Video Article

Mouse Model of Intraluminal MCAO: Cerebral Infarct Evaluation by Cresyl Violet Staining

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Abstract

Stroke is the third cause of mortality and the leading cause of disability in the World. Ischemic stroke accounts for approximately 80% of all strokes. However, the thrombolytic tissue plasminogen activator (tPA) is the only treatment of acute ischemic stroke that exists. This led researchers to develop several ischemic stroke models in a variety of species. Two major types of rodent models have been developed: models of global cerebral ischemia or focal cerebral ischemia. To mimic ischemic stroke in patients, in whom approximately 80% thrombotic or embolic strokes occur in the territory of the middle cerebral artery (MCA), the intraluminal middle cerebral artery occlusion (MCAO) model is quite relevant for stroke studies. This model was first developed in rats by Koizumi et al. in 1986¹. Because of the ease of genetic manipulation in mice, these models have also been developed in this species²³. Herein, we present the transient MCA occlusion procedure in C57/Bl6 mice. Previous studies have reported that physical properties of the occluder such as tip diameter, length, shape, and flexibility are critical for the reproducibility of the infarct volume⁴. Herein, a commercial silicon coated monofilaments (Doccol Corporation) have been used. Another great advantage is that this monofilament reduces the risk to induce subarachnoid hemorrhages. Using the Zeiss stereo-microscope Stemi 2000, the silicon coated monofilament was introduced into the internal carotid artery (ICA) via a cut in the external carotid artery (ECA) until the monofilament occludes the base of the MCA. Blood flow was restored 1 hour later by removal of the monofilament to mimic the restoration of blood flow after lysis of a thromboembolic clot in humans. The extent of cerebral infarct may be evaluated first by a neurologic score and by the measurement of the infarct volume. Ischemic mice were thus analyzed for their neurologic score at different post-reperfusion times. To evaluate the infarct volume, staining with 2,3,5-triphenyltetrazolium chloride (TTC) was usually performed. Herein, we used cresyl violet staining since it offers the opportunity to test many critical markers by immunohistochemistry. In this video, we report the MCAO procedure; neurological scores and the evaluation of the infarct volume by cresyl violet staining.

Video Link

The video component of this article can be found at https://www.jove.com/video/4038/

Protocol

Transient Middle Cerebral Artery Occlusion (MCAO)

1. Surgery Procedure (Figure 1)

Transient middle cerebral artery occlusion (tMCAO) is performed on 2- to 3-month old male C57Bl/6 mice (22-28g). This protocol was approved by the IRCM bioethics committee animal care. Surgical tools were sterilized by autoclaving (121 °C at 15 psi for 60 min). Between each animal, they were sterilized using the hot bead sterilizer (15 sec). Surgery table and other equipment are sanitized using 70% ethanol.

Two hours before surgery, mice were analgesized with buprenorphine (0.03 mg/kg b.w. i.p.).

Figure 1A.

