PURPOSE OF THIS DOCUMENT

This is a handout that accompanies a hands-on mouse biomethodology workshop in the Laboratory Animal Resources Center (LARC).

OBJECTIVES

A. Instruct participants in methods of safe, humane handling and restraint.

B. Instruct participants in substance administration to include {intramuscular (IM), intraperitoneal (IP), subcutaneous (SC), and intravenous (IV)} as well as the technique of gavage.

C. Instruct participants in techniques associated with the collection of blood samples.

D. Instruct participants in the areas of sedation, anesthesia, and analgesia.

E. Instruct participants in methods of euthanasia.
**BASIC INFORMATION ABOUT WORKING WITH MICE**

A. Wear a minimum of a clean laboratory coat and gloves. The use of surgical masks or respirators may assist in reducing allergen exposure.

B. Keep records of each procedure performed on each mouse or group of mice.

C. If bitten:
   - Don’t punish the mouse for its natural response! Calmly return the animal to its cage.
   - Wash the wound thoroughly with an antiseptic soap and water.
   - Cover the wound with a bandage.
   - Notify your immediate supervisor of the bite so that procedures appropriate to the injury can be followed.

D. Mouse psychology:
   - Mice respond positively to quiet, gentle handling. They are normally not aggressive (except for some strains), but if frightened or distressed mice can inflict painful bites.
   - Like any animal, mice are creatures of habit. Everyday events do not tend to stress or excite mice. However, out of the ordinary events such as being picked up, handled, and restrained are stressful and can result in the mice being fractious. Conditioning the mice to handling and restraint will prevent the mice from associating being handled with “negative” things (like being stuck with a needle) and often makes the animals much easier with which to work.
   - Work quietly among the animals, and try to avoid performing procedures in the animal housing room. This will minimize the excitement experienced by the mice from smells and noises, and will allow you to perform your tasks on a more tractable, less stressed animal.

**GENERAL INFORMATION**

**General Biology**

The laboratory mouse (*Mus musculus*) is a mammal of the order Rodentia. The laboratory mouse has been domesticated by man for many generations. Other notable biological characteristics are their very acute hearing, well developed sense of smell, poor vision, small size and short generation interval. Mice are by far the most common laboratory animal used for research.

**Behavior**

The laboratory mouse can be easily handled with appropriate training. Animals that grow up together or those grouped at weaning usually coexist peacefully. However, some strains of mice (i.e. BALB/CJ, SJL/J, HRS/J) may begin to fight even if grouped at weaning. Breeding males that have been removed from breeding cages and then caged together will usually fight. Wounds on the tail or along the back are a common sign of aggression between cage mates.

The *Guide for the Care and Use of Animals* indicates that animals that form social groups (such as mice) should be housed with conspecifics (group housed). Mice can be single housed only for
clinical reasons or if scientifically justified (and approved in the IACUC protocol).

**Biological Characteristics and Data**

Mice, like most species, have a circadian rhythm. Investigators should be aware that this may affect biological data, and it is best to standardize the time of day that samples/measurements are taken to avoid this effect. The standard light/dark cycle in LARC animal rooms is 12/12. This light cycle can be modified upon request by the investigator based on their research needs.

The adult mouse weighs approximately 25-30 grams. The small size and relatively large surface area/body weight ratio makes mice susceptible to changes in environmental conditions. The core body temperature is easily affected by small changes in ambient temperature, which may modify the physiologic responses of the animal. The acute hearing of mice makes them highly sensitive to ultrasounds and high pitched noises inducing a stress response that has been empirically related to cannibalism of pups by their dams. The well developed sense of smell is used to detect pheromones used in social interactions. Mice have poor vision and are unable to detect color. Red light is often used to observe animals during the dark cycle.

The following tables provide general data for many mice, however, for strained specific characteristics and normal physiological data, clinical chemistry, hematology, etc. go to http://jaxmice.jax.org/info/popular.html, http://www.taconic.com/wmspage.cfm?parm1=893, and http://www.criver.com/flex_content_area/documents/rm_rm_n_techbul_spring_99.pdf

<table>
<thead>
<tr>
<th>Basic Biological Data</th>
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<tbody>
<tr>
<td>Adult body weight: male &amp; female</td>
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<tr>
<td>Birth weight</td>
</tr>
<tr>
<td>Body surface area (cm²)</td>
</tr>
<tr>
<td>Life span</td>
</tr>
<tr>
<td>Food consumption</td>
</tr>
<tr>
<td>Water consumption</td>
</tr>
<tr>
<td>Breeding onset: male</td>
</tr>
<tr>
<td>Breeding onset: female</td>
</tr>
<tr>
<td>Estrous cycle</td>
</tr>
<tr>
<td>Gestation Period</td>
</tr>
<tr>
<td>Milk Spot</td>
</tr>
<tr>
<td>Ears open</td>
</tr>
<tr>
<td>Eyes open</td>
</tr>
<tr>
<td>Puberty</td>
</tr>
<tr>
<td>Optimal breeding age</td>
</tr>
<tr>
<td>Body Temperature</td>
</tr>
<tr>
<td>Heart rate</td>
</tr>
<tr>
<td>Respiratory Rate</td>
</tr>
<tr>
<td>Diploid chromosomes</td>
</tr>
<tr>
<td>Metabolic rate</td>
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### Urinalysis

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Approximate normal value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Color</td>
<td>Clear or slightly yellow</td>
</tr>
<tr>
<td>Volume</td>
<td>0.5-2.5 ml/24 h⁽⁷⁾; 0.5-1.0 ml/24 h⁽⁶⁾</td>
</tr>
<tr>
<td>Specific gravity</td>
<td>1.030⁽⁷⁾; 1.058⁽⁶⁾</td>
</tr>
<tr>
<td>pH</td>
<td>5.0⁽⁷⁾; 7.3-8.5⁽⁶⁾</td>
</tr>
<tr>
<td>Glucose</td>
<td>0.5-3.0 mg/24 h</td>
</tr>
<tr>
<td>Protein</td>
<td>0.6-2.6 mg/24 h</td>
</tr>
</tbody>
</table>

