

**Video Article**

**A Mouse Model of Orthopedic Surgery to Study Postoperative Cognitive Dysfunction and Tissue Regeneration**

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**Abstract**

Surgery is commonly used to improve and maintain quality of life. Unfortunately, in vulnerable patients such as the elderly, complications may occur and significantly diminish the outcome. Indeed, after routine orthopedic surgery to repair a fracture, as many as 50% of elderly patients suffer from neurologic complications like delirium. Also, the capacity to heal and regenerate tissue after surgery decreases with age, and can impact the quality of fracture repair and even osseous integration of implants. Thus, a better understanding of mechanisms that drive these age-dependent changes could provide strategic targets to minimize risk for such complications and optimize outcomes. Here, we introduce a clinically relevant mouse model of tibial fracture. The postoperative changes in these mice mimic some of the cognitive impairments commonly observed after routine orthopedic surgery in humans. Briefly, an incision is performed in the right hind limb under strictly aseptic conditions. Muscles are disassociated, and a 0.38-mm stainless steel pin is inserted into the upper crest of the tibia, inside the intramedullary canal. Osteotomy is then performed, and the wound is stapled. We have used this model to investigate the effects of surgical trauma on postoperative neuroinflammation and behavioral changes. By applying this fracture model in combination with parabiosis, a surgical model in which 2 mice are anastomosed, we have studied cells and secreted factors that systemically rejuvenate organ function and tissue regeneration after injury. By following our step-by-step protocol, these models can be reproduced with high fidelity, and can be adapted to interrogate many biologic pathways that are altered by surgical trauma.

**Video Link**

The video component of this article can be found at https://www.jove.com/video/56701/

**Introduction**

Surgery has transformed the medical healthcare system and is continuously contributing to cutting edge technology, improved safety, and maintained quality of life. Unfortunately, surgery also induces pathophysiologic responses that can lead to postoperative complications including wound infections, neurologic impairments, and even mortality, particularly in elderly patients. Orthopedic surgery is performed routinely, especially in older adults, to improve quality of life and repair common bone injuries. However, up to 50% of orthopedic surgery patients who are 65 years and older experience neurologic impairments such as postoperative delirium. This consistently correlates with poor prognosis, i.e., 5-fold increased risk for mortality at 6 months, persistent functional decline, increased nursing time per patient, increased length of hospital stay, and higher rates of nursing home placement. Some risk factors, including advanced age, have been identified, but little is known about the mechanisms responsible for neurologic impairments after surgery.

Since fractures are very common in the elderly, we have established a mouse model of tibial fracture to determine the impact of peripheral trauma on postoperative recovery, including neuroinflammation and brain health (cognitive function). This model, originally described by Harry et al., consists of intramedullary pinning and tibial fracture under general anesthesia and analgesia, and thus mimics the skin injury, muscle trauma, and bone repair associated with common long-bone fractures and repair in humans. After this procedure, mice demonstrate changes in inflammatory markers similar to changes observed in humans, as well as microglial activation in the hippocampus, which is associated with deficits in declarative memory and hippocampal neuropaen/ulcity. We have previously combined this fracture model with parabiosis. Parabiosis is a surgical model in which 2 mice are anastomosed, and thus share a circulatory system. This model has provided a breakthrough in the understanding of regulatory effects of circulating cells and humoral factors on organ function in the context of age and disease. Using this approach we recently discovered systemic factors associated with age-dependent fracture healing.
Here, we introduce a protocol that combines the tibial fracture model with parabiosis to study bone-to-brain age-dependent mechanisms, which are relevant to regenerative medicine and neuroimmunology. Protocol 1A describes the parabiosis procedure, and Protocol 1B details the tibial fracture procedure (Figure 1A). These can be performed independently or in combination, depending on the nature of the interrogation.

### Protocol

All animal experiments were conducted in accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals, and approved by the Institutional Animal Care & Use Committee (IACUC) at Duke University.

#### 1. Experimental Animals

1. Keep mice in an air-conditioned environment with 12-h light/dark cycles and proper access to standard food and water. House no more than 5 littermates per cage, and avoid conditions that could encourage fighting.
2. Use females C57BL6/J mice at 3 months of age (young) or 18 months of age (old).
3. For parabiosis, acclimate 2 mice together for at least two weeks before surgery. Mice are either wild-type C57BL6/J or eGFP+.
4. Check the body condition and overall appearance of mice daily.

