

# Instructions for Use

# PRESSIO<sup>®</sup> MONITORING KIT, PARENCHYMAL WITH BOLT

ICP monitoring: Model PSO-PB ICP and ICT monitoring: Model PSO-PBT

> NT530\_NCE(USA) Rev001 - 2023-12

# Table of contents

1.	Indications		
2.	Contraindications		
3.	Description   4     3.1. Catheters   4     3.2. Accessories   4		
4.	Sterilisation - Decontamination		
5.	Setting-up the Monitoring System		
6.	Preparing, Implanting and Monitoring 5   6.1. Preparing the Catheter Implantation Site 5   6.2. Zeroing the Catheter 6   6.3. Implanting the Catheter 7   6.4. Monitoring, Caring and Nursing of the Patient 9   6.5. Explanting the Catheter 9		
7.	Behaviour During an MRI Examination 9   7.1. General Points 10   7.2. Positioning of the Monitoring Kits and the MRI Support 10   7.3. Positioning of the Monitoring Kits without the MRI Support 11		
7.	Behaviour During an MRI Examination 9   7.1. General Points 10   7.2. Positioning of the Monitoring Kits and the MRI Support 10		
	Behaviour During an MRI Examination 9   7.1. General Points 10   7.2. Positioning of the Monitoring Kits and the MRI Support 10   7.3. Positioning of the Monitoring Kits without the MRI Support 11   7.4. In Vitro Test Conclusions 11   Complications / Side Effects 11   8.1. Infection 11		
8.	Behaviour During an MRI Examination 9   7.1. General Points 10   7.2. Positioning of the Monitoring Kits and the MRI Support 10   7.3. Positioning of the Monitoring Kits without the MRI Support 11   7.4. In Vitro Test Conclusions 11   Complications / Side Effects 11   8.1. Infection 11   8.2. Cerebral Hemorrhage 11		
8.	Behaviour During an MRI Examination 9   7.1. General Points 10   7.2. Positioning of the Monitoring Kits and the MRI Support 10   7.3. Positioning of the Monitoring Kits without the MRI Support 11   7.4. In Vitro Test Conclusions 11   Complications / Side Effects 11   8.1. Infection 11   8.2. Cerebral Hemorrhage 11   Destruction or Return of Product 11		
8. 9. 10.	Behaviour During an MRI Examination 9   7.1. General Points 10   7.2. Positioning of the Monitoring Kits and the MRI Support 10   7.3. Positioning of the Monitoring Kits without the MRI Support 11   7.4. In Vitro Test Conclusions 11   Complications / Side Effects 11   8.1. Infection 11   8.2. Cerebral Hemorrhage 11   Destruction or Return of Product 11   Warranty 11		

#### FIGURE 1: PRESSIO<sup>®</sup> ICP MONITORING KIT, PARENCHYMAL WITH BOLT (MODEL PSO-PB)

A: PRESSIO<sup>®</sup> CATHETER FOR INTRACRANIAL PRESSURE MEASUREMENT WITH ITS DONGLE B: GUIDE STYLET C: BOLT D: ALLEN KEY E: DRILL BIT WITH ADJUSTABLE STOP

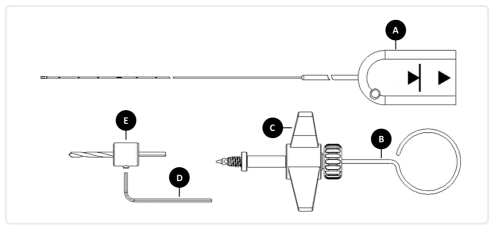
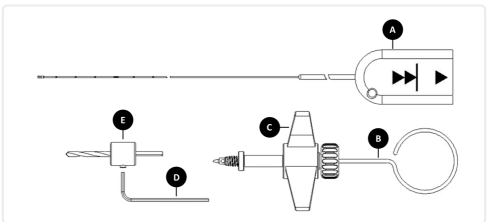


FIGURE 2: PRESSIO® ICP AND ICT MONITORING KIT, PARENCHYMAL WITH BOLT (MODEL PSO-PBT)

A: PRESSIO<sup>®</sup> CATHETER FOR INTRACRANIAL PRESSURE AND TEMPERATURE MEASUREMENT WITH ITS DONGLE B: GUIDE STYLET C: BOLT D: ALLEN KEY E: DRILL BIT WITH ADJUSTABLE STOP



#### Warning

Read the instructions for use carefully before using the Pressio<sup>®</sup> Monitoring Kit, Parenchymal with Bolt (PSO-PB or PSO-PBT).

Read the instructions for use before using the Pressio<sup>®</sup> ICP Monitor (PSO-3000), the Pressio<sup>®</sup> 2 ICP Monitor (PSO-4000), or the Pressio<sup>®</sup> Interface (PSO-IN00).

#### Caution

Federal (USA) law restricts this device to sale by or on the order of a physician.