### Hematology

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>RBCs</td>
<td>8.7-10.5 x 10⁶/µl⁽⁴⁾</td>
</tr>
<tr>
<td>Hemoglobin</td>
<td>12.2-16.2 g/dl</td>
</tr>
<tr>
<td>Hematocrit</td>
<td>39-43%</td>
</tr>
<tr>
<td>Total leucocyte count</td>
<td>5.1-11.6 x 10⁹/µl</td>
</tr>
<tr>
<td>Neutrophils</td>
<td>6.7-37.2%</td>
</tr>
<tr>
<td>Eosinophils</td>
<td>0.9-3.8%</td>
</tr>
<tr>
<td>Basophils</td>
<td>0-1.5%</td>
</tr>
<tr>
<td>Lymphocytes</td>
<td>63-75%</td>
</tr>
<tr>
<td>Monocytes</td>
<td>0.7-2.6%</td>
</tr>
<tr>
<td>Platelets</td>
<td>100-1000 x 10⁹/µl</td>
</tr>
<tr>
<td>Partial thromboplastin time</td>
<td>55-110 s</td>
</tr>
<tr>
<td>Prothrombin time</td>
<td>7-19 s</td>
</tr>
<tr>
<td>Clotting time</td>
<td>2-10 min</td>
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</table>

### Clinical chemistry (n ± SD)

<table>
<thead>
<tr>
<th></th>
<th>CD-1</th>
<th>C57BL/6</th>
<th>BALB/cBy</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>M</td>
<td>F</td>
<td>M</td>
</tr>
<tr>
<td>Glucose (mg/dl)</td>
<td>112 ± 38.1</td>
<td>97 ± 39.9</td>
<td>121.7 ± 33.2</td>
</tr>
<tr>
<td>BUN (mg/dl)</td>
<td>38 ± 20.1</td>
<td>37 ± 16</td>
<td>32.7 ± 3.5</td>
</tr>
<tr>
<td>Creatinine (mg/dl)</td>
<td>1.1 ± 0.45</td>
<td>0.5 ± 0.08</td>
<td>0.84 ± 0.298</td>
</tr>
<tr>
<td>Na (mEq/l)</td>
<td>166 ± 8.6</td>
<td>166 ± 4.1</td>
<td>166.7 ± 8.9</td>
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<tr>
<td>K(mEq/l)</td>
<td>8.0 ± 0.85</td>
<td>7.8 ± 0.75</td>
<td>8.1 ± 0.8</td>
</tr>
<tr>
<td>Cl (mEq/l)</td>
<td>125 ± 7.2</td>
<td>130 ± 3.9</td>
<td>8.0 ± 1.46</td>
</tr>
<tr>
<td>Ca (mEq/l)</td>
<td>8.9 ± 2.06</td>
<td>10.3 ± 1.58</td>
<td>8.9 ± 2.06</td>
</tr>
<tr>
<td>P (mEq/L)</td>
<td>8.30 ± 1.46</td>
<td>8.00 ± 1.85</td>
<td>8.30 ± 1.46</td>
</tr>
<tr>
<td>Mg (mEq/L)</td>
<td>3.11 ± 0.37</td>
<td>3.13 ± 0.28</td>
<td>3.11 ± 0.37</td>
</tr>
<tr>
<td>Fe (mEq/L)</td>
<td>474 ± 44</td>
<td>473 ± 16</td>
<td>474 ± 44</td>
</tr>
<tr>
<td>ALT (IU/L)</td>
<td>99 ± 86.3</td>
<td>49 ± 22.6</td>
<td>41.4 ± 15.4</td>
</tr>
<tr>
<td>AST (IU/L)</td>
<td>196 ± 132.6</td>
<td>128 ± 60.6</td>
<td>196 ± 132.6</td>
</tr>
<tr>
<td>Alk phos (IU/L)</td>
<td>39 ± 25.7</td>
<td>51 ± 27.3</td>
<td>39 ± 25.7</td>
</tr>
<tr>
<td>LDH (IU/L)</td>
<td>378 ± 269</td>
<td>378 ± 269</td>
<td>378 ± 269</td>
</tr>
<tr>
<td>Protein (g/L)</td>
<td>44 ± 11.0</td>
<td>48 ± 0.85</td>
<td>53.9 ± 7.5</td>
</tr>
<tr>
<td>Albumin (g/L)</td>
<td>36.7 ± 5.2</td>
<td>46.4 ± 7.0</td>
<td>36.7 ± 5.2</td>
</tr>
<tr>
<td>Total cholesterol (mg/dl)</td>
<td>114 ± 56.3</td>
<td>72 ± 20.1</td>
<td>94.8 ± 16.9</td>
</tr>
<tr>
<td>Triglycerides (mg/dl)</td>
<td>91 ± 58.5</td>
<td>53 ± 23.6</td>
<td>97 ± 21.1</td>
</tr>
<tr>
<td>Total bilirubin (mg/dl)</td>
<td>0.4 ± 0.2</td>
<td>0.5 ± 0.35</td>
<td>0.4 ± 0.2</td>
</tr>
</tbody>
</table>
BUN, blood urea nitrogen; ALT, alanine aminotransferase (also known as serum glutamate pyruvate transaminase or SGPT); AST, aspartate aminotransferase (also known as serum glutamic-oxaloacetic transaminase or SGOT); Alk phos, alkaline phosphatase; LDH, lactate dehydrogenase.

Basic Husbandry

Most mice are housed in shoe box cages composed of a plastic (polycarbonate or polysulfone) material with a stainless steel wire bar lid used to hold the water bottle and feed. Bedding is placed directly into the shoe box cage allowing the absorption of urine.

When removing the lid from this type of cage it is important to remove the water bottle to prevent spillage (unless automated watering is used). If the cage is to be transported the bottle should be turned sipper tube up to prevent spillage during transport. However, you should remember to turn the bottle back over to allow access to water after transport.