#### 2. Preoperative Setup for Parabiosis and Orthopedic Surgery

1. Weigh the 2 mice.
2. Administer general anesthesia via an anesthesia system at a constant O\textsubscript{2} flow rate of 0.2 L/min. Use 5% isoflurane for induction in an induction chamber.
   NOTE: The depth of anesthesia can be confirmed using a toe pinch.
3. Place the mice on a heated pad in a supine position. Use a rectal probe to monitor body temperature throughout the surgical procedure.
4. To maintain anesthesia, lower the inhaled concentration of isoflurane to 2.0% in 0.2 L/min through a facemask. Monitor to ensure adequate depth of anesthesia.
5. Monitor physiologic parameters (respiratory rate, oxygen saturation, and heart beat) non-invasively using a pulse oximeter (optional).

#### 3. Parabiosis Surgery (Protocol 1A)

1. Administer analgesia (buprenorphine, SR/slow release, 0.1 mg/kg subcutaneously) after induction and before surgical manipulation. Inject bupivacaine (0.25%) at the flanks just before opening. Apply eye lubricant.
   NOTE: The parabiosis protocol can be performed independently from the tibial fracture.
2. For all internal body suturing, use 4-0 polydioxanone suture material. For all external suturing, use 4-0 polypropylene suture material.
3. Shave each of the 2 mice along a contiguous line from the elbow, the flank, and the knee on the side to be joined. Disinfect with iodine + 70% alcohol skin scrub over 3 alternating cycles in preparation for incision.
   Used autoclaved instruments and maintain a sterile field.
   NOTE: Place the wild-type mouse to the right of the eGFP mouse, and prepare the left flank of the wild-type mouse and the right flank of the eGFP mouse for surgery (Figure 1).
4. On each mouse, use a scissor to make a skin incision along the flank, ranging proximal to the knee to just proximal to the elbow, and without disturbing the muscles underneath the skin.
5. Join the triceps of the animals with 2 interrupted sutures.
6. Join the body walls along the flanks with a running, continuous suture of 7-9 passes.
7. Join the quadriceps of the animal with 2 interrupted sutures.
8. Close the skin of the 2 parabionts with interrupted sutures.
9. Allow the mice to awake in ambient air.
10. To ensure success of this protocol, the health of the pairs must be maintained: monitor the pairs daily; perform body condition scoring twice weekly; and weigh the pairs twice weekly.
   NOTE: Each pair is housed individually.
   NOTE: If using regular buprenorphine (ie not SR/slow release) then administer 0.1 mg/kg subcutaneously in 1 mL of saline per mouse twice daily for 3 days.
11. Allow 4 weeks of recovery time to ensure shared circulation between parabionts if performing tibial fracture surgery.

#### 4. Tibial Fracture Surgery (Protocol 1B)

1. Administer analgesia (buprenorphine, 0.1 mg/kg subcutaneously) after induction and before surgical manipulation. Inject bupivacaine (0.25%) at the surgical site proximal to the knee just before the opening. Apply eye lubricant.
2. Shave the medial aspect of the right hind limb of the mouse to expose the surgical area, and disinfect with iodine + 70% alcohol skin scrub over 3 alternating cycles. Maintain a sterile surgical field throughout the procedure. To limit contamination; use autoclaved instruments and gloves; complete the surgery under a dissecting microscope (optional); and use a heating pad to maintain body temperature.
   NOTE: If the protocol is performed on parabionts, only 1 mouse is fractured in the pair (the right tibia of the right mouse). See Figure 1B for a schematic diagram of the fracture on parabionts.
3. Use a scissor to make a skin incision along the medial aspect of the right hind limb proximal to the knee down to the midshaft of the tibia on the right mouse.
4. Expose the midshaft of the tibia and visually locate the diaphysis. Flex the knee and visualize the tibial plateau using the patella-femoral ligament as a landmark.
5. Visualize the patellar tendon; manually drill by rolling thumb and index finger a 0.5-mm hole in the intramedullary canal using a 25-gauge needle. 
   NOTE: The drilled hole will run parallel along the tibia, in through the tibial plateau.
6. Insert a 0.38-mm stainless steel pin through the hole about 15 mm into the medullary cavity until resistance is felt, and cut flush with the tibial plateau using a wire cutter (see Supplementary Video 1 for a 3D reconstruction).
7. Using straight Bonn scissors, fracture the tibia midshaft (diaphysis). See Figure 1C for a schematic diagram of the fracture.
8. Visually observe the fracture site and adjacent tissues to inspect for stabilization of the fracture sight.
9. Close with dermal staples.
10. Place the mice on heated pads to recover before returning them to a clean home cage. Inject 1 mL prewarmed (37 °C) normal saline subcutaneously in each mouse for fluid replacement.
11. Check mice daily for signs of lameness, infection, or bleeding. 
   NOTE: If using regular buprenorphine (ie not SR/slow release) then administer 0.1 mg/kg subcutaneously in 1 mL of saline daily for 3 days.