1. Deeply anaesthetize mice with isoflurane 5% and then maintain the anesthesia at 2.5% isoflurane. Body temperature of the mice is maintained constant during surgery with a heating pad.
2. Disinfect the fur and skin with 70% ethanol or Betadine. Since shaving produces hair fragment release, microabrasions and inflammation that may impact on stroke pathophysiology, we do not shave mouse fur.
3. Make a midline neck incision and gently pull apart the soft tissues.
4. Under a stereo microscope (Stemi 2000, Zeiss), a blunt dissection is performed to expose the trachea and retract the muscles to locate the carotid artery.
5. The left common carotid artery (CCA) is carefully dissected from surrounding tissue and the CCA is temporarily occluded by a temporary suture (1) using 5-0 silk suture cut into 20 mm segments. Great care should be exercised not to harm the vagal nerve.
6. Separate the bifurcation of the left internal common carotid artery (ICA) and external common carotid artery (ECA). A permanent suture (2) is placed around the ECA, as distally as possible, and another temporary suture slightly tight (3) is placed on the ECA distal to the bifurcation.
7. Clip the left ICA (4) using a reverse-action tweezers to avoid bleeding. Great care should be exercised not to harm the vagal nerve.
8. Cut a small hole into ECA (5) between permanent (2) and temporary (3) sutures.
9. Introduce a 12 mm-long 6-0 silicon-coated (about 9-10mm is coated silicon) monofilament suture (Doccol Corporation) into the ECA (6), completely cut the ECA distal to the permanent suture and invert the occluder into the ICA. The suture is tightly tied around the monofilament to prevent bleeding and the reverse-action tweezers are removed.
10. The occluder is introduced to occlude the origin of the MCA in the circle of Willis. Stop its insertion around 9-10 mm beyond the bifurcation of ECA and CCA. The occluder is blocked and cannot move anymore. Care must be exercised not to penetrate the pterygopalatine artery. The suture (3) on the ECA is tightly tied to fix the monofilament in position.
11. Close the skin with an autoclip wound closing system.
12. Inject 1 ml of saline solution subcutaneously and place mice under an infrared heating lamp during all the post-occlusion period (60 min).

2. Restoration of Middle Cerebral Artery Blood Flow

Before reanesthesia, the neuroscore can be checked to evaluate the success of the surgery.

1. Anesthetize the mice as previously described and remove the autoclips.
2. Slightly open the suture (3) onto the ECA to allow monofilament withdrawal and blood reperfusion.
3. Permanently tie off the temporary suture on the ECA to prevent blood loss. The monofilament is kept for reuse.
4. Remove the temporary suture (1) onto the CCA to allow blood recirculation.
5. Close the wound. The mice should receive another 1 ml saline solution subcutaneously.
6. Place the mice under the infrared heating lamp for 1 hr. After checking that the animal regains mobility, the mouse is returned to its cage
7. On the next day, mice were injected with in situ 3H-D-thymidine (1 µCi/ml saline) to label dividing cells.
8. Place the mice under the infrared heating lamp for 1 hr. After checking that the animal regains mobility, the mouse is returned to its cage
9. Eight hours later, the mice are killed with an overdose of buprenorphine (buprenorphine (0.03 mg/kg b.w. i.p.)
10. The neuroscore can be checked to evaluate the success of the surgery.

3. Sham Operation

For sham operations, all procedures are identical except that the occluder is not inserted.

4. Neuroscore

Neurological deficits allow the evaluation of the success of tMCAO just after reperfusion and later the estimation of the degree of severity of the injury. Neurological deficits are scored as previously described and performed at 1, 24, 48 and 72 hr post-reperfusion. An expanded six-point scale is used:

• 0: normal.
• 1: mild circling behavior with or without inconsistent rotation when picked up by the tail, <50% attempts to rotate to the contralateral side.
• 2: mild consistent circling, >50% attempts to rotate to the contralateral side.
• 3: consistent strong and immediate circling, the mouse holds a rotation position for more than 1-2 sec, with its nose almost reaching its tail.
• 4: severe rotation progressing into barreling, loss of walking or righting reflex.
• 5: comatose or moribund.

5. Cresyl Violet Staining (Figure 2)

Following PBS solution perfusion, brains are quickly frozen in isopentane and stored at -80 °C. Mouse brains may be also perfused with 4% paraformaldehyde (PFA) depending on the planned immunohistochemistry studies. Cryostat-cuts of coronal brain sections (17 μm) are performed. One section out of every 30 is collected on the same slide to have a representative cerebral injury. For volume quantification, 2 slides are stained by cresyl violet staining and an image analysis system (Scion Corporation, Frederick, MD) was used to evaluate the lesion. The injury volume was calculated in arbitrary units (pixels), and expressed as a percentage of the contralateral non-lesioned area for each section. Other slides are kept at -80 °C for immunohistochemistry studies. Alternatively, PFA-fixed brain can be cut at 30-50 μm using vibratome and the infarct volume measured as recommended by Han et al. (2009).
the study. However, the neuroscore evaluation does not allow us to distinguish between mice with an injured hippocampus and those with an intact hippocampus.

