is placed around the Peltier element (Fig. 3a) and attached to the CPU cooler using two 4 mm long M3 screws (Fig. 3b). The bed thermocouple is inserted into groove of the bed aluminium plate (Fig. 3c), the bed plate placed on top of the Peltier element enclosing the thermocouple and secured with 10 mm long countersunk M3



Fig. 2. Required parts for the bed build (a). Application of thermal compound to top of cooler block (b). Peltier element placed on top of thermal compound with cold side facing up (location of red and black wires used as reference) (c). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

screws and nuts (Fig. 3d-e). The bed is attached to the base and base spacer using two M4 screws with a compression spring between the base and the Peltier holder (Fig. 3f-h). These springs allow for easy adjustment of bed height.

Electronics holder assembly

The OLED display and the ON/OFF switch are the final components to be soldered on the board. It is recommended to apply an isolating layer (electric or polyimide tape) to the back of the display before soldering it on the board (Fig. 4a).

The firmware must be uploaded to the board before assembling it in the 3D printed electronics holder since the micro-USB port cannot be accessed afterwards. To upload the firmware on the board, follow the next steps (Windows):

1. Install the Arduino IDE on your computer (link)
2. If necessary, install the SiLabs CP2104 drivers (link)
3. Install the ESP32 package for the Arduino IDE (Version 1.0.6 recommended) (link)
4. Download the sketch from the repository (link)
5. Download libraries:
 - i. ESP32_analog_write (link)
 - ii. Encoder.h (link)
 - iii. Adafruit_MAX31855.h (link)