The animal care staff change the cages on a fixed schedule (frequency depends upon the type of housing, bedding and research needs), thereby providing the animal a clean cage with new bedding, food, and water. Water bottles and feed hoppers are checked daily by caretakers to insure the provision of food and water and to monitor for health or other problems. Some mice are housed on wire mesh bottom cages if scientifically justified and approved in the IACUC protocol. This type of housing is not preferred and is used only when dictated by experimental design.

Pelleted diets are used to feed all rodents and are composed primarily of cereal grains supplemented with additional protein, vitamins and minerals. The water provided to the mice is treated to make it as pure as possible. For mice housed under sterile conditions, the water is autoclaved.

Two levels of barrier housing of mice are available. One level involves the housing of mice in a cage as described above. However, in addition, the cage is fitted with a filter top (microisolater top). This filter top allows pathogen exclusion and containment. LARC also maintains ventilated cage racks which provide HEPA filtered supply and exhaust air to each individual cage. This type of housing also provides for pathogen exclusion and containment. Depending upon the needs of the investigator, a change-out hood can be placed in the animal room to allow for pathogen containment and exclusion when the filter top is removed for cage change-outs or animal manipulations. Also, arrangements can be made to have the entire cage and its contents autoclaved to allow for a sterile environment in which the mice can live. Please contact LARC to discuss the details of these housing options.

A health surveillance program is in place utilizing sentinel animals to detect the presence of rodent pathogens. Rodent pathogens often do not produce clinical signs in affected animals but their presence serves as an unwanted research variable.

Identification

Cage cards are utilized to identify the strain of mouse, sex, number, principal investigator, and IACUC protocol #. Cage cards should not be removed from the cage to avoid misidentification.
of the animals. Temporary identification of individual mice can be accomplished by pen marks on the tail, hair clipping or dyeing the fur. Pen marks will only last a few days whereas hair clipping may last up to 14 days. Ear punch identification and ear tags can be utilized but may be obliterated by fighting between individuals. Microchips and tattoos have also been used for identification.

**Handling (General Information)**

When handling mice it is advisable to wear gloves to prevent the development of allergies due to direct contact with animal allergens. Mice are usually caught and lifted by the tail. The tail should be grasped between its midpoint and the mouse's body. The tail may be grasped with the thumb and forefinger or by the use of smooth-tipped forceps. With this simple method of holding, they may be transferred to another cage, a balance, identified, or casually examined. Pregnant mice or very obese mice may be handled by this method, but they should be supported by use of the second hand placed under their feet. However, such restraint is not sufficient for treatment and close examination. For more effective control, the mouse may be held by the tail and placed on a table or other surface, (preferably one such as a wire cage lid that the mouse can grasp) and the loose skin over neck and shoulders grasped with thumb and fingers (see “handling and restraint exercises” section). It is necessary to perform this maneuver expeditiously, or the mouse may turn and bite. Once the mouse is grasped correctly, the head is adequately controlled. Restraint is improved if the tail or the tail and rear legs are held by the third and little fingers of the same hand or with the other hand (see “handling and restraint exercises” section). Mice should not be dropped into the cage as this may result in spinal fracture. Rather, they should be lowered into the cage and released upon contact with the bedding.

Mice less than two weeks of age can be handled by grasping the loose skin over the neck and shoulder with thumb and forefinger or smooth tipped forceps. Handling neonatal mice should be avoided especially during the first few days after birth to avoid triggering cannibalism or litter abandonment by the dam. If it is necessary to handle the litter, remove the dam to a separate cage and handle the neonates using gloves or better yet with forceps to avoid contamination with human scent. Multiparous females are less likely to cannibalize if they have historically been successful mothers.

Numerous types of restraint devices are commercially available to restrain mice. Quality devices prevent the animal from turning around yet allow easy access to strategic parts of the mouse. Devices should also be easy to clean and provide adequate ventilation.

**Sexing**

Male and female mice can be differentiated by observing the distance from the anus and genital papilla (called the anogenital distance) which is greater in males. This difference is also present in neonatal mice.

In addition, one can usually determine gender by looking for the presence of testicles. However,
one must realize that rodents have the ability to retract their testicles into the abdominal cavity (thus the apparent absence of testicles does not necessarily mean the mouse is a female).

![5-day old Mice]

**HANDLING AND RESTRAINT EXERCISES**

**A  Mouse Restraint Technique I - For removal from caging**

Materials: Disposable gloves

Procedures:

1. Grasp mouse near base of tail (grasping near the tip of the tail may induce a “de-gloving” injury in which the skin on the tip of the tail is removed).
2. Lift animal out of cage and place in new caging or on firm surface.
3. DO NOT suspend mouse by the tail for a prolonged time period.

**B  Mouse Restraint Technique II - For technical manipulation**

Materials: Disposable gloves

Procedures:

2. Lift animal out of cage and place on firm surface.
3. Grasp nape of neck with opposite hand.
4. Place tail between fingers to secure and control animal. Be careful not to grasp too tight as this may compromised breathing.
5. Mouse is now ready for technique manipulations.
C. Mouse Restraint Technique III - For technical manipulation using mechanical restraint

Materials: Disposal gloves, Plexiglas restraint box

Procedures:

1. Restrain mouse by grasping near base of tail.
2. Grasp nape of neck with opposite hand.
3. Place mouse’s head into opening of the restraint box.
4. Release hold on neck while maintaining grasp on tail.
5. Place securing block on appropriate slot for necessary restraint.

D. Mouse Restraint Technique IV - For technical manipulation using DecapiCone®

Materials: Disposable gloves, DecapiCone®

Procedures:

1. Grab the base of the tail with the thumb and index finger.
2. With the head first, place animal in DecapiCone.
3. Gentle restraint is done by wrapping DecapiCone around snug around the animal.
4. The animal is now ready for technical manipulations such as IP injections or decapitation. Note that decapitation of conscious rodents can only be performed when scientifically justified and approved in the IACUC protocol.
INJECTION TECHNIQUES AND BLOOD WITHDRAWAL

Always use sterile syringes and needles for all procedures. To insure aseptic techniques and sharp needles, the one time use of disposable supplies is strongly recommended. When administering injections, select the smallest gauge needle possible to minimize tissue trauma and discomfort. Before injecting the solution, always check for correct placement of the needle by slightly pulling back the plunger of the syringe to create a vacuum. This is known as aspiration. The signs to look for will vary with the injection site. If blood or other fluids are aspirated, placement may be incorrect.

