

## 5-1 Transport of 2-Cell Embryos at Cold Temperature

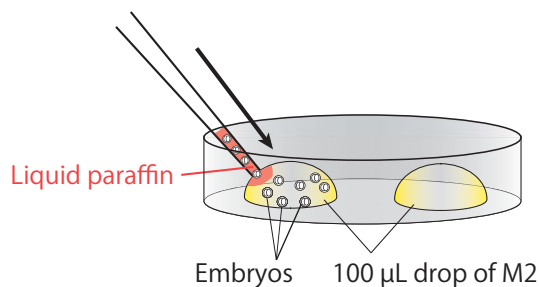
### Materials and Equipment

- 2-cell embryos (adaptable for fresh and frozen/thawed embryos)
- Plastic dishes (35 mm X 10 mm Cat. No. 430588; CORNING)
- Gel-loading tip (Cat. No. 010-R204S; Bio Medical Instrument)
- M2 (Cat. No. M7167; Sigma)
- 0.5 mL tube (Fisherbrand Flip Cap Microtubes 0.5 mL; Fisher Scientific Cat. No. FS-MCT-060 -C)
- Transfer pipettes
- KSOM/AA
- Liquid paraffin
- Temperature data logger (Thermochron iButton Cat. No. DS1921G; Maxim Integrated Products)
- CARD Cold Transport Kit (Cat. No. KYD-006-EX, Cosmo Bio Co., Ltd.)
  - Thermos bottle (Cat. No. JMK-501; Thermos K.K.)
  - Paper box (in which a 0.5 mL tube can stand)
  - Cotton wool
  - Cold packs (small and large)
  - Polystyrene foam transport box (Cat. No. KC-3, KARUX)

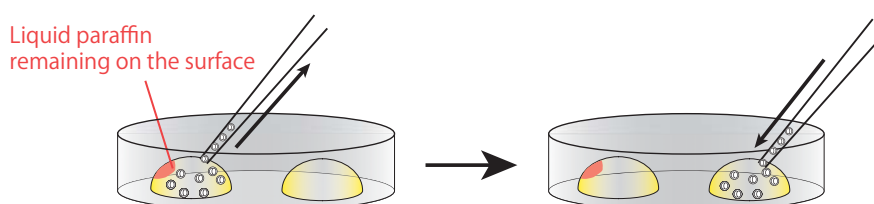
### Procedures

#### Cold Storage of 2-cell Embryos

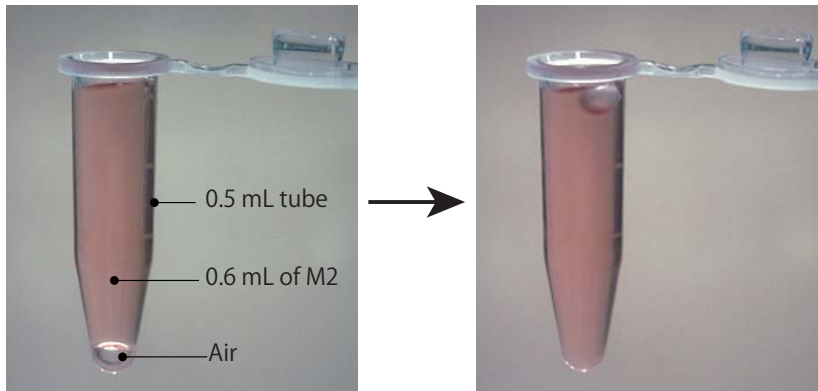
- Place two 100  $\mu$ L drops of M2 on a plastic dish.
- Transfer the 2-cell embryos from the culture medium to the drop of M2.



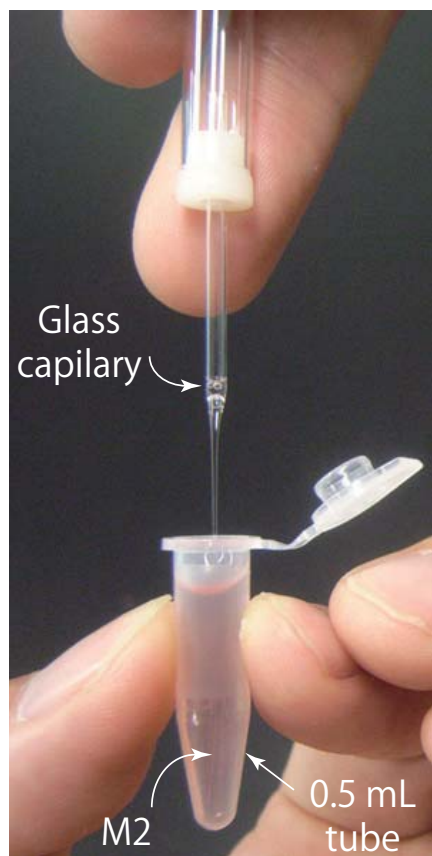
- Exchange the capillary for embryo manipulation and aspirate the embryos into the new capillary, making sure to avoid dropping liquid paraffin onto the drop of M2. Transfer the embryos into the drop of M2 made in step 1.