No mortality was observed during the surgery day, suggesting that subarachnoid hemorrhages did not occur. When subarachnoid hemorrhage is identified in a mouse, this one is systematically excluded from analysis. The use of silicon-coated monofilament from Doccol Corporation, which is smoother than home-prepared monofilaments, increases the success of tMCAO and reduces subarachnoid hemorrhages. Mortality between 24 hr and 72 hr is 14% in this model, as generally reported for 60 min of occlusion. The mortality observed is probably due to a large infarct volume in this mouse strain. A strong lesion reproducibility has also been observed (standard deviation is 15%), which is very interesting to study neuroprotection molecules where their effect(s) could be hidden by the variability of the model.

To evaluate the extent of brain injury following tMCAO, we opted to use cresyl violet staining (Figure 2) rather than TTC in order to have a lot of materials to test relevant markers\(^6\). The extent of the lesion is relatively consistent. However, we noticed that in some mice, the hippocampus was injured (around 30% of mice). It is interesting to note that cresyl violet staining can be applied up to 1 week later. In the literature, the percentage of brain infarct varies from one study to the other. It depends on the choice of mouse strain, the anesthesia, the thickness of brain sections, the monofilament used, or the staining used\(^6,7\).

![Figure 1. Scheme of the occlusion of the middle cerebral artery using silicon-coated intraluminal monofilament.](image)

**Figure 1. Scheme of the occlusion of the middle cerebral artery using silicon-coated intraluminal monofilament.** A. Simplified scheme of mouse brain and cerebral arteries showing successive sutures and clip to prepare the introduction of silicon-coated monofilament. B. The position of monofilament through the circle of Willis is represented. The monofilament is introduced into ICA via ECA to occlude the base of the MCA. ACA, anterior cerebral artery; BA, basilar artery; CCA, common carotid artery; C. Willis, Circle of Willis; ECA, external carotid artery; ICA, internal carotid artery; MCA, middle cerebral artery; PCA, posterior communicating artery; PPA, pterygopalatine artery.
Discussion

Different stroke models have been developed to mimic stroke consequences in patients. The choice of the stroke model depends on the biological question. The intraluminal MCAO model mimics one of the most common types of ischemic stroke in patients and is less invasive and more consistent than the Tamura model. It is a really interesting model for neuroprotection, neurorepair and cell death analyses. The success of the intraluminal model depends on many factors such as animal sex, age and weight, temperature, anesthesia and the time of surgery, which have to be controlled. The physical properties of the occluder (tip diameter, length, shape and flexibility) are critical for the consistency of the MCAO. Herein, we use the 12 mm-long 6-0 monofilament coated with silicon on 9-10 mm from Doccol Corporation. The great advantage of this monofilament is to reduce subarachnoid hemorrhages and because its length covered all the ICA length, residual blood flow, which comes from the anterior and posterior communicating arteries of the Circle of Willis and PPA, is prevented and the variability in the infarct volume decreases (15% of variability in our hands). The suture of CCA also decreases the variability of the infarct volume.

The extent of brain injury following tMCAO can be assessed by different ways. Neurological deficits can be measured as previously mentioned. However, it is difficult to have an efficient measure by this way. The extent of the infarct is commonly performed by TTC staining. More recently, magnetic resonance techniques are also used, a particularly interesting technology for neuroprotective treatments. To understand the underlying cellular mechanisms involved in stroke such as neurorepair, cell death or cell proliferation, the study of different markers by immunohistochemistry is critical and very informative. Thus, our protocol of brain section preparations has the great advantage of allowing these analyses, and minimizes the number of mice used. Moreover, Tureyen et al. have reported that there is good correlation between cresyl violet staining and TTC.

Disclosures

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