Due to the small muscle mass of many rodents, an intramuscular injection may cause discomfort and local tissue irritation, especially if too large a volume of a solution or a solution with an acidic or alkaline pH is administered. An understanding of anatomy and careful technique are necessary to avoid the ischiatic (sciatic) nerve in the hind leg, just behind the femur. Injection into or close to the nerve may lead to unnecessary discomfort, temporary lameness, or permanent paralysis of the leg. As a result of nerve damage, an animal may chew off the affected extremity.

If too much blood is withdrawn too rapidly, or too frequently without replacement, one may induce hypovolemic shock and/or anemia. As a general guide, up to 10% of the circulating blood volume can be taken on a single occasion from normal healthy mice with minimal adverse effect (mice blood volume = 70 ml/kg body weight). This volume may be repeated after 2-3 weeks. For repeat bleeds at shorter intervals, a maximum of 1% of an animal's circulating blood volume can be removed every 24 hours. However, care should be taken in these calculations, as the percentage of circulating blood will be about 15% lower in obese and older animals.

INJECTIONS EXERCISES

BASIC PROCEDURE

- Wipe the top of the drug bottle with alcohol before withdrawing the dose.
- Slowly withdraw the dose and tap the air bubbles out of the syringe. Air bubbles injected into vessels can potentially cause air emboli and associated problems.
- Always check route of administration for the particular drug to be injected.

A Intramuscular (IM) Injection

Materials: Disposable gloves, Syringe (1 ml), Hypodermic needle (22-30 ga), Injection article, Isopropyl alcohol, Gauze

Procedures:

★ Maximum injection volume = 0.05 ml.

1. Fill syringe with appropriate amount of article to be administered.
2. Restrain mouse.
3. Prep area with alcohol swab.
4. Insert needle into caudal hind leg muscles.
5. Aspirate syringe to insure proper placement. Any sign of blood in the syringe indicates improper placement – reposition.
6. Administer article in a steady, fluid motion. DO NOT administer rapidly because of tissue trauma.

B Subcutaneous (SC) Injection

Materials: Disposable gloves, Syringe (1-3 ml), Hypodermic needle (21-30 ga), Injection article, Isopropyl alcohol, Gauze

Procedures:

★★Maximum injection volume = 2-3ml.

1. Fill syringe with appropriate amount of article to be administered.
2. Restrain mouse.
3. Prep area with alcohol swab.
4. Insert needle at base of skin fold between thumb and opposing finger.
5. Aspirate syringe to insure proper placement. Any sign of blood indicates improper placement; also, a lack of negative pressure in the syringe indicates the needle has punctured out through the opposite side of the skin – reposition.
6. Administer article in a steady, fluid motion.

C Intraperitoneal (IP) Injection

Material: Disposable gloves, Syringe (1-3 ml), Hypodermic needle (22-30 ga), Injection article, Isopropyl alcohol, Gauze

Procedures:

★★Maximum injection volume = 3 ml

1. Fill syringe with appropriate amount of article to be administered.
2. Restrain mouse for technical manipulation. Tilt the body at a 45° angle with the head down. This will position the intestines cranial to the injection site.
3. Prep area with alcohol swab.
4. Insert needle into the mouse’s right lower quadrant of the abdomen at a 30° angle.
5. Aspirate syringe to insure proper placement. Any sign of blood or other fluid indicates improper placement. To prevent inducing peritonitis, remove syringe, discard, and use new syringe, needle, and article in the event that fluids other than blood are aspirated.
6. Administer article in a steady, fluid motion.

**D Intradermal (ID) Injection**

Materials: Anesthetic, Disposable gloves, Syringe (1 ml), Hypodermic needle (25-30 ga), Gauze, Clippers, #40 blade, Isopropyl alcohol

Procedures:

Maximum injection volume = 0.05-0.1 ml depending on thickness of skin.

1. Intradermal injection MUST be done UNDER ANESTHESIA!
2. Clip hair on injection area and prep with alcohol swab.
3. Insert needle between layers of skin on the back at about 20° angle.
4. Aspirate syringe to insure proper placement. Any sign of blood or other fluid indicates improper placement- reposition.
5. Administer article slowly to avoid tissue trauma. Successful injection results in a small circular skin welt.

**E. Intravenous (IV) Injection Utilizing Lateral Tail Veins**

Materials: Disposable gloves, Plexiglas restraint box, Syringe (1 ml), Hypodermic needle (27-30 ga), Injection article, Isopropyl alcohol, Gauze, Warming source

Note: The lateral tail veins of the tail are the most frequently used site for intravenous injection. The secret of successful injection of the tail vein is to dilate the veins. This has been accomplished in various ways such as the following: Placing the tail in warm water (47 °C for about 1 minute (do not exceed 47 °C as this can result in thermal injury to the tail); warming gauze in the warm water bath as stated above and then wrap the tail on the gauze for about 1 minute; placing the animal in an incubator (37° C) for 10 – 15 minutes; or
wrapping the tail in an electric heating pad that is warm (not hot) to the touch. In addition one can place a tourniquet around the base of the tail to facilitate visualization of the vein. See illustration on tourniquet preparation later in this handout. A rubber band and mosquito hemostat also make suitable tourniquets.

The veins can be seen when the tip of the tail is lifted and rotated slightly in either direction. The tip of the needle can be followed visually as it penetrates the vein. Trial injection verifies proper needle placement. Also, accurate placement can be confirmed when the vessel is visually flushed when the compound is administered. The formation of a bleb at the site indicates improper placement of the needle. A second attempt can be performed by removing the needle and trying a site on the same vessel in a more proximal (closer to the animal’s body) location on the tail. Practice is essential.