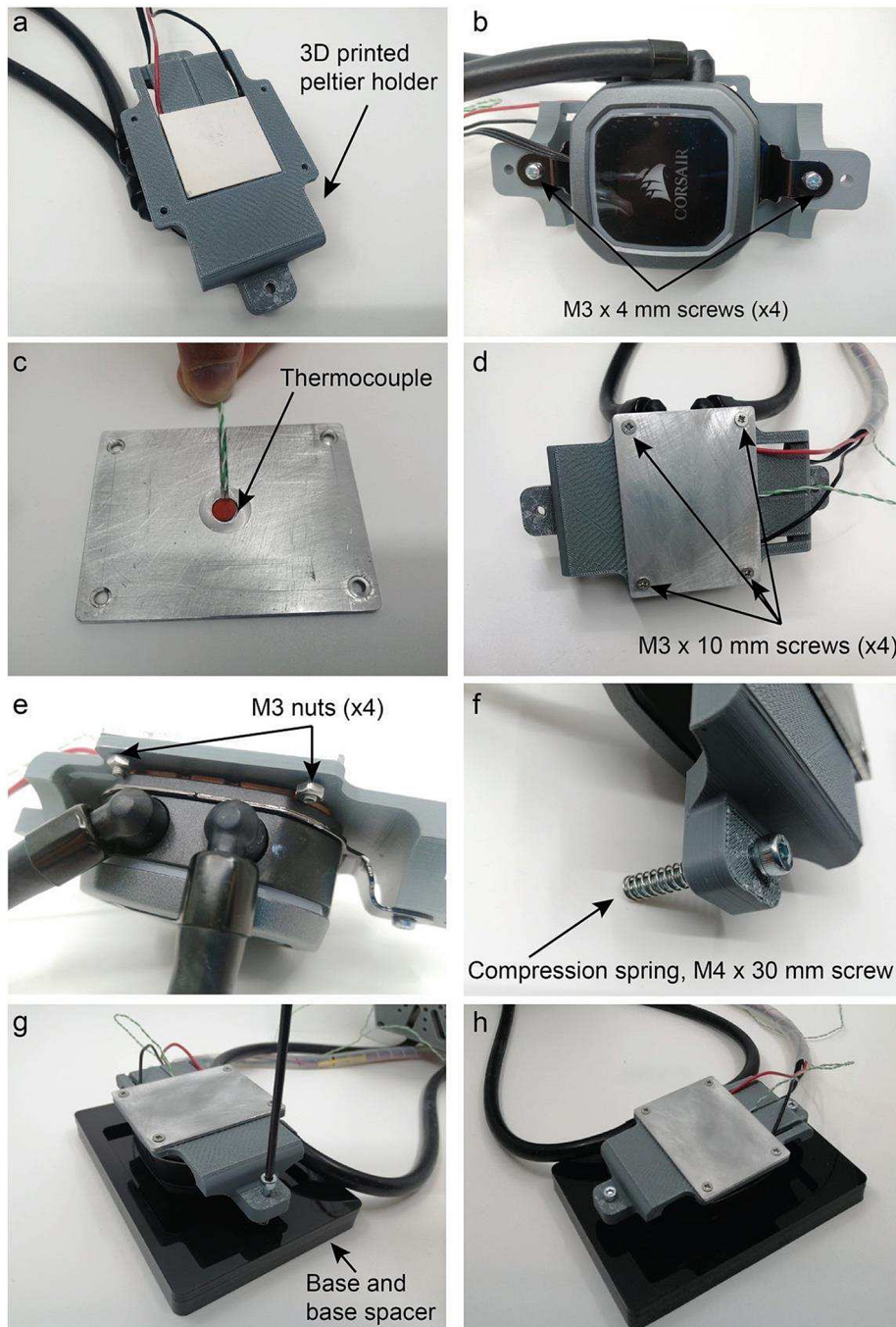


Fig. 3. Installation of the thermoelectric module in the holder (a). Peltier holder attached to CPU cooling block (b). Thermocouple placed between Peltier element and bed aluminium plate (c). Aluminium plate attached to Peltier holder (d). Bed plate and Peltier holder (e). Compression springs provide counterforce to adjust height and to level the bed (f). Bed attached to laser cut base and base spacer (g). Finished assembly (h).

- iv. [Adafruit_GFX.h](#) (link)
- v. [Adafruit_SSD1306.h](#) (link)
5. Connect the CryoPup board to the computer using a micro-USB cable and check “Device Manager” for the name of the COM port.
6. Select the board on Tools -> boards -> ESP32 Arduino -> Adafruit ESP32 Feather
7. Select the board on Tools -> ports -> the COM port listed in Device Manager.
8. Open the sketch *Cold_bed_2.ino* and upload it to the board (link)