Representative Results

After transverse osteotomies were carefully performed under sterile conditions with steady insertion of the pins and proper suturing, mice showed no signs of lameness, infection, or bleeding in the left leg after surgery. Orthopedic healing was assessed using radiographic analysis and histomorphometry after Safranin-O staining (Figure 2). Radiographs of midshaft fractured tibiae indicated more tissue deposition in the fracture calluses of young mice than in the fracture calluses of aged mice. Fracture calluses were decalcified and embedded in paraffin in preparation for histological analysis. Sections were stained with Safranin-O and tissue deposition was quantified using histomorphometry analysis. Fracture calluses of young mice contained more bone and less fibrotic tissue than fracture calluses from aged mice.

Tibial fracture induces systemic and central inflammation\(^6,7,11,15,16\). Indeed, peripheral levels of pro-inflammatory cytokines and danger-associated molecular patterns (DAMPs) are rapidly elevated after orthopedic surgery, both in mice and humans\(^7,17,18\). This contributes to activation of microglial cells in the brain via several signaling mechanisms involving humoral, cellular, and neuronal pathways\(^7,15,19,20,21\). Following surgery, endothelial dysfunction, blood-brain barrier opening, and peripheral macrophage infiltration contribute to acute hippocampal neuroinflammation in wild-types and Ccr2\(^{RFP/+}\) Cx3cr1\(^{GFP/+}\) adult mice\(^15,19\), and have been associated with subsequent memory deficits that resemble human delirium and postoperative cognitive dysfunction\(^15,19,22\). This neuroinflammatory response is exacerbated in aged animals, with significant changes in microglial morphology, as detected by IBA-1 immunostaining (Figure 3).

Using the tibial fracture model described here, we also found transiently impaired hippocampal neurogenesis, as evidenced by a reduction in doublecortin (DCX) immunostaining in the dentate gyrus\(^23\). Electrophysiologic measurements of long-term potentiation (LTP), a surrogate for memory function, revealed a time-dependent disruption in neuroplasticity postoperatively\(^11\). Mice after surgery also display impairments in hippocampal-dependent memory function, for example using fear conditioning behavioral assessment (Figure 4). In fear conditioning, mice are placed in a chamber and exposed to an auditory cue followed by an aversive stimulus (i.e., footshock). Three days after tibial surgery, mice are tested in the conditioning chamber, this time without any auditory or aversive stimulation, and freezing behavior is recorded as an index of memory (for a detailed protocol see\(^23\)).

Advanced age is a critical risk factor for memory decline. Yet, as we age, the capacity for tissue repair and regeneration also diminishes although these mechanisms remain poorly understood. Thus, we used the protocols herein described to perform heterochronic parabiosis (pairing young/old animals) and assessed fracture healing within the aged mouse (for a detailed parabiosis protocol see\(^24\)). Blood-sharing between parabiotic pairs was confirmed, and found to be equal\(^12\). Exposure to a youthful circulation enhanced bone repair with earlier union, increased bone deposition, and decreased fibrosis (Figure 5)\(^12\). This rejuvenation of bone regeneration occurred independent of the endogenous osteocalcin-positive osteoblasts (Figure 5, brown cells), but rather relied on CD45-positive cells that migrated from the young parabiont (Figure 5, blue cells). These results indicate that the CD45-positive hematopoietic cells secrete a youthful and healthy niche, which is able to signal to the aged osteoblastic cells inducing them to become more active.
Figure 1: Schematic representation of the parabiosis and tibial fracture surgery. (A) Timeline for performing parabiosis and tibial fracture (protocol 1A) or tibial fracture alone (protocol 1B). (B) Tibiae of 20-month-old mice in isochronic or heterochronic parabiotic pairs were fractured (right leg of the right mouse is fractured, as denoted with an X). Grey mice depict wildtype mice while green mice depict eGFP mice. (C) Schematic diagram of tibial fracture model with intramedullary pinning and mid-shaft fracturing. See also Supplementary Video 1 for a 3D reconstruction of the hind limb and pinning of the tibia. Please click here to view a larger version of this figure.
Figure 2: Tibial fracture healing within young and old mouse models. The tibiae of young or old mice were fractured and fracture calluses were investigated 21 days post injury. (A) Radiographic imaging and histologic staining (Safranin-O) were used to evaluate the fracture calluses 21 days post fracture. Staining shows collagenous tissues in blue and proteoglycans (contained within cartilage) in red. Dashed lines indicate approximate location of the fracture callus. (B) Histomorphometry was used to assess the amount of bone, cartilage, and fibrotic tissue deposited within the fracture callus 21 days post fracture. Data are expressed as mean ± 95% confidence intervals, *P<0.05, statistically significant (one-way ANOVA, Dunnett’s test), scale bars represent 2 mm, and images were obtained using a microscope, 1.25x objective. n = 9 for each sample. Please click here to view a larger version of this figure.