Procedures:

★Maximum injection volume = 0.2 ml.

1. Restrain mouse in plexiglass restrainer.
2. Warm the tail as described above to facilitate dilation.
3. Needle placement should be no closer to the body than half the length of the tail.
4. With tail under tension, insert needle into skin approximately parallel with the vein.
5. Insure proper placement by inserting needle at least 2-3 mm into lumen of vein.
6. Administer article in a slow fluid motion to avoid rupture of vessel.
7. Upon completion, insure good hemostasis before returning to cage.
GAVAGE EXERCISES

Gavaging the Mouse

Materials: Disposable gloves, Gavage tubes, Syringes (1-3 ml), Injection article

Procedures:

- Maximum administration volume = 0.2 ml (10 ml/kg considered best practice & 50 ml/kg as possible max vol.) for typical adult mouse. Large dose volumes (40 ml/kg) have been shown to overload the stomach capacity, pass immediately into the small bowel (Hejgaard et al 1999) and may also reflux into the esophagus.

1. Measure the distance from the tip of nose to the last rib. This is the length the needle should be inserted.
2. Fill syringe with appropriate amount of article to be dosed.
3. Restrain mouse (Refer to Restraint Technique II).
4. Place tip of needle in the rear of the rat’s mouth to induce swallowing, while gently pressing the shaft of the needle against the roof of the mouth to extend the neck and align the mouth and esophagus.
5. Slide tip down back of mouth, moving tip forward in one fluid motion.
6. Take your time, any resistance felt indicates improper placement. Needle should slide down into esophagus easily. A violent reaction (coughing, gasping) usually follows accidental introduction of the tube into the larynx or trachea.
7. Once the needle is properly placed, administer the article.

**Recommended Standard Gavage Tube Sizes for Mice**

<table>
<thead>
<tr>
<th>Wt. range (grams)</th>
<th>Gauge</th>
<th>Length (inches)</th>
<th>Ball Diameter (mm)</th>
<th>Shape</th>
</tr>
</thead>
<tbody>
<tr>
<td>To 14</td>
<td>24</td>
<td>1</td>
<td>1 ¼</td>
<td>Straight</td>
</tr>
<tr>
<td>15-20</td>
<td>22</td>
<td>1, 1 ½</td>
<td>1 ¼</td>
<td>Straight</td>
</tr>
<tr>
<td>20-25</td>
<td>20</td>
<td>1, 1 ½, 2</td>
<td>2 ¼</td>
<td>Straight, Curved</td>
</tr>
<tr>
<td>25-30</td>
<td>18</td>
<td>1, 1 ½, 3</td>
<td>2 ¼</td>
<td>Straight, Curved</td>
</tr>
<tr>
<td>30-35</td>
<td>18</td>
<td>2, 3</td>
<td>2 ¼</td>
<td>Straight, Curved</td>
</tr>
</tbody>
</table>
BLOOD COLLECTION EXERCISES

A. Lateral Saphenous Vein

Materials: +/- Anesthetic, Disposable gloves, Hypodermic needle (22 gauge), Gauze, Electric clippers, #40 blade, Hematocrit tube or Microvette, petrolatum-based lubricant (can be applied to the puncture site to make blood bead for easier collection)

Procedures:

1. Restrain or anesthetize mouse.
2. Clip hair from lateral aspect of lower leg (between the ankle and knee joints). When clipping the leg, be sure to use small clippers like you will use in the lab. Large clippers can easily induce trauma by cutting the leg.
3. Prep with alcohol swab.
4. Wipe dry with clean gauze.
5. Apply a thin film of petrolatum-based lubricant (such as KY Jelly) over puncture site.
6. Constrict lateral saphenous vein above knee joint between the thumb & index finger.
7. Puncture the vein with a needle. Collect the blood via a hematocrit tube or Microvette.
8. Upon completion, insure good hemostasis by applying gentle pressure to the collection site until blood flow stops.

B. Blood Withdrawal Utilizing Periorbital Venous Sinus*

*Note: This technique has been largely replaced with less invasive blood collection techniques such as from the lateral saphenous vein.

Materials: Anesthetic (systemic and local), Disposable gloves, Hematocrit tubes, Collection vessel, Gauze

Procedures:

1. Anesthetize mouse. After the mouse is anesthetized, place a drop of the proparacaine or tetracaine (local anesthetic) in the eye from which the sample is to be collected. The local anesthetic takes effect in
about 30 seconds and lasts for about 5-10 minutes.
2. Place hematocrit tube at the medial canthus of the eye and insert behind the eye.
3. Rotate tube on its axis on back of orbit until you feel a “pop” and blood flows. Please note that this is a finesse procedure and does not require force.
4. Instill sterile eye ointment when finished.
5. Upon completion, insure good hemostasis by holding eyelids closed.

C. Intracardiac (IC) Puncture

Materials: Anesthetic, Disposable gloves, Syringe (1-3 ml), Hypodermic needle (21-25g), Isopropyl alcohol, Gauze

Procedures:

1. Anesthetize mouse.
2. Prep area with alcohol swab.
3. Insert needle at base of sternum on a 20 degree angle under or just left to the xiphoid process.
4. Aspirate syringe slowly. A good puncture should allow the collection of approximately 1 ml of blood from an adult mouse.
5. This procedure must be followed by euthanasia as it is only permissible as a terminal procedure unless it is scientifically justified and approved in the IACUC protocol.

D. Axillary Cutdown

Materials: Anesthetic, Disposable gloves, Syringe (1-3 ml), Isopropyl alcohol, Gauze, Scissors

Procedures:

1. Anesthetize mouse.
2. With the mouse in dorsal recumbency (lying on its back), prep axillary (armpit) area with alcohol swab.
3. Cut axillary region with scissors or a scalpel blade to expose the subclavian artery and vein which are deep in the armpit.
4. Cut the subclavian artery and vein with the scissors or a scalpel blade.
5. Collect the blood sample with the syringe (no needle) as the blood pools in the axillary region. This is a terminal procedure.
D. Mandibular Puncture by the Golden Rod Animal Lancet Bleeding Technique


Procedures:

1. Restrain the animal securely.
2. Locate the back of the jaw bone, the mandibular area.
3. Prep area with alcohol swab.
4. Puncture the vein with the Golden Rod Animal Lancet.
5. Collect the sample.
6. Press a clean compress to the site for a few seconds.