### 1. Indications

The use of a Pressio Monitoring Kit (hereinafter referred to as Monitoring Kit) is indicated in patients requiring continuous monitoring of intracranial pressure (ICP). Depending on the type of catheter used, the Monitoring Kit can also be indicated in patients requiring continuous monitoring of intracranial temperature (ICT):

- Monitoring Kit, Parenchymal with Bolt (PSO-PB) for parenchymal ICP monitoring,
- Monitoring Kit, Parenchymal with Bolt (PSO-PBT) for parenchymal ICP and ICT monitoring.

The Monitoring Kit is indicated for use by trained personnel of (neuro) intensive care units and neurosurgery departments.

#### Note

The ICP and ICT values indicated on the  ${\rm Pressio}^{\odot}$  Monitoring Systems do not prejudge the condition of the patient.

# 2. Contraindications

Contraindications to the use of a Monitoring Kit are the following:

- Established or suspected infections in the tissues in direct contact with Monitoring System components (meningitis, ventriculitis, septicaemia or bacteraemia), or any infection present in any part whatsoever of the body.
- Patients on anticoagulant therapy or presenting with bleeding diathesis.
- Young children, because the bony plate of their skull is not thick enough to use the fixation with the bolt.

#### Warning

Do not use the Monitoring Kit if there are no trained personnel available to provide continuous observation.

### 3. Description

The Monitoring Kits are made up of:

- a Pressio<sup>®</sup> Catheter (hereinafter referred to as Catheter),
- accessories allowing the Catheter to be implanted and kept in position.

#### 3.1. CATHETERS

Catheters are single-use implantable devices intended to be used only with a Pressio Monitoring System (hereinafter referred to as Monitoring System):

- a Pressio ICP Monitor PSO-3000 or PSO-4000 (hereinafter referred to as Monitor).
- a Pressio Interface PSO-IN00 (hereinafter referred to as Interface).

The Catheters are of the "BF" type.



#### Warning

Catheters are not protected against defibrillation and may be damaged as a result.

Before defibrillation, withdraw the Catheter. If this is not possible, for safety reasons, change the Catheter after defibrillation to continue monitoring.

The Monitoring Kit, **Parenchymal with Bolt PSO-PB** (Figure 1) allows continuous monitoring of ICP.

The Monitoring Kit, **Parenchymal with Bolt PSO-PBT** (Figure 2) allows continuous monitoring of ICP and ICT.

It is designed to be implanted in the cerebral parenchyma.

The two kits can be distinguished visually by their

respective arrows on the Catheter dongle. The presence of a red arrow indicates that it is a Catheter for monitoring both ICP and ICT.

In both kits, the Catheter has a marking every centimeter between 1 and 10 cm from the proximal end, and a marker at 15 cm. The marking at 4 cm is thicker.

The end of the Catheter, opposite the sensor, has a connector to connect the Catheter to a Monitoring System via a Catheter extension cable:

- Catheter extension cable PSO-EC20 to connect a Catheter to a Pressio ICP Monitor or a Pressio Interface.
- Catheter extension cable PSO-EC30 to connect a Catheter to a Pressio 2 ICP Monitor.

The Catheter extension cable transmits the measurement signals from the sensor(s) to the Monitoring System, in the form of analog signals.

The Catheter also exchanges a digital signal with the Monitoring System. This signal contains data, such as sensor calibration, or value and date on which the Catheter was zeroed.

Data is memorised in the Catheter dongle and is thus independent of the Monitoring System used.

#### Note

If a Pressio 2 ICP Monitor is used, the ICP or ICP + ICT data for the first fifteen days of monitoring are recorded in the Catheter memory. Beyond 15 days, no more new data is recorded in the Catheter memory.

This data may be consulted only with a Pressio 2 ICP Monitor by using the "History" function.

#### Specific case for the PSO-PBT

To monitor both ICP and ICT, the Monitoring System must either be a Pressio ICP monitor, software version V2, or a Pressio 2 ICP Monitor.

#### Note

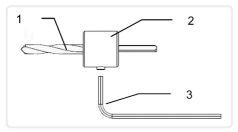
The software version can be identified when powering on the Monitor: the Sophysa logo and software version display on the Monitor touch screen.

If you do not have a Pressio 2 ICP Monitor or a Pressio ICP monitor, software version V2, you can still use the Catheter on one of the other Monitoring Systems (Pressio ICP monitor, software version V1 or Pressio Interface). In this case, only the pressure will be displayed, the temperature will not be displayed. The performances of the pressure sensor will be identical.

### 3.2. ACCESSORIES

The different accessories described below enable the Catheters to be implanted.

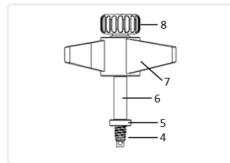
- [1] Drill bit for intracranial access.
- [2] Adjustable stop and its locking screw to set a drilling depth.
- [3] Allen key to set the position of the adjustable stop.



The bolt enables the introduction and fixation of the Catheter in the parenchyma.

