**Brain Removal Protocol for RNAscope**

**10% Sucrose Solution**

* *Sucrose (C12H22O11) 2 g*
* *1x PBS to 20 mL*

**20% Sucrose Solution**

* *Sucrose (C12H22O11) 4 g*
* *1x PBS to 20 mL*

**30% Sucrose Solution**

* *Sucrose (C12H22O11) 6 g*
* *1x PBS to 20 mL*

**Solutions needed***RNAse-free 4% PFA 20 mL*

*10% sucrose solution 20 mL*

*20% sucrose solution 20 mL*

*30% sucrose solution 20 mL*

**Materials needed**

Absorbent bench liner

Falcon tubes

Metal spatula

Metal tweezers (sharp & blunt)

Petri dish

Stereotax (optional)

Surgical microscope (optional)

*Note: Only use items marked with “F” when working with fixative. Wear gloves!*

**Protocol**

1. Set up the working area by putting all materials needed within easy reach of the surgical microscope. Turn microscope on to halfway.
   1. If removing the brain without the microscope and stereotax, jump to step 4.
2. Place the head in the stereotax, using the bird’s ear holes to suspend the head, and the beak around the metal beak bar. Use a screwdriver to tighten the screws.
3. Adjust the microscrope settings.
4. Break and pull away the first two layers of skull (pia mater and arachnoid mater) with a tweezer so that only the last layer is exposed.
   1. These layers are hard and brittle compared to the soft and shiny third layer, the dura mater. The dura is best left intact so that it can be pulled away in one go at the end.
   2. The arachnoid layer is much thicker at the caudal end of the brain, work carefully here so as not to accidentally injure the delicate floculli.
5. Pull back the muscle layers at the back of the neck and cut the white spinal cord.
6. At some point the head will have to be removed from the stereotax in order to reach all sides of brain. When this happens, move the head to a petri dish to work on it.
7. Work around the sides of the skull, cutting the two optic nerves that go from the base of each eye and meet at the ventral side of the brain.
8. When most of the skull is removed and the optic nerves are cut, gently roll the brain out of the head.
9. Place the brain into a new falcon tube with 20 mL of RNAse-free 4% PFA for 24 HRS at 4°C.
10. Move the brain into a new falcon tube with 20 mL of 10% sucrose in 1X PBS at 4°C until the brain sinks to the bottom of the container (~18 HRS for brain tissue).
11. Move the brain into a new falcon tube with 20 mL of 20% sucrose in 1X PBS at 4°C until the brain sinks to the bottom of the container.
12. Move the brain into a new falcon tube with 20 mL of 30% sucrose in 1X PBS at 4°C until the brain sinks to the bottom of the container. Store in the fridge at 4°C.
13. Weigh the brain just prior to flash freezing it.
14. Flash freeze the tissue in OCT (Optimal Cutting Temperature) embedding media with crushed dry ice (see *OCT-embedding-and-flash-freezing-protocol* for detailed instructions).
15. Store tissue blocks in an airtight container at -80°C.