Proper lancet point length corresponds to the age of the mouse and the volume of blood needed (http://www.medipoint.com/html/animal_lancets.html)

4 mm - 2 to 6 weeks
5 mm - 2 to 6 months
5.5 mm - over 6 months

TOURNIQUET PREPARATION

The following figure displays a practical and inexpensive method of creating a tourniquet for blood collection from the lateral saphenous vein or to prepare the tail for injection into the tail vein.

1. Disassemble a 3 cc syringe. Thread a loop of 2-0 suture material through the syringe.

2. Sew one end of the suture material into the hub of the syringe.

3. Place a needle into the loop and pull the loop tight against the tip of the syringe. Tie the long ends of the suture material together.

4. Cut the long ends of the suture material and reassemble the syringe.

Lateral saphenous blood collection using a syringe & suture tourniquet
ANESTHESIA AND ANALGESIA

METHODS OF ANESTHETIC DELIVERY/EQUIPMENT (OVERVIEW)

There are generally two methods of anesthetic delivery to rodents, parenteral and inhalation.

A. Parenteral anesthesia involves the injectable routes of administration.

B. Inhalation anesthesia involves the delivery of volatile anesthetic agents to the patient via the respiratory tract.

METHODS OF DELIVERY OF INHALANT AGENTS TO RODENTS

The best method for the delivery of volatile agents to rodents involves the use of a precision vaporizer and an anesthesia chamber alone or in combination with a face mask appropriately sized for rodents. **LARC has the equipment to safely and effectively administer inhalant anesthetics (isoflurane) to rodents using a precision vaporizer. Please contact LARC for details regarding use of this equipment.** The rodent is placed within the chamber for induction. Once anesthetized, the animal is removed from the chamber with anesthesia maintained by delivery through a face mask. Both chamber and mask delivery incorporate the use of a precision vaporizer for precise control of the concentration of anesthetic gas delivered to the patient. Because oxygen flow is required to volatilize the liquid anesthetic placed within the vaporizer, oxygen is also delivered to the patient and helps to maintain the blood oxygen saturation. Because fairly high fresh gas flows are required for either chamber or mask delivery, adequate scavenging of waste anesthetic gases is necessary to avoid exposure to personnel. In general, isoflurane anesthesia is superior to injectable anesthesia. Animals are more quickly induced and recovered, and close to 100% of the gas is eliminated through the lungs without being metabolized, (<1% of isoflurane is metabolized). This allows for greater control of the anesthetic depth and tends to minimize experimental variables.

Precision vaporizers must be recertified at the manufacturer’s recommended interval. In the absence of a manufacturer’s recommendation, certification must be performed on an annual basis.

If gas is delivered without a precision vaporizer (called the “Open Drop” method), the following guidelines should be considered:

For induction, a concentration of 2-5% concentration of isoflurane gas is normally adequate. To use anesthetic gas accurately, the induction chamber volume must be known. After determining the chamber volume (it is recommended to record this permanently somewhere easily retrievable), add 0.1-0.2 ml of gas (in liquid form from the bottle) for each liter of chamber capacity. This can be done by applying the gas in liquid phase from its bottle to a cotton ball below the false floor of the container. For
small containers, a piece of cotton can be enclosed in a histology tissue cassette and the agent may be poured or applied onto the cotton in the cassette. Use of 0.2 ml liquid agent per 1000 ml chamber volume will give about a 4% concentration of gas. In the experience of the veterinary staff at Emory University, using nine naïve ICR mice (5 males & 4 females; 2 months of age) introduced to the chamber sequentially after the introduction of isoflurane (0.2 ml/L chamber volume), recumbency was obtained in 57 +/- 21 seconds. However, for rapid and effective induction, the agent had to be replenished in the chamber approximately every 3 mice. Gas delivered by this method must be done under a chemical fume hood or type IIB Biosafety cabinets that are vented to the outside.

<table>
<thead>
<tr>
<th>Volume of liquid agent/1000 ml chamber volume</th>
<th>Approximate concentration of isoflurane or halothane</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.05 ml</td>
<td>1%</td>
</tr>
<tr>
<td>0.1 ml</td>
<td>2%</td>
</tr>
<tr>
<td>0.2 ml</td>
<td>4%</td>
</tr>
<tr>
<td>0.3 ml</td>
<td>6%</td>
</tr>
</tbody>
</table>

**ANESTHETIC MONITORING OF RODENTS**

Parameters that can be used to assess the depth of anesthesia in rodents include:

- recumbency and loss of purposeful movements
- muscle relaxation
- lack of vocalization
- loss of response to aversive stimulation (e.g. pinching the toes)

In most instances, cardiovascular and respiratory assessments are limited to observations of chest wall movement to determine respiratory rate and palpation of the apical pulse through the chest wall.

Because the ratio of body surface area to body mass is greater in rodents than in larger species, thermal support is critical to the successful recovery of rodents from anesthesia. Body heat may be dissipated from the tail, soles of the feet and ears with a resultant profound decline in the core and surface body temperature. This hypothermia may, in turn, lead to a decline in both anesthetic metabolism and any urinary excretion of the anesthetic agent.

**SUPPORTIVE CARE OF ANESTHETIZED RODENTS**

Methods to minimize heat loss to the environment during anesthesia of rodents include increasing the ambient temperature of the operating room; placement of a thermal blanket (e.g. recirculating warm water blanket) or drape between the animal and the stainless steel operating table; use of heat lamps (carefully placed!); minimization of organ exposure from body cavities during surgery; recovery of the animal on a warming blanket or within a temperature-supported
cage; administration of warmed subcutaneous or intraperitoneal fluids before, during or after the anesthetic episode; housing on bedding during recovery to provide thermal insulation; and recovery with cage mates to permit animals to huddle together and thus provide thermoregulation. Do not place an unconscious mouse in a cage with an awake mouse as the alert mouse will tend to mutilate the anesthetized mouse.