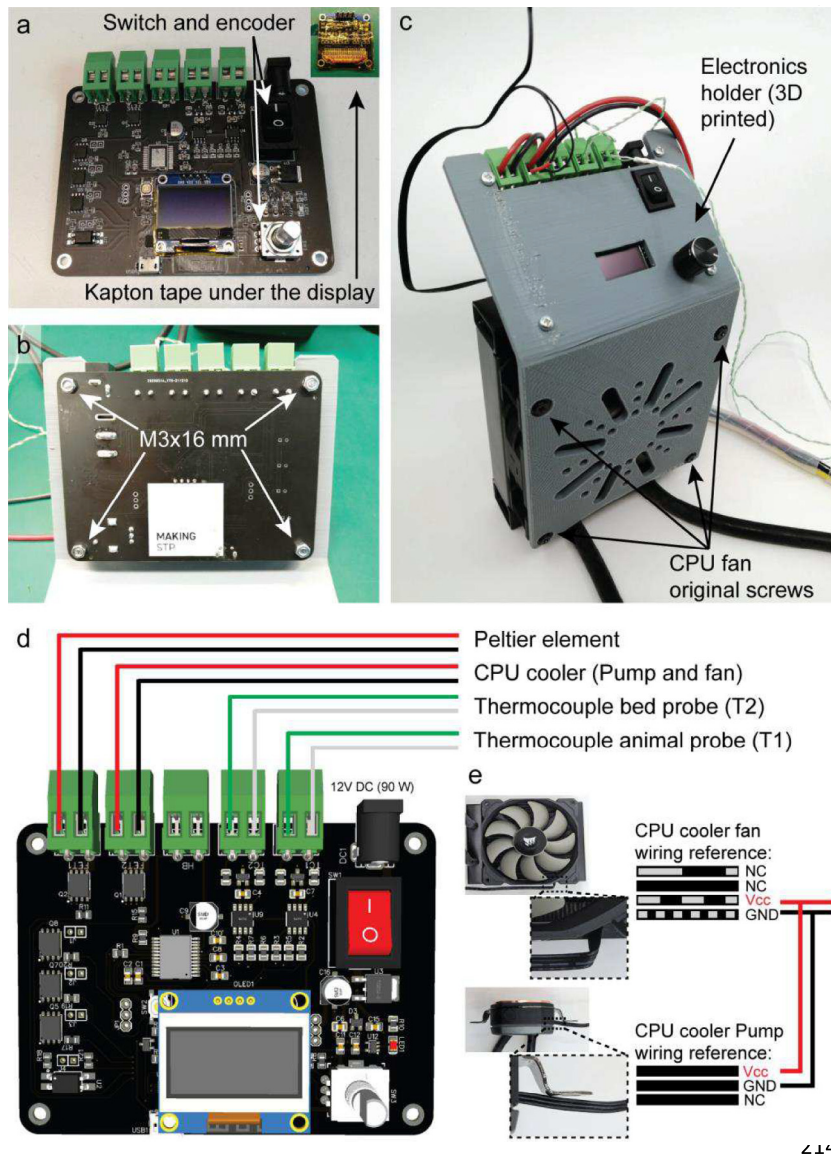


Fig. 4. PCB with components and insulation between back of screen and microcontroller housing. PCB is fixed to the 3D printed electronics holder (b). Electronics holder attached to heat exchanger using the original fan screws (c). Electrical connections to/from PCB screw terminals (d). Heat exchanger fan and pump wire connections (e).

After uploading the firmware, the PCB is attached to the 3D printed electronics holder using four 16 mm long M3 screws and nuts (Fig. 4b). The original fan screws are subsequently used to attach the electronics holder to the fan shroud (Fig. 4c). The frame also serves as a finger (and cable) guard. Finally, all electrical connections (for two thermocouples, CPU fan and pump, and Peltier element) are completed as specified in Fig. 4d. Since the CPU closed loop water cooler consists of two main elements (heat exchanger, i.e., fan and radiator, and pump/cooling block attached with the bed) which have different wiring connectors, they must be re-wired and connected to the same screw terminal as outlined in Fig. 4e.

Using the external thermocouple as a temperature probe (optional)

The external thermocouple can in principle be used as an internal (rectal) probe to monitor animal temperature in adults. For this purpose, the end of the external thermocouple probe can be covered with a 2 cm piece of Tygon[®] E-3603 (3.2 mm ID). This device is implantation biocompatible (USP class VI). However, we have not validated this functionality in animals.

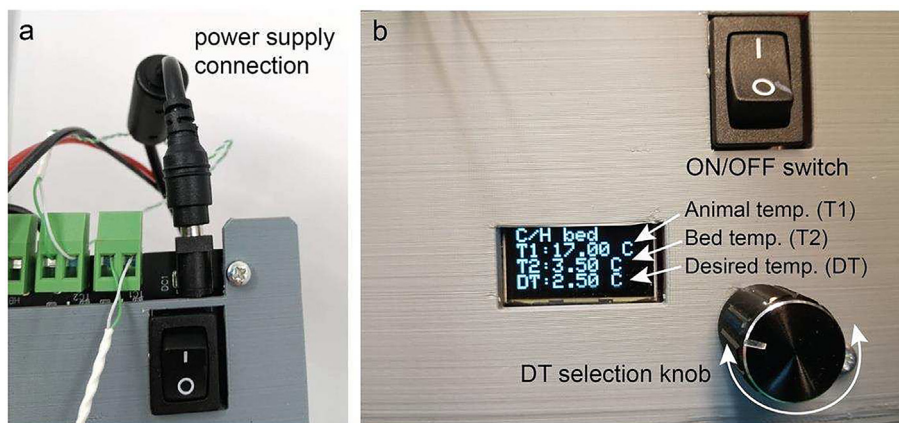


Fig. 5. CryoPup is powered from a 12 V DC 90 W center positive power supply via a standard 2 mm DC barrel jack connector (a). CryoPup control panel (b).