- [4] Screw thread to be inserted into the burr hole.
- [5] Spacer ring, mounted on the bolt, to adjust the seating depth for the bolt, depending on the thickness of the skull bone of the patient.
- [6] Bolt shoulder.
- [7] Wing nut to help screw the bolt into the skull bone.
- [8] Tightening nut to fix the Catheter at the required depth in the parenchyma.

4: SCREW THREAD 5: SPACER RING 6: BOLT SHOULDER 7: WING NUT 8: TIGHTENING NUT



The guide stylet is inserted into the bolt (nut [5] unscrewed). It pierces the dura mater to facilitate the introduction of the Catheter.



A sterile disposable drill (PSO-DR) to perforate the skull is also available to order. It can be used in combination with the drill bit supplied in each Monitoring Kit.

# 4. Sterilisation - Decontamination

The Monitoring Kits are packed individually in double peeloff, sterile, pyrogen-free packaging. They are sterilized with ethylene oxide.

#### Warning

Do not use the products if the sterile packaging is open or damaged, or if the expiry date has passed.

This product is **for single use only**. It is intended to be used once only for a single patient.

Do not re-sterilize or re-use after opening the packaging and/or after explantation.

Resterilization may damage the product, which could cause lesions to the patient.

Reusing this device may change its mechanical or biological features and may provoke failure, or the development of allergic reactions or bacterial infections.

#### Note

Sophysa cannot be held responsible for the performance of any product that has been re-sterilized, or for any complications which might result from this.

## 5. Setting-up the Monitoring System

#### Precaution

The Monitoring System should only be used by trained personnel.

- Power on the Monitoring System.
- Connect the Catheter extension cable to the Monitoring System.

# 6. Preparing, Implanting and Monitoring

#### Warning

Read the instructions for use of the Monitoring System in use before using a Monitoring Kit.

#### Caution

The implantation of the Catheter should be performed **immediately** after the Catheter is zeroed. Therefore, prepare the Monitoring System and the implantation site **before** zeroing the Catheter.

Do not perform the implantation of a Catheter without having a replacement Monitoring Kit available in case it is required.

Only connect Catheters to calibrated Monitoring Systems.

# 6.1. PREPARING THE CATHETER IMPLANTATION SITE

Observe aseptic neurosurgical techniques to prepare the Catheter implantation site and the insertion of the bolt.

Surgeons will perform the most appropriate technique depending upon their experience and the clinical status of the patient.

Use the insertion accessories supplied by Sophysa in the Monitoring Kit.

#### 6.1.1. Choice of the Implantation Area

As an example, the right and left prefrontal areas are the main implantation areas. These areas enable the patient to turn his/her head from one side to the other whilst remaining in decubitus without interfering with the intracranial pressure surveillance function.

In most cases, the incision is made behind the hairline, which is acceptable from an aesthetic point of view.

#### 6.1.2. Intracranial Access

Once the implantation site has been chosen, the area is shaved and prepared aseptically. A local anesthetic is applied in the incision area, generally 2 to 3 centimeters in front of the coronal suture on the mid-pupillary line.

- 1. Make an incision of about 1 cm down to the bone.
- 2. Expose the bony plate and perform haemostasis on the wound edges.
- Loosen the locking screw with the Allen key to set the position of the adjustable stop on the drill bit, depending upon the drilling depth chosen.
- 4. Once the adjustable stop is correctly positioned, retighten the locking screw.

#### Warning

Incomplete tightening of the locking screw will prevent the adjustable stop from playing its role, with the risk of drilling too deeply.

- 5. Fix the helical drill bit to a drill.
- Perforate the internal and external skull plates avoiding any possibility of a parenchymal injury when passing through the internal plate.

#### Warning

Make sure to make a completely round hole with the drill bit.

Do not move the shaft of the drill during perforation, as this could affect the hold of the bolt in the burr hole.

- 7. Having crossed the internal plate, remove the drill bit.
- 8. Irrigate the burr hole with sterile saline solution and remove the debris.

### 6.1.3. Putting the Bolt in Place

Screw the bolt into the skull using the wing nut. The spacer ring may be used to reduce the implantation depth of the bolt, otherwise remove and discard it.

#### Precaution

Make sure the bolt is not screwed too deeply. The bolt shoulder (8) and its spacer ring are a simple visual marker and not a stop. They will not prevent deeper perforation when screwing the bolt.

It is the surgeon's responsibility to judge the depth of the bolt seating depending on the thickness of the skull.



#### Precaution

Do not unscrew and then retighten the bolt in the same burr hole. This could affect the hold of the bolt in the burr hole.

# 6.2. ZEROING THE CATHETER

Zeroing the Catheter involves calibrating the Catheter in relation to atmospheric pressure.

#### Warning

Perform the zeroing procedure on each new Catheter, **before** implanting it in the patient.

#### Caution

Observe aseptic technique throughout the entire zeroing procedure.

#### Notice

Do not allow the Catheter extension cable connector to come into contact with liquid.

#### Note

The ICT sensor is calibrated in the factory. Therefore, the temperature setting does not need prior zeroing.