Rodents have high energy requirements due to their small size and high metabolic rate, yet they have minimal fat reservoirs which can be mobilized to supply needed energy. Nutritional support is critical upon recovery to avoid hypoglycemia. Nutritional support can be provided by simply providing a high-quality pelleted rodent diet as soon as the animal has recovered sufficiently to ambulate and eat (remember - rodents do not vomit so pre-anesthetic fasting is not typically performed).

Fluid deficits can be corrected by subcutaneous or intraperitoneal injection of warmed saline, Lactated Ringers solution or replacement fluids (e.g., Normosol®). Recommended fluid replacement for mice is 17–33 ml/kg (~0.3-0.7 ml for a 20 g mouse) SC and 33 ml/kg (~0.7 ml for a 20 g mouse) IP.

Because rodents are frequently anesthetized with injectable agents that inhibit blinking (e.g., ketamine), ocular lubrication is important to protect against corneal ulceration.

### Mouse Anesthetics, Anesthetic Cocktails and Sedatives

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dose</th>
<th>Route</th>
<th>Indications &amp; Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ketamine</td>
<td>90-120</td>
<td>mg/kg</td>
<td>IP, IM Surgical anesthesia</td>
</tr>
<tr>
<td>Xylazine</td>
<td>10.00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ketamine</td>
<td>50-75</td>
<td>mg/kg</td>
<td>IP Moderate surgical anesthesia. Not for major surgery</td>
</tr>
<tr>
<td>Medetomidine</td>
<td>1.00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ketamine</td>
<td>30.00</td>
<td>mg/kg</td>
<td>IP Surgical anesthesia</td>
</tr>
<tr>
<td>Xylazine</td>
<td>6.00</td>
<td>mg/kg</td>
<td>IM Surgical anesthesia</td>
</tr>
<tr>
<td>Acetylpromazine</td>
<td>1.00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tribromoethanol</td>
<td>125-250</td>
<td>mg/kg</td>
<td>IP 15-45 min surgical anesthesia, 60-120 min sleep time</td>
</tr>
<tr>
<td>Pentobarbital</td>
<td>40-85</td>
<td>mg/kg</td>
<td>IP Surgical anesthesia</td>
</tr>
<tr>
<td>Yohimbine</td>
<td>2.1</td>
<td>mg/kg</td>
<td>IP Xylazine antagonist for reversal of ketamine/xylazine anesthesia</td>
</tr>
<tr>
<td>Atipamezole</td>
<td>1</td>
<td>mg/kg</td>
<td>IP, SC, IV Medetomidine antagonist for reversal of ketamine/medetomidine anesthesia</td>
</tr>
<tr>
<td>Acetylpromazine</td>
<td>2-5</td>
<td>mg/kg</td>
<td>IP, IP Sedation</td>
</tr>
<tr>
<td>Diazepam</td>
<td>5</td>
<td>mg/kg</td>
<td>IP Sedation</td>
</tr>
<tr>
<td>Xylazine</td>
<td>10</td>
<td>mg/kg</td>
<td>IP Sedation</td>
</tr>
</tbody>
</table>

IM = intramuscular
IP = intraperitoneal
SQ or SC = subcutaneous
IV = intravenous
PO = orally

<table>
<thead>
<tr>
<th>Induction</th>
<th>Maintenance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Isoflurane (Aerrane®, Isoflo®)</td>
<td>Up to 5%</td>
</tr>
</tbody>
</table>

When using isoflurane in a calibrated vaporizer, an oxygen flow rate of 1 liter/minute should be used in conjunction with a non-rebreathing system.

**Analgesic Considerations in Mice**

For *Guidelines for Evaluating and Treating Postoperative Pain in Mice and Rats* (GDLARC001) please visit the LARC website ([http://vpr.utsa.edu/larc.php](http://vpr.utsa.edu/larc.php)).

**Commonly Used Systemic Analgesics in Mice**

<table>
<thead>
<tr>
<th>Class</th>
<th>Buprenorphine* 0.05-0.1 mg/kg SC q 12 h</th>
<th>Butorphanol 1-2 mg/kg SC q 4 h</th>
<th>Morphine 2-5 mg/kg SC q 4 h</th>
<th>Ketoprofen 5 mg/kg SC q 24 h</th>
<th>Carprofen 5 mg/kg SC q 24 h</th>
<th>Flunixin 2.5 mg/kg SC, IM q 12 h</th>
<th>Meloxicam 1-2 mg/kg SC, PO q 12 h</th>
</tr>
</thead>
</table>

* Buprenorphine is the only opioid with long duration effect in rodents.


**NSAID dosing caution:**
1. Ensure that animals are adequately hydrated (skin pinch test, or serum Total Protein test) before administering an NSAID to avoid renal damage.
2. NSAIDs must be used with caution beyond 3 days as it may have deleterious effect on the gastrointestinal mucosa. This may be especially true when using ketoprofen and flunixin.

**Opioid dosing caution:**
1. Opioid agents enhance sedative and respiratory depressive effects of anesthetics.
2. For rodents anesthetized without respiratory support (intubation, ventilation and oxygen supplementation), you may wish consider opioid administration until the end of surgery.
3. In this case, an NSAID may be the preemptive analgesic of choice.
4. If ventilatory support can be provided and an opioid is used as a preemptive analgesic agent, expect to reduce the dose of anesthetic agent (e.g. pentobarbital, isoflurane) by 30-50%.

**Oral dosing caution:**

1. Animals should be acclimated to oral medications before the surgery. When added to the drinking water, rodents will initially refuse to drink until they become adjusted to the flavor, which could be disastrous postoperatively.

2. When used in drinking water, analgesics should generally be administered 5-7 days prior to the anticipated pain insult.