Operation instructions

This section describes detailed steps for operating CryoPup.

1. Connect the 12 V power supply to the barrel jack (Fig. 5a, b).
2. Power up using the ON/OFF switch (Fig. 5b).
3. The display will show three parameters: the temperature from the external thermocouple (T1), the temperature from the bed (T2) and the desired temperature of the bed (DT). DT can be adjusted by the user at any time using the selection knob (default = 0 °C).

Validation and characterization

To test the functionality of CryoPup, we used it to anaesthetise neonatal mice (postnatal day 2–3, $n = 9$) (Fig. 6a). For this purpose, the CryoPup device was slotted into a commonly used stereotactic frame (Kopf, Model 942). A 3D printed head holder (optional, see Table 1) was inserted into the frame's mouse animal adaptor (Model 926, Traditional, Fig. 6a, b). CryoPup was set to a desired temperature (DT) of 2 °C to ensure a bed temperature (T2) of 4 °C. A DT of 2 °C is necessary to achieve a bed temperature of 4 °C due to the temperature difference between the ambient temperature and the temperature measured by the thermocouple, which is directly in contact with the Peltier element. This 2 °C temperature difference is valid for lab environments which are temperature-controlled at 20–22 °C. We chose a target of 4 °C since this temperature reliably induced cryoanesthesia. This is warmer than immersion of pups in ice (0 °C) or in water containing crushed ice (2–3 °C) [13], thus reducing the risk of adverse effects. Mouse pups were gently placed onto the device bed with surgical tape to ensure that the bed was in close contact with the pups' chest/abdomen (Fig. 6a), and the blunt ends of standard ear bars (Kopf #1921) were used to gently fix the head in place (Fig. 6a, b). The pedal withdrawal reflex was tested every 30 s with blunt forceps on alternating feet, and mice were assessed to have a level of anesthesia appropriate for surgery (surgical plane) when this reflex was absent from both feet. This was achieved in 123.8 ± 8.3 s (mean \pm SEM; Fig. 6c, e).

Following induction, pups were maintained at 4 °C for 15 min before DT was set to room temperature (21 °C). The pedal withdrawal reflex was again monitored, and recovery was considered complete when this reflex had returned (140.2 ± 7.6 s, mean \pm SEM; Fig. 6d, e). Animals were subsequently transferred to a standard heated recovery chamber to fully assess their mobility before returning them to the dam.

CryoPup is thus capable of (1) rapidly inducing cryoanesthesia, (2) stably maintaining this state for periods of time compatible with surgical procedures such as intracranial stereotaxic injections (which can be performed in ~ 5 min per coordinate in our experience), and (3) safely returning experimental subjects to wakefulness. Since both the control unit and the fan are potentially significant sources of electrical noise, appropriate grounding and/or shielding might be necessary when performing electrophysiological recordings while using this device.

Notes on safe use.

- (1) Since direct skin contact with cold aluminium carries a potential risk of skin damage, it is essential to ensure that bed temperature does not decrease below 4 °C. A thin protective layer (e.g. surgical drape) can be inserted between the animal and the metal bed to further minimize this risk, but this might result in poorer cooling efficiency. In all cases, confirming optimal temperature settings is essential before using this device on live animals.