#### Prerequisites

- Power on the Monitoring System.
- Connect the Catheter extension cable to the Monitoring System:
  - if using a Monitor, a message inviting you to connect the Catheter displays on the Monitor screen,
  - if using an Interface, the zeroing diode flashes slowly.

#### Procedure

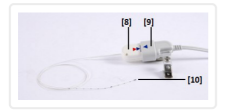
- 1. Unpack the Catheter (sterile) within sterile field.
- Prepare a shallow cup of sterile saline solution (less than 5 mm).

#### Caution

Using a deep receptacle could lead to hydrostatic pressure greater than atmospheric zero, which would then lead to an erroneous reference zero.

3. Observing aseptic technique, connect the Catheter extension cable (non-sterile) to the Catheter, aligning

the blue arrow on the Catheter dongle [8] with the blue arrow on the Catheter extension cable connector [9].



The Catheter dongle must be completely inserted into the Catheter extension cable connector. A blue line on the Catheter dongle indicates the point of complete insertion.

When correctly inserted:

- if using a Monitor, a message displays on the screen,
- if using an Interface, the zeroing diode flashes quickly.

#### Note

The appearance of error codes, "E001", "E002", or "E005" on the Monitor after connecting the Catheter to the Catheter extension cable may be a sign of an incorrect connection. In this case, make sure that the Catheter dongle is pushed right up to the stop in the Catheter extension cable, and that the latter is correctly connected to the Monitor.

 Within the sterile field, immerse the sensor (metal tip) of the Catheter [10] in the cup of sterile saline solution [11], without touching the cup.



#### Precautions

Avoid any contact with the sensor during zeroing procedure, as this could adversely affect the calibration, resulting in an inaccurate zero reference.

Keep the sensor immersed in sterile saline solution during the zeroing procedure. Zeroing the sensor in conditions other than those recommended may cause inaccurate pressure readings.

5. While the sensor is immersed, press the Zero button on the Monitor touch screen, or on the Interface:



Zeroing the Catheter requires approximatively 4 seconds. Do not move the Catheter during the zeroing procedure.

- If the zeroing procedure is successful:
  - the Monitor touch screen displays a message indicating that the Catheter is ready to be implanted,
  - the Interface zeroing diode remains lit.
- If the zeroing procedure is not successful:
  - follow the instructions displayed on the Monitor touch screen,
  - the Interface emits a series of beeps and sends a pressure value of 360 mmHg to the patient bedside monitor. For more information, refer to the instructions for use of the Interface.

#### Note

Zeroing the Catheter can only be performed once. Upon completion, the zero calibration information is stored in the Catheter dongle. It allows the Catheter to be disconnected from a Monitoring System, and re-connected to any Monitoring System, without losing the zero calibration information.

#### **6.3. IMPLANTING THE CATHETER**

Before implanting the Catheter, check that the screen of the Monitor (or of the patient bedside monitor if the Interface is used) displays the value "0 mmHg".

#### Warning

Limit repeated intracerebral implantation of Catheters. Frequent perforation of the brain to allow Catheter insertion may predispose the brain to oedema and intracerebral haemorrhage resulting in an increase in intracranial pressure.

#### Caution

Observe aseptic neurosurgical technique when implanting a Catheter.

Surgeons will perform the most appropriate technique depending upon their experience and the patient clinical status.

The final implantation of the device must satisfy the conditions for optimal positioning of the sensor in the parenchyma.

#### Prerequisites

The Catheter must be zeroed.

#### Procedure

Use the insertion accessories supplied by Sophysa in the Monitoring Kit.

1. Turn the tightening nut counter-clockwise to open the bolt.  Carefully introduce the guide stylet into the bolt to create a passage through the parenchyma, to facilitate the introduction of the Catheter.



3. Remove the guide stylet and irrigate the bolt with sterile saline solution.



- 4. Introduce the zeroed Catheter into the bolt.
- 5. Push the Catheter through the parenchyma. Position it at the desired depth using the centimeter marking.



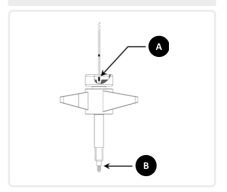
#### Precaution

To perform a parenchymal measurement of intracranial pressure, the implantation depth must be 1 to 3 cm in the cerebral parenchyma.

#### Note

To check the implantation depth, the Catheter has centimeter markings (see figure below). A thicker marker located 4 cm from the Catheter end represents the length of the bolt (A). Once the Catheter is introduced into the bolt, this marker is no longer visible when the sensor at the end of the Catheter just emerges from the bolt (B).

Push the Catheter 1 to 3 centimeters further into the cerebral parenchyma in relation to the thicker marker.



 Once the chosen introduction depth is reached, turn the tightening nut on the bolt clockwise to hold the Catheter in place.



#### Note

The tightening nut must be fully tightened to ensure optimal hold of the Catheter in the bolt.

Once the Catheter is implanted, the Monitor (or patient bedside monitor) displays the mean ICP value in millimeters of mercury (mmHg).