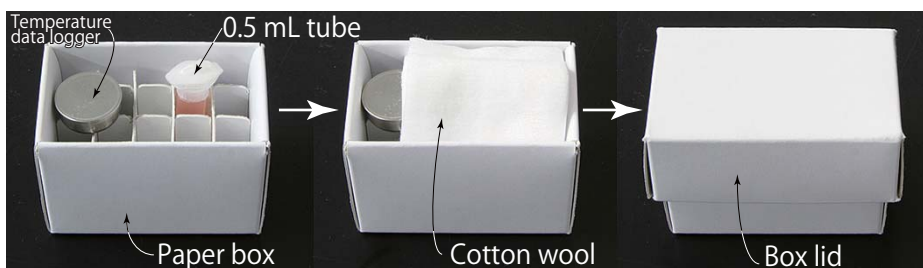
- Fill a 0.5 mL tube with 0.6 mL of M2 at room temperature. If there is a bubble in the bottom of the tube, tap the tip of the tube to release the bubble.



- Collect and transfer the embryos into the bottom of the tube (40 embryos/tube).



- Place the tube containing the embryos, a temperature data logger and a piece of cotton wool in the paper box.



- Store the paper box in the refrigerator (4-8°C).

### Comment

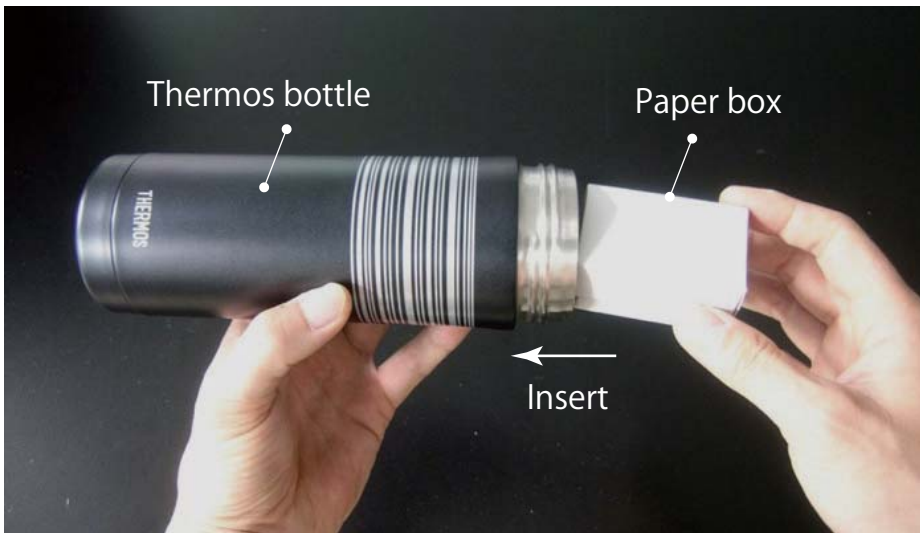
The embryos will maintain developmental ability for up to 72 hours.

**Package and Transport of 2-cell Embryos**

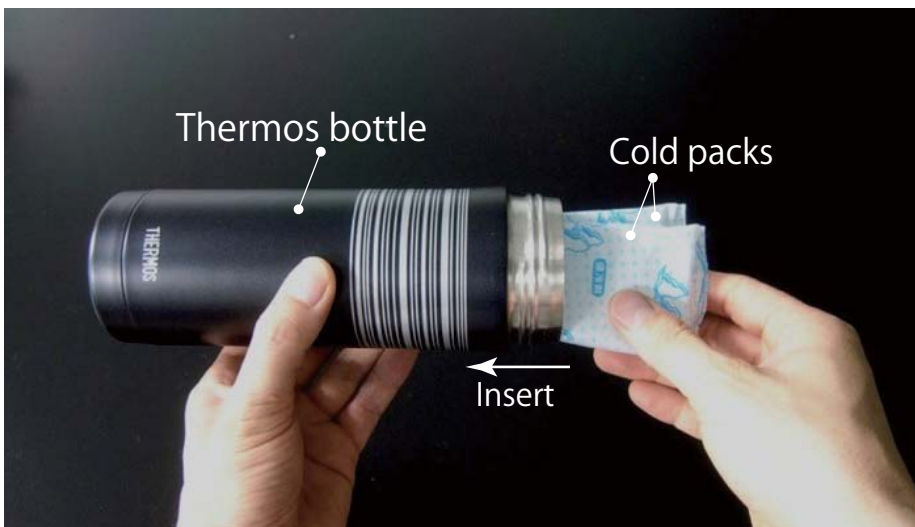
Prepare a paper box containing 2-cell embryos in the same manner as described before (Cold Storage of 2-cell Embryos).