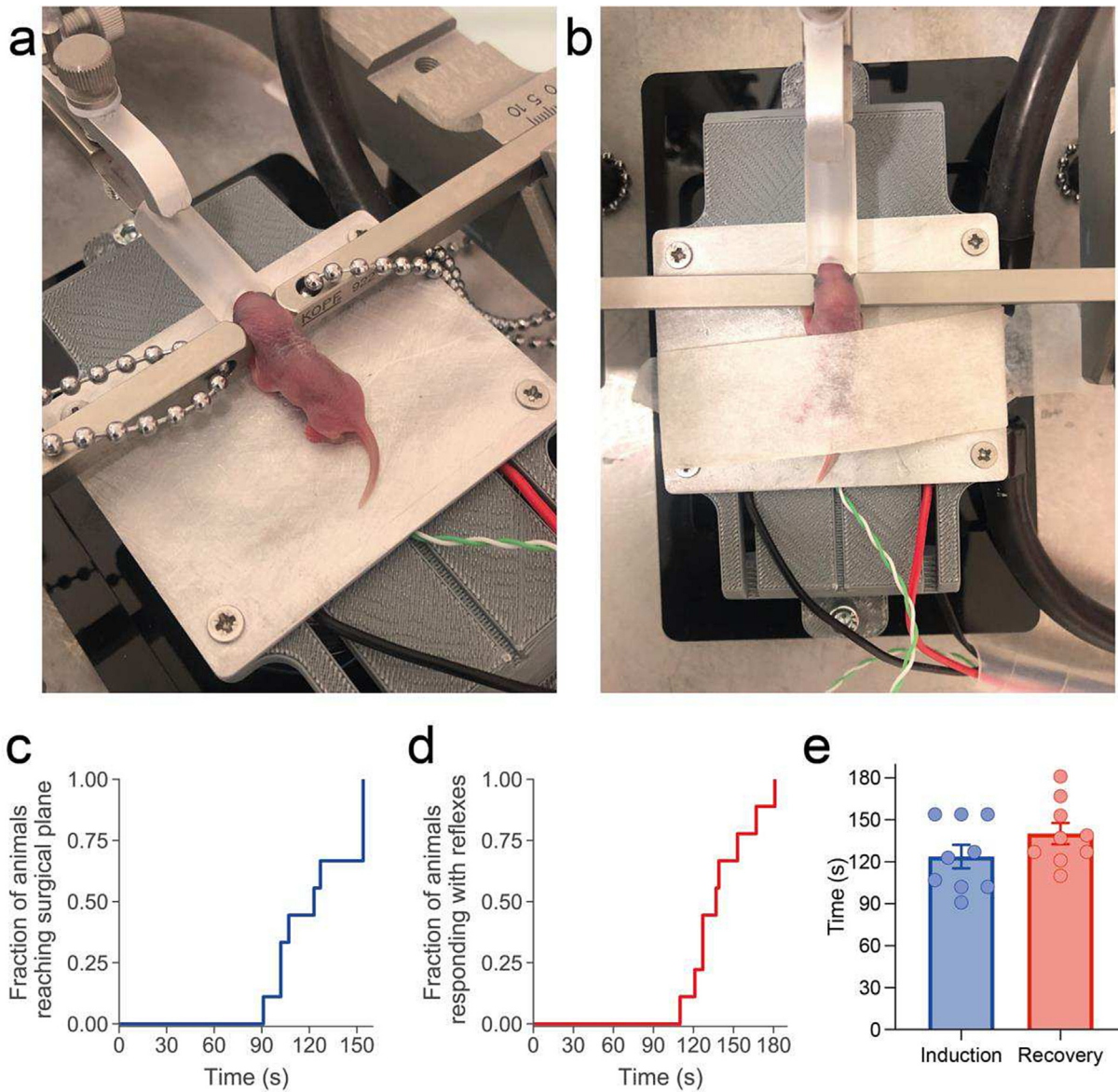


Fig. 6. Example images of neonatal mice during cryoanesthesia on CryoPup (a, b). Survival curves showing the time of induction to reach surgical plane (c), and recovery rate following cessation of cryoanesthesia (d). Average induction and recovery latencies (e). Data presented as mean \pm SEM.

Table 1
List of design files.