1 mmHg corresponds to 13.60 mmH<sub>2</sub>O and to 133 Pa.

#### Note

The appearance of error codes "- - -", "999", "E001", "E002", "E005", or the message "Connect sensor" on the Monitor after the Catheter implantation, when the Catheter is correctly connected may be a sign of damage to the dongle or to the sensor located on the tip of the Catheter.

#### Specific case for the PSO-PBT

The temperature is also displayed and will stabilise at its

#### Note

A temperature measurement can only be accurate if the sensor is implanted. The sensor is not suitable for measuring the temperature in air.

When the temperature read by the sensor is less than 20 °C or higher than 45 °C, the Monitor displays: " - - . - ". This means that the measurement is in the sensor measuring range but outside the display ranges of the Monitor.

# 6.4. MONITORING, CARING AND NURSING OF THE PATIENT

Once the Catheter is correctly implanted and connected to the Monitoring System, take into account the precautions listed below to ensure optimum monitoring.

#### Precautions

The patient should only be monitored by trained and qualified personnel.

Attach the Catheter extension cable fixation clips to the bed sheets, or to the patient's clothing to limit traction on the implanted Catheter, and reduce the risk of Catheter disconnection.

Move the patient with care in order to avoid disconnecting any cables, or any movement of the implanted Catheter.

After moving the patient, check the connection of the Catheter to the Catheter extension cable, and the connection of the Catheter extension cable to the Monitoring System.

Do not use a Monitoring System and the implanted Catheter at the same time as a high frequency electrosurgical instrument or a defibrillator. The Catheter and/or the Monitoring System could be damaged or have their operation disrupted.

Do not allow any of the connectors to come into contact with liquid, especially during care of the patient.

During monitoring, observe the Catheter implantation site in compliance with standard hospital procedures.

After the first 24 hours, the Monitor displays the duration of the Catheter implantation in the form of the message "Implantation: X days", at the top left corner of the touch screen, for the entire duration of its implantation.

#### Caution

Catheters are recommended to be implanted for up to 6 days (144 hours).

When this duration is exceeded:

- on a Monitor, the message "Implantation: X days" flashes at the top left corner of the touch screen of the Monitor,
- on the Interface, the diode showing the implantation duration for the Catheter flashes.

The accuracy of the displayed ICP value is no longer guaranteed.

#### Note

If a Pressio 2 ICP Monitor is used, it is possible to access the records for the last 15 days of monitoring by using the "History" function on the Monitor.

#### 6.5. EXPLANTING THE CATHETER

- 1. Stop the Monitor, or disconnect the Interface.
- Disconnect the Catheter from the Catheter extension cable.
- 3. Completely unscrew the bolt tightening nut.

#### Precaution

Before explanting the Catheter, make sure that the bolt tightening nut is completely unscrewed.

- 4. Carefully explant the Catheter.
- 5. Check the integrity of the explanted Catheter.
- 6. Unscrew the bolt very carefully and remove it.

#### Precaution

When the bolt is removed, make sure that the spacer ring is also removed.

# 7. Behaviour During an MRI Examination

#### **MRI Safety Information**

Non-clinical testing has demonstrated the Monitoring Kit, Parenchymal with Bolt (PSO-PB) for parenchymal ICP monitoring and the Monitoring Kit, Parenchymal with Bolt (PSO-PBT) for parenchymal ICP and ICT monitoring are MR Conditional. A patient with these devices can be safely scanned in an MR system meeting the following conditions:

- Static magnetic field of 1.5 T and 3.0 T.
- Maximum spatial field gradient of 1,900 gauss/cm (19 T/m) (extrapolated).
- Maximum MR system reported, whole body averaged specific absorption rate (SAR) of 2 W/kg (Normal Operating Mode).
- The non-implanted part of the catheter must be wrapped around the Pressio® MRI Support when there is one available and the MRI Support must be centered in the scanner beforehand. Refer to the information in section 7.2 for the positioning of the Catheters with the MRI Support and section 7.3 for Manual Coiling when there is no MRI Support available.

Under the scan conditions defined above, the Monitoring Kit, Parenchymal with Bolt (PSO-PB) for parenchymal ICP monitoring or the Monitoring Kit, Parenchymal with Bolt (PSO-PBT) for parenchymal ICP and ICT monitoring is expected to produce a maximum temperature rise of less than 2.2 °C after 15 minutes of continuous scanning.

In non-clinical testing, the image artifact caused by the length of the implanted Catheter extends approximately 55 mm from the Monitoring Kit, Parenchymal with Bolt (PSO-PB) for parenchymal ICP monitoring or the Monitoring Kit, Parenchymal With Bolt (PSO-PBT) for parenchymal ICP and ICT monitoring when imaged with a gradient echo pulse sequence and a 3.0 T MRI system.

#### Warning

Disconnect the Catheter from the Monitor before any  $\mathsf{MRI}$  examination.