3. Consideration to the use of analgesics in drinking water must take into account that postoperatively, animals may decrease fluid intake and may therefore not receive the intended analgesic dose.

**Local anesthesia/analgesia**

Preoperative (preferred) or intraoperative infiltration of the incision site and underlying tissues with 1-2% lidocaine/0.25-0.5% bupivacaine (50/50 mix by volume) is an effective local anesthesia cocktail to block surgical pain from the offset and for up to 6 hours. Lidocaine provides immediate pain control, while bupivacaine provides longer pain control.

<table>
<thead>
<tr>
<th>Local Anesthetic</th>
<th>Onset</th>
<th>Duration</th>
<th>Do not exceed (toxic dose)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lidocaine (xylocaine)</td>
<td>1-3 minutes</td>
<td>20-40 minutes</td>
<td>10 mg/kg</td>
</tr>
<tr>
<td>Bupivacaine</td>
<td>~20 minutes</td>
<td>4-6 hours</td>
<td>6 mg/kg</td>
</tr>
</tbody>
</table>

**Adjuvants:** Adding epinephrine (1:50:000 to 1:200,000) to plain solutions of local anesthetics just before administration shortens the onset time and prolongs the duration of action. A 1:200,000 dilution is obtained by adding 0.1 ml of 1:1000 epinephrine (with a tuberculin syringe) to 20 ml of local anesthetic. **Epinephrine should not be used for peripheral nerve blocks in areas with poor collateral circulation e.g., digits, tails. Use caution if patient has cardiac problems.**
EUTHANASIA

Proper euthanasia technique includes a follow-up exam to confirm death or the absence of a heartbeat. Monitoring respiration is not considered sufficient since with some euthanasia techniques heartbeat may be maintained after visible respiration has ceased.

The need to minimize fear and apprehension must be considered in determining the method of euthanasia. Distress vocalizations, fearful behavior, and release of certain odors or pheromones by a frightened animal may cause anxiety and apprehension in other animals. Therefore, whenever possible, animals should not be exposed to euthanasia of others.

The euthanasia methods listed in table below are consistent with the American Veterinary Medical Association (AVMA) Panel on Euthanasia, 2007.

MICE EUTHANASIA METHODS

For more details on rodent euthanasia, please refer to the IACUC Euthanasia Policy by going to http://vpr.utsa.edu/oric/iacuc/programpolicies.php

<table>
<thead>
<tr>
<th>Method of Euthanasia</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbon dioxide*</td>
<td>Acceptable method</td>
</tr>
<tr>
<td>Pentobarbital overdose (≥130 mg/kg IV or IP)</td>
<td>Acceptable method</td>
</tr>
<tr>
<td>Inhalant Anesthetic overdose</td>
<td>Acceptable method</td>
</tr>
<tr>
<td>Exsanguinations in anesthetized animal</td>
<td>Acceptable method</td>
</tr>
<tr>
<td>Decapitation in anesthetized animal</td>
<td>Acceptable method</td>
</tr>
<tr>
<td>Cervical dislocation in anesthetized animal</td>
<td>Acceptable method</td>
</tr>
<tr>
<td>Decapitation in awake animal</td>
<td>Acceptable only with scientific justification in writing on the IACUC protocol and subsequent IACUC approval</td>
</tr>
<tr>
<td>Cervical dislocation in awake animal</td>
<td>Acceptable only with scientific justification in writing on the IACUC protocol and subsequent IACUC approval</td>
</tr>
</tbody>
</table>

*Carbon dioxide (CO₂), when used properly, is classified by the 2007 Report of the American Veterinary Medical Association Panel on Euthanasia as a safe method of euthanasia for many small laboratory animals. CO₂ has many advantages including: (1) rapid depressant, analgesic, and anesthetic effects; (2) easy availability in compressed gas cylinders; and (3) inexpensive, nonflammable, nonexplosive, and poses minimal hazard to personnel when used with properly designed equipment.

Although CO₂ is generally considered an acceptable form of euthanasia for small laboratory animals when properly administered, its acceptability is predicated on the following:

The euthanasia chamber should not be prefilled with CO₂, since high concentrations (>70%) can cause nasal irritation, discomfort, and excitability. Rather, the animals should first be placed into the chamber, followed by the addition of CO₂ at a low flow rate (e.g. a rate sufficient to displace
approximately 20% of the chamber volume per minute) to complete the process. In general the low flow rate can be assessed if a very low hissing sound is heard as the chamber is filled with CO₂. Rapid gas flows should be avoided since excessive noises (“winds”) can develop and induce excitement and distress in the animals. In addition, exposure to high CO₂ concentrations is painful as the CO₂ mixes with moisture in the nasal passages to form carbonic acid. Gas flow should be maintained for at least 2 minutes after apparent clinical death (e.g. at least one minute after the animal has quit breathing). Unintended recovery must be obviated by the use of appropriate CO₂ concentrations and exposure times, and followed by a physical method such as cervical dislocation, bilateral thoracotomy, exsanguination or decapitation.

According to the 2007 Report of the AVMA Panel on Euthanasia, ‘Compressed CO₂ gas in cylinders is the only recommended source of carbon dioxide because the inflow to the chamber can be regulated precisely. CO₂ generated by other methods such as from dry ice, fire extinguishers, or chemical means (e.g. antacids) is unacceptable.’ Only one species at a time should be placed into a chamber, and the chamber must not be overcrowded. When placed into the chamber, all animals must have floor space. Euthanasia should always be done in cohorts (live animals should not be placed in the chamber with dead animals). Chambers should be kept clean to minimize odors that might distress animals prior to euthanasia. Animals must not be euthanized in animal housing rooms, except under special circumstances such as during quarantine for infectious disease agents.

Neonates: Since the time period for euthanasia is substantially prolonged in neonatal animals due to their inherent resistance to hypoxia, CO₂ narcosis is generally not recommended. Other methods (see table 2) should be used. The IACUC policy on euthanasia (http://vpr.utsa.edu/oric/iacuc/programpolicies.php) provides further instruction on the euthanasia of the rodent fetus and newborn.