Design filename	File type	Open source license	Location of the file
Control board	Easy EDA/ Gerber/csv	GPL-3.0 License	OSF
Peltier holder	Autodesk Inventor/STL/Step	GPL-3.0 License	OSF
Bed plate	Autodesk Inventor/STL/Step	GPL-3.0 License	OSF
Electronics holder	Autodesk Inventor/STL/Step	GPL-3.0 License	OSF
Base	Autodesk Inventor/DXF/STL/Step	GPL-3.0 License	OSF
Base spacer	Autodesk Inventor/DXF/STL/Step	GPL-3.0 License	OSF
Head holder	STL/Step	GPL-3.0 License	OSF

Table 2
Components required for CryoPup.

Designator	Component	Number	Cost per unit (£)	Total cost (£)	Source of materials	Material type
Peltier element	100 W TEC1-12709 Thermoelectric Cooler	1	8.89	8.89	Amazon	Semiconductor / Ceramic
CPU liquid cooler	Peltier 12 V 40 × 40 mm Corsair Hydro H60 Liquid CPU Cooler	1	65.99	65.99	Amazon	Copper / Aluminium / Polymers / Water-Glycol
OLED display	0.96-inch OLED Display SSD1306	1	4.99	4.99	Amazon	Glass / Semiconductor
ON/OFF switch	128 × 64 Pixels RS PRO SPST, Off-On Rocker Switch Panel Mount	1	1.36	1.36	RS Components	Polymer/ Copper
Type K Thermocouple (Bed)	Type K Thermocouple 1 m Length, 6.35 mm Diameter + 200 °C	1	11.93	11.93	RS Components	Copper / Metal Alloys
Type K Thermocouple (Animal)	RS PRO Type K Thermocouple 1 m Length, 2 mm Diameter + 250 °C	1	3.59	3.59	RS Components	Copper / Metal Alloys
Compression Spring	Hilka 79,552,020 Spring Assortment, 200 Piece (5.6 × 17.5 mm).	2	6.00	6.00	Amazon	Steel
Thermal compound	Metal Oxide Thermal Paste, 0.65 W/m-K	1	6.24	6.24	RS Components	Silicone / Zinc Oxide
Power supply 12 V 90 W	RS PRO 12 V DC Power Supply, 90 W, 7.5A, C14 Connector	1	51.23	51.23	RS Components	Polymers / Semiconductors / Metals
M3 screw × 4 mm	RS PRO, Pozidriv Pan Steel, Machine Screw DIN 7985, M3 × 4 mm	2	0.025	0.050	RS Components	Steel
M3 screw × 10 mm countersunk	RS PRO, Pozidriv Countersunk A4 316 Stainless Steel, Machine Screw DIN 965, M3 × 10 mm	4	0.056	0.224	RS Components	Steel
M3 screw × 16 mm	RS PRO, Pozidriv Pan Steel, Machine Screw DIN 7985, M3 × 16 mm	4	0.036	0.144	RS Components	Steel
M4 screw × 30 mm	RS PRO Plain Stainless-Steel Hex Socket Cap Screw, DIN 912 M4 × 30 mm	2	0.6	1.2	RS Components	Steel
M3 nut	RS PRO Stainless Steel, Hex Nut, M3	8	0.03	0.24	RS Components	Steel
M3 threaded insert	RS PRO, M3 Brass Threaded Insert diameter 4 mm Depth 4.78 mm	2	0.13	0.26	RS Components	Steel
M4 threaded insert	RS PRO, M4 Brass Threaded Insert diameter 5.6 mm Depth 7.95 mm	2	0.14	0.28	RS Components	Steel

- (2) While careful application and removal of surgical tape does not lead to any adverse effects in our experience, a small soft tissue pad should ideally be added between tape and animal.
- (3) We have not systematically assessed the maximal safe duration of cryoanesthesia using the CryoPup device. In our experience, ~15 min of cryoanesthesia result in full recovery, as observed previously by others [18]. While longer cryoanesthesia might be required for other applications, the maximal safe duration of cryoanesthesia is poorly documented and thus needs to be carefully established by experimenters.
- (4) For recovery, DT should be set to 21 °C on the CryoPup device.
- (5) We recommend transferring animals to a standard recovery chamber before returning them to the maternal nest in order to fully assess their mobility.