The "MR Conditional" status only concerns Monitoring Kits for ICP or ICP and ICT. The Monitors, Interface, cables and the temperature module are considered to be "*MR Unsafe*" (not MRI-compatible) and must not be exposed to an MRI environment.

#### Note

The ICP or ICP and ICT Monitoring Kits are considered to be "MR Conditional" according to the definition of the ASTM F2503 standard.

## 7.1. GENERAL POINTS

A patient implanted with an ICP or ICP and ICT Monitoring Kit may undergo an MRI examination after the implantation of the ICP or ICP and ICT Monitoring Kit .

The results of *in-vitro* tests on ICP or ICP and ICT Monitoring Kit have shown that the ICP or ICP and ICT Monitoring Kits do not present any danger for patients in the following conditions:

- Satic magnetic field of 1.5 Tesla or 3 Tesla.

- Maximum spacial field gradient of 1,900 gauss/cm

(19 T/m) (extrapolated).

 $-\,$  Maximum MR system reported, whole body averaged specific absorption rate (SAR) of 2 W/kg (Normal Operating Mode) for 15 minutes exposure at 1.5 or 3 T.

 The non-inserted part of the Catheter is coiled around the MRI Support (when you have one available), and the Pressio MRI Support must be centered in the scanner beforehand.

For more information, refer to the indications mentioned in section "Positioning of the Monitoring Kits and the MRI Support" below.

If you do not have a Pressio MRI Support available,

you can opt for a manual coiling of the Catheter.

For more information, refer to the indications mentioned in section "Positioning of the Monitoring Kits without the MRI Support" below.

— Do not use an emission/reception head coil (RF emission coil for the head) or a simple RF emission head coil. Only use an RF emission/reception whole body coil or an RF emission whole body coil with a simple RF reception head coil.

- Do not scan a patient with a high body temperature.

# 7.2. POSITIONING OF THE MONITORING KITS AND THE MRI SUPPORT

#### Warning

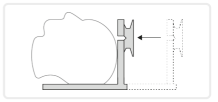
Read the instructions for use before using the MRI Support.

The Catheter must be disconnected from the Pressio®2 Monitor before any MRI examination.

Do not perform an MRI leaving the Catheter uncoiled or in a straight line along the patient's body. This configuration could cause serious injuries to the patient.

Once the patient is installed on the MRI bed:

- Place the MRI support on the bed, the base of the support in contact with the bed.
- 2. Place the support as close as possible to the patient's head.



Move the MRI Support so that it is in the center of the bed.

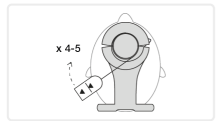


 Pass the non-implanted part of the Catheter through one of the side notches on the body of the MRI Support, and coil it. Depending on the Catheter length, it may be coiled 4 or 5 times round the MRI Support.

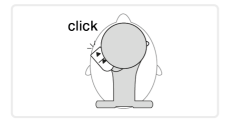
#### Caution

Handle the Catheter with care when coiling it to avoid any excessive traction on the implanted Catheter.

When handling the Catheter, avoid any liquid spray on the connector as this may damage it.



5. Fix the Catheter dongle by inserting it, as shown below, between the flanges of the MRI Support spool.



 Before starting the MRI examination, make sure that the MRI Support is correctly centered. If not centered, adjust its position.

# 7.3. POSITIONING OF THE MONITORING KITS WITHOUT THE MRI SUPPORT

#### Warning

The Catheter must be disconnected from the Pressio®2 Monitor before any MRI examination.

Do not perform an MRI leaving the Catheter uncoiled or in a straight line along the patient's body. This configuration could cause serious injuries to the patient.

When there is no MRI Support available, you can opt for the manual coiling of the Catheter.

Once the patient is installed on the MRI bed, take the non-implanted part of the Catheter and coil it behind the top of the patient's head, in 5 cm loops (up to 4 or 5 loops) and perpendicular to the primary magnetic field.



# 7.4. IN VITRO TEST CONCLUSIONS

The *in vitro* tests conducted in accordance with the ASTM F2182 standard have shown that in the specific conditions mentioned previously, the increase in temperature resulting from exposure to an MRI examination at 1.5 and 3 Tesla is negligible and has no clinical consequences for the patient (maximum increase measured +  $2^{\circ}C + / 0.3^{\circ}C$ ).

The *in vitro* tests conducted in accordance with the ASTM F2052 standard have shown that the displacement forces induced by a magnetic field of 3 Tesla or less do not present any risks for the patient.

The *in vitro* tests conducted in accordance with the ASTM F2213 standard have shown that the torque forces induced by a spatial gradient of the magnetic field less than or equal to 19 T/m do not present any risks for the patient.

The size of the artifacts has been evaluated in accordance with the ASTM F2119 standard.

#### Note

The *in vitro* tests have shown that the operation of the ICP or ICP and ICT Monitoring Kits is not affected by exposure to a magnetic field of 1.5 to 3 Tesla (30,000 Gauss).