Figure 3: Surgery induced age-dependent microglial activation in the hippocampus. Tibial fracture surgery induces greater hippocampal neuroinflammation in aged mice (20-month-old) compared to 4-month-old C57BL6/J mice. Brain-section staining with microglial marker IBA-1 shows more positive cells and morphologic changes in surgical groups 24 h after surgery. Images were obtained with an epifluorescent microscope with a magnification of 100x.; scale bar represents 10 µm. Please click here to view a larger version of this figure.
Discussion

Fractures are a common clinical problem, and remain a leading cause of morbidity, particularly in the rapidly growing senior population. Here, we introduce a step-by-step protocol for a mouse model of tibial fracture to study mechanisms responsible for postoperative neuroinflammation and cognitive impairment. This model can be combined with parabiosis surgery to study neuro-immune interactions, tissue regeneration, and other signaling processes. Understanding these mechanisms will provide strategic targets to minimize the risks for postoperative complications and optimize outcomes.
Several orthopedic models have been developed to study bone repair in rodents. We adopted and modified this tibial fracture procedure, originally described by Harry et al., to study effects of orthopedic surgery on brain function. We have also used this fracture model in combination with our parabiosis model to investigate factors that are responsible for bone healing and age-dependent tissue regeneration. When performed under volatile general anesthesia, this tibial fracture procedure requires only about 15 min per animal, results in zero to minimal mortality (depending on the age of the mouse and underlying genetic susceptibilities), and recapitulates common insults associated with long-bone fracture and orthopedic surgical trauma. Thus, this model is ideal for interrogating biologic pathways and performing longitudinal assessments. However, it is critical that the osteotomy and pinning are reproducible, and that damage to the soft tissue is consistent. Soft tissue damage can be modulated, for example by stripping the periosteum and pinching surrounding muscles to make the surgery more traumatic. Models of traumatic fracture induced by non-fixed blunt trauma or three-point bending would not provide such consistency or accuracy. These procedures often result in re-injury, which leads to prolonged inflammatory response. Conversely, models of fracture involving rigid fixation have a more moderate inflammation, which does not fully recapitulate the damage associated with orthopedic surgery.

Other models using titanium-alloy pinning have been developed to closely mimic human arthroplasty and may be more relevant to interrogate prosthetic instability, osteolysis, and prosthesis-associated complications in mice. Drill-hole models such as the one presented here, provide adequate stabilization, and mice can be tested in behavioral paradigms without significant deficits that could confound such tasks as fear conditioning or open field locomotion/ anxiety testing. However, rotational deformities can occur if the fixation is not properly locked. Some models use an external fixator, which provides superior stabilization but is challenging to implement in a mouse tibia, although it can be successfully implemented in a mouse femur.

Cognitive impairments, including delirium and postoperative cognitive dysfunction, are common complications after orthopedic surgery for fracture repair, especially in elderly and frail patients. This clinically relevant mouse model of tibial fracture surgery demonstrates that postoperative systemic cytokine release, impaired blood-brain barrier function, and altered microglial morphology, contribute to the memory impairment, and may represent critical features of the postoperative neurologic complications seen in many patients after orthopedic surgery. It is important to note that other surgical procedures have been used to model postoperative cognitive dysfunction in mice. These include abdominal and vascular surgery, as well as superficial trauma. The parabiosis technique is applicable to all of these models, which share similar endpoints, including inflammation, glial activation, and behavioral deficits, that may be mediated by common mechanisms.

Studies that include parabiosis have revealed novel roles for circulating factors that can impact cognitive function, neuroinflammation, and tissue rejuvenation in aged animals. We have shown that parabiosis can be successfully combined with the tibial fracture model described here to interrogate regenerative pathways and study mechanisms involving blood-borne factors that influence healing and fracture repair.

Here, we demonstrated that the fracture-repair capacity of an aged animal can be rejuvenated when the aged animal is anastomosed to a young animal. This reversal of age was rooted in the engraftment of hematopoietic cells at the fracture site. Interestingly, such rejuvenation could also be achieved through transplantation of young bone marrow into aged mice. In this regard, bone marrow transplantation can be considered a more direct and simpler alternative approach to parabiosis. However, parabiosis is a more robust model for investigating the function of circulating cells and factors. We expect that a combination of parabiosis and orthopedic surgery models will play an important role in answering critical questions in perioperative care and aging biology.

In summary, we introduce a step-by-step protocol for a mouse model of tibial fracture to study mechanisms responsible for postoperative neuroinflammation and cognitive impairment after orthopedic surgical procedures. This model can be combined with a parabiosis procedure to study neuro-immune interactions, tissue regeneration, and other pathways. Defining these mechanisms will provide strategic targets to minimize the risks for postoperative complications and optimize outcomes.

Disclosures

The authors have nothing to disclose.

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