Conclusion

We describe the design for the hardware and software of the CryoPup hypothermic anesthesia device. We provide manufacturing instructions for CryoPup and validate its use as an anesthetic device for neonatal rodents. CryoPup will enable researchers to carry out controlled hypothermic anesthesia on neonatal rodents and can seamlessly integrate into existing stereotaxic frames. This device thus provides a simple and cost-effective manner for stereotaxic surgery in neonatal rodents, facilitating the study of developing brain circuits.

CRediT authorship contribution statement

Bradley B. Jamieson: Conceptualization, Methodology, Validation, Formal analysis, Investigation, Data curation, Writing – original draft, Writing – review & editing, Visualization. **Xavier Cano-Ferrer:** Conceptualization, Methodology, Software, Data curation, Writing – original draft, Writing – review & editing, Visualization. **George Konstantinou:** Conceptualization,

Methodology, Software, Data curation, Writing – original draft, Writing – review & editing, Visualization. **Elisa de Launoit:** Resources, Validation. **Nicolas Renier:** Resources, Validation, Project administration. **Albane Imbert:** Writing – review & editing, Project administration, Supervision. **Johannes Kohl:** Conceptualization, Methodology, Resources, Writing – original draft, Writing – review & editing, Project administration, Supervision.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Ethics statements

Mice were bred and maintained at the specific pathogen-free mouse facility of Paris Brain Institute, with controlled temperature (21 ± 1 °C) and humidity (40-70%), *ad libitum* access to standard mouse chow and water, and on a 12 h:12 h light/dark cycle (lights on at 8 am). Lighting cycles with a controlled rising and decreasing gradient at 8 am/8 pm respectively, simulating natural light. All procedures followed the European legislation for animal experimentation (directive 2010/63/EU).