# 8. Complications / Side Effects

Complications, which may result from the implantation of a Catheter, include the inherent risks in any surgical procedure and the insertion of a foreign body.

These complications require the rapid intervention of a physician.

#### 8.1. INFECTION

An infection is the major complication associated with this type of monitoring.

Risks of infection can be reduced by observing aseptic techniques when handling and implanting the Catheter, and by respecting the maximum duration for the Catheter implantation (6 days or 144 hours). If monitoring must be continued after this duration, implant a new Catheter on another site.

#### Warning

In case of infection, remove the Catheter and start a specific treatment by a general or intrathecal route.

## 8.2. CEREBRAL HEMORRHAGE

Cerebral hemorrhage may also be observed with this type of monitoring.

Risks of hemorrhage may be reduced by limiting the number of cerebral incisions during the introduction procedure, and by ensuring that this procedure is only performed by trained and qualified personnel.

# 9. Destruction or Return of Product

#### Destruction After Use

An unpacked, used or explanted Monitoring Kit should be destroyed in accordance with the procedures in force in the medical establishment.

#### **Return of Products**

#### Note

For more effective analysis, avoid cleaning the device.

Carefully remove the Catheter from the bolt.

If an explanted Monitoring Kit needs to be returned to Sophysa for analysis, indicate whether cleaning has been performed.

In order to assess the returned product properly, it must be accompanied by an explanatory Return to Manufacturer Authorization form.

# 10. Warranty

The performance of the Monitoring Kits is only warranted with Monitoring Systems and accessories designed, tested and manufactured by Sophysa.

Sophysa warranties that the Monitoring Kits are free of any material and manufacturing defects. Apart from this warranty, Sophysa does not provide any other warranty, express or implicit, including commercialization or adaptation for a particular use. Sophysa cannot be held responsible for any incident, complication, damage or prejudice occurring directly or indirectly from the use of this device. Sophysa does not authorize anyone whomsoever to take responsibility on its behalf for its products.

# 11. Symbols



Meaning
Catalog reference



Lot number

Serial number



Manufacturer



Sterilization Method using Ethylene Oxide



Do not re-use



Do not re-sterilize



The Catheters are not protected against the effects of a cardiac defibrillator.



Refer to the instructions for use



Consult Instructions for use on our website: www.sophysa.us



Use until



BF TYPE EQUIPMENT: Providing an appropriate degree of protection against electric shock, having a Type BF insulated applied section (floating).

122°

Temperature conditions for storage and transport



Keep away from liquids

Fragile, handle with care



Symbol

Meaning MR Conditional.

It has been demonstrated that the use of ICP or ICP and ICT Monitoring Kits within an MRI setting does not pose any risks as long as the specific conditions of use are complied with.

# 12. Technical Specifications

General specifications		
Item	Specification	
Sensor capsule diameter	1.2 mm	
Sensor capsule material	Titanium	
	2.1 Fr (0.7 mm)	
Catheter diameter	D(mm) = Fr / 3 Fr = D(mm) x 3	
Catheter length	1,000 mm (radiopaque on the entire length)	
Catheter sheath material	Polyamide	
Depth markers	Marking every centimeter between 1 and 10 cm from the proximal end. Thicker marking at 4 cm. Additional marking at 15 cm.	
Leakage current	Less than 10 µA to 120 V AC	
Weight	11 g	
Drill bit diameter	2.7 mm, adapted to the screw thread of the bolt	
Note		

The Monitoring Kits are latex- and phthalate-free.

Pressure sensor specifications		
Item	Specification	
Type of sensor	Piezoresistive with strain gauge type on silicon	
Reference pressure	Atmospheric pressure	
Display range (complete system)	-40 mmHg to +100 mmHg (PSO-3000) -40 mmHg to +150 mmHg (PSO-4000)	
Accuracy of the pressure measurement (linearity and hysteresis) (complete system)	$\pm$ 2% reading in the range 0 to 100 mmHg	
Bandwith	> 100 Hz	
Functional range of overpressure without damage	-700 mmHg to +1,250 mmHg	
Temperature coefficient	0.1 mmHg/°C max	
Input resistance	667 Ohms	
Output resistance	810 Ohms	
Excitation voltage	1 to 8 V AC or DC	
Drift from zero	Less than 1 mmHg during the first day, at 37°C Less than 2 mmHg during the first week, at 37°C	

Temperature sensor specifications	
Item	Specification
Type of sensor	Thermistor
Unit	Celsius (°C) or Fahrenheit (°F) °F = °C x (9/5) + 32 °C = (°F - 32) x (5/9) Equivalence in Kelvin (K): K = °C + 273.15 K = °F x (5/9) + 255.37

Temperature sensor specifications	
Item	Specification
Display range (complete system)	+20 °C to +45 °C (+68 °F to +113 °F)
Maximum tolerated error (accuracy) in the reference conditions (complete system)	$\pm$ 0.2 °C max. from 25 °C to 45 °C ( $\pm$ 0.4 °F max. from 77 °F to 113 °F) $\pm$ 0.4 °C max. from 20 °C to 25 °C ( $\pm$ 0.7 °F max. from 68 °F to 77 °F)
Resolution	0.1 °C/0.1 °F

Environmental specifications		
Item	Specification	
Temperature	Operation Storage, transport	+10 °C (50 °F) to +40 °C (104 °F) 0 °C (32 °F) to +50 °C (122 °F)
Humidity	Operation Storage, transport	30% to 75% relative humidity 20% to 95% relative humidity
Atmospheric pressure	Operation Storage, transport	500 hPa to 1060 hPa 500 hPa to 1060 hPa

Catheter history specifications	
Item	Specification
Data accessible	First 15 days of monitoring
ICP data	A value every 20 seconds. This value is an average of the previous 20 seconds.
ICT data	A value every 60 seconds. This value is an average of the previous 60 seconds.
Note Data can only be consulte	ad with a Pressio 2 ICP Monitor.

# 13. References

## PRESSIO<sup>®</sup> CATHETER KITS

PSO-PB	$\operatorname{Pressio}^{\otimes}\operatorname{ICP}\nolimits\operatorname{Monitoring}\nolimits\operatorname{Kit},\operatorname{Parenchymal}$ with Bolt
PSO-PBT	$Pressio^{\otimes}\operatorname{ICP}$ and ICT Monitoring Kit, Parenchymal with Bolt
PSO-PT	Pressio <sup>®</sup> ICP Monitoring Kit, Parenchymal Tunneling
PSO-PTT	$\operatorname{Pressio}^{\otimes}\operatorname{ICP}$ and ICT Monitoring Kit, Parenchymal Tunneling
PSO-VT	Pressio <sup>®</sup> ICP Monitoring Kit, Ventricular Tunneling with external CSF drainage function
PSO-VTT	Pressio <sup>®</sup> ICP and ICT Monitoring Kit, Ventricular Tunneling with external CSF drainage function

### PRESSIO<sup>®</sup> MONITORING SYSTEM

PSO-3000	Pressio <sup>®</sup> ICP Monitor
	Power cable and Catheter extension cable (PSO-EC20) included
PSO-4000	Pressio <sup>®</sup> 2 ICP Monitor
	Power cable and Catheter extension cable (PSO-EC30) included
PSO-IN00	Pressio <sup>®</sup> ICP Monitoring Interface
	Catheter extension cable (PSO-EC20) included

#### PRESSIO<sup>®</sup> ACCESSORIES

PSO-EC20	Catheter extension cable Use only with a Pressio <sup>®</sup> ICP Monitor (PSO-3000) or a Pressio <sup>®</sup> Interface (PSO-IN00)
PSO-EC30	Catheter extension cable Use only with a Pressio <sup>®</sup> 2 ICP Monitor (PSO-4000)
PSO-MCxx	Pressure cable   Use only with a Pressio® 2 ICP Monitor (PSO-4000)   - MC01: PHILIPS (AGILENT) - 12 pins   - MC02: SIEMENS (SIRECUST) - 10 pins   - MC03: SPACELABS & MINDRAY - 6 pins   - MC04: GE DATEX-Ohmeda - 10 pins   - MC05: GE Solar (MARQUETTE) - 11 pins   - MC08: NIHON KOHDEN - 5 pins
PSO-MCT-Y	Temperature cable   Use only with a Pressio <sup>®</sup> 2 ICP Monitor (PSO-4000)   MCT-A: PHILIPS (AGILENT) - 2 pins   MCT-B: SIEMENS - 7 pins   MCT-C: SPACELABS - 10 pins   MCT-F: GE Solar (MARQUETTE), GE DATEX-Ohmeda - 11 pins   MCT-F: HELLIGE, DATEX-Ohmeda, NIHON KOHDEN, MINDRAY & DATASCOPE - JACK 6.35 mm
PSO-MT00	Intracranial temperature module Enables temperature values to be displayed on the Pressio <sup>®</sup> ICP Monitor (PSO-3000). Standard transmitter
PSO-TX00	Enables the pressure and temperature value to be transmitted to a computer. Use only with a Pressio <sup>®</sup> ICP Monitor (PSO-3000)
PSO-DR PSO-MRI	Single use, disposable drill Pressio <sup>®</sup> MRI Support For positioning Pressio <sup>®</sup> Catheter during MRI examination

Technical specifications and list of product references may be modified without notice. Availability may vary according to country.

Page intentionally left blank

Page intentionally left blank

Page intentionally left blank



# Sophysa

5, rue Guy Moquet 91400 Orsay France Tel.: +33 (0)1 69 35 35 00 Fax: +33 (0)1 69 35 36 90 contact@sophysa.com

# Sophysa USA

503 E Summit Street, Suite 5 Crown Point, IN 46307 USA Tel.: +1 219 663 7711 Fax: +1 219 663 7741 contact@sophysa.us

www.sophysa.us

Pressio<sup>®</sup> is a registered trademark of Sophysa. ©2023 Sophysa. All rights reserved.