References

- [1] L. Tenenbaum, A. Chtarto, E. Lehtonen, T. Velu, J. Brotchi, M. Levivier, Recombinant AAV-mediated gene delivery to the central nervous system, *J. Gene Med.* 6 (2004) S212–S222, <https://doi.org/10.1002/jgm.506>.
- [2] A. Cetin, S. Komai, M. Eliava, P.H. Seeburg, P. Osten, Stereotaxic gene delivery in the rodent brain, *Nat. Protoc.* 1 (2006) 3166–3173, <https://doi.org/10.1038/nprot.2006.450>.
- [3] B. Mathon, M. Nassar, J. Simonnet, C. Le Duigou, S. Clemenceau, R. Miles, D. Fricker, Increasing the effectiveness of intracerebral injections in adult and neonatal mice: a neurosurgical point of view, *Neurosci. Bull.* 31 (2015) 685–696, <https://doi.org/10.1007/s12264-015-1558-0>.
- [4] M.R. Zimmer, A.H.O. Fonseca, O. Iyilikci, R.D. Pra, M.O. Dietrich, Functional Ontogeny of Hypothalamic Agrp Neurons in Neonatal Mouse Behaviors, *Cell* 178 (2019) 44–59.e7, <https://doi.org/10.1016/j.cell.2019.04.026>.
- [5] H. Ho, A. Fowle, M. Coetzee, I.H. Greger, J.F. Watson, An inhalation anaesthesia approach for neonatal mice allowing streamlined stereotactic injection in the brain, *J. Neurosci. Methods* 342 (2020), <https://doi.org/10.1016/j.jneumeth.2020.108824> 108824.
- [6] D. Ortolani, B. Manot-Sailliet, D. Orduz, F.C. Ortiz, M.C. Angulo, In vivo Optogenetic Approach to Study Neuron-Oligodendroglia Interactions in Mouse Pups, *Front. Cell. Neurosci.* 12 (2018) 477, <https://doi.org/10.3389/fncel.2018.00477>.
- [7] C.E.J. Cheetham, B.D. Grier, L. Belluscio, Bulk regional viral injection in neonatal mice enables structural and functional interrogation of defined neuronal populations throughout targeted brain areas, *Front. Neural Circuits* 9 (2015), <https://doi.org/10.3389/fncir.2015.00072>.
- [8] P.R. Olivetti, C.O. Lacefield, C. Kellendonk, A device for stereotaxic viral delivery into the brains of neonatal mice, *Biotechniques* 69 (2020) 307–312, <https://doi.org/10.2144/btn-2020-0050>.
- [9] S.E. Maloney, C.M. Yuede, C.E. Creeley, S.L. Williams, J.N. Huffman, G.T. Taylor, K.N. Noguchi, D.F. Wozniak, Repeated neonatal isoflurane exposures in the mouse induce apoptotic degenerative changes in the brain and relatively mild long-term behavioral deficits, *Sci. Rep.* 9 (2019) 2779, <https://doi.org/10.1038/s41598-019-39174-6>.
- [10] C. Zuo, J. Ma, Y. Pan, D. Zheng, C. Chen, N. Ruan, Y. Su, H. Nan, Q. Lian, H. Lin, Isoflurane and Sevoflurane Induce Cognitive Impairment in Neonatal Rats by Inhibiting Neural Stem Cell Development Through Microglial Activation, Neuroinflammation, and Suppression of VEGFR2 Signaling Pathway, *Neurotox. Res.* 40 (2022) 775–790, <https://doi.org/10.1007/s12640-022-00511-9>.
- [11] M.L. Schaefer, P.J. Perez, M. Wang, C. Gray, C. Krall, X. Sun, E. Hunter, J. Skinner, R.A. Johns, Neonatal Isoflurane Anesthesia or Disruption of Postsynaptic Density-95 Protein Interactions Change Dendritic Spine Densities and Cognitive Function in Juvenile Mice, *Anesthesiology* 133 (2020) 812–823, <https://doi.org/10.1097/ALN.0000000000003482>.
- [12] V. Jevtovic-Todorovic, R.E. Hartman, Y. Izumi, N.D. Benshoff, K. Dikranian, C.F. Zorumski, J.W. Olney, D.F. Wozniak, Early exposure to common anesthetic agents causes widespread neurodegeneration in the developing rat brain and persistent learning deficits, *J. Neurosci.* 23 (3) (2003) 876–882.
- [13] C.B. Phifer, L.M. Terry, Use of hypothermia for general anesthesia in preweanling rodents, *Physiol. Behav.* 38 (1986) 887–890, [https://doi.org/10.1016/0031-9384\(86\)90058-2](https://doi.org/10.1016/0031-9384(86)90058-2).
- [14] S.-Y. Chen, H.-Y. Kuo, F.-C. Liu, Stereotaxic Surgery for Genetic Manipulation in Striatal Cells of Neonatal Mouse Brains, *JoVE.* (2018) 57270, <https://doi.org/10.3791/57270>.
- [15] A. De la Rossa, D. Jabaudon, In vivo rapid gene delivery into postmitotic neocortical neurons using iontoporation, *Nat. Protoc.* 10 (2015) 25–32, <https://doi.org/10.1038/nprot.2015.001>.
- [16] K. Herrmann, P. Flecknell, Retrospective review of anesthetic and analgesic regimens used in animal research proposals, *ALTEX - Alternatives to Animal Exp.* 36 (2019) 65–80, <https://doi.org/10.14573/altex.1804011>.
- [17] N. Madrahimov, R. Natanov, A. Khalikov, E.C. Boyle, D. Jonigk, A.-K. Knoefel, T. Siemeni, A. Haverich, Warming and cooling device using thermoelectric Peltier elements tested on male mice, *Lab Anim.* 54 (2020) 443–451, <https://doi.org/10.1177/0023677219873687>.
- [18] C. Janus, T. Golde, The effect of brief neonatal cryoanesthesia on physical development and adult cognitive function in mice, *Behav. Brain Res.* 259 (2014) 253–260.



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