The cold packs (large) and a foam transport box must be pre-cooled to 4-8°C before use. Use the cold packs (small) and a thermos bottle at room temperature.

1. Insert the paper box containing the embryos into a thermos bottle.



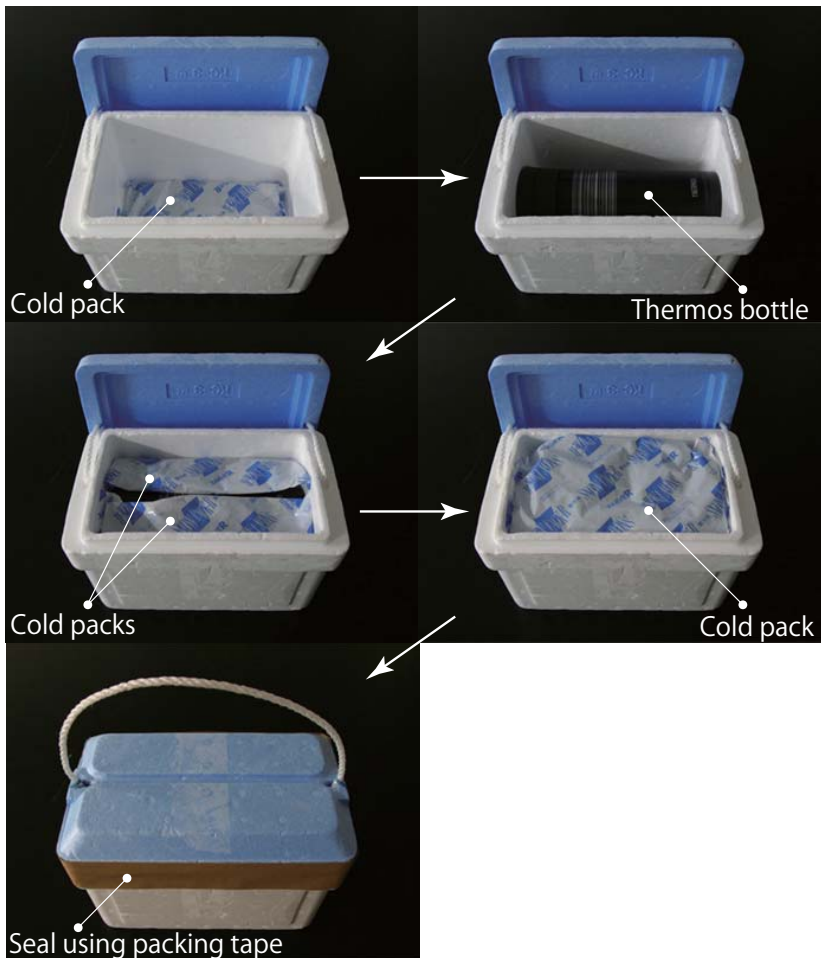
2. Insert two cold packs (small) into the thermos bottle.



3. Close the bottle cap.



4. Put a cold pack (large) in the bottom of a foam transport box, then put the thermos bottle on top of it.
5. Pack one cold pack (large) on either side of the bottle, then set a further pack (large) on top and close the lid.
6. Seal the lid of the foam transport box using packing tape.



7. Keep the foam transport box in the refrigerator until a courier comes to pick it up.
8. Send the samples via regular courier service.

Note

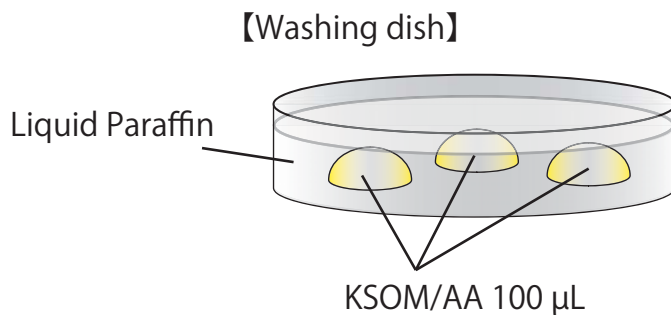
Take care not to place the paper box upside down.

Note

It is only possible to place the thermos bottle in the center of the foam transport box and not the actual bottom, because the length of the thermos bottle is the same as that of the inner length of the foam transport box. This is to protect the thermos bottle during shipping.

**Collection of 2-cell Embryos from the Transport Box**

- Put 3 drops (100  $\mu\text{L}$  / drop) of KSOM/AA into a dish and cover them with liquid paraffin. Place the dish in an incubator (37°C, 5%  $\text{CO}_2$  in air) for at least 30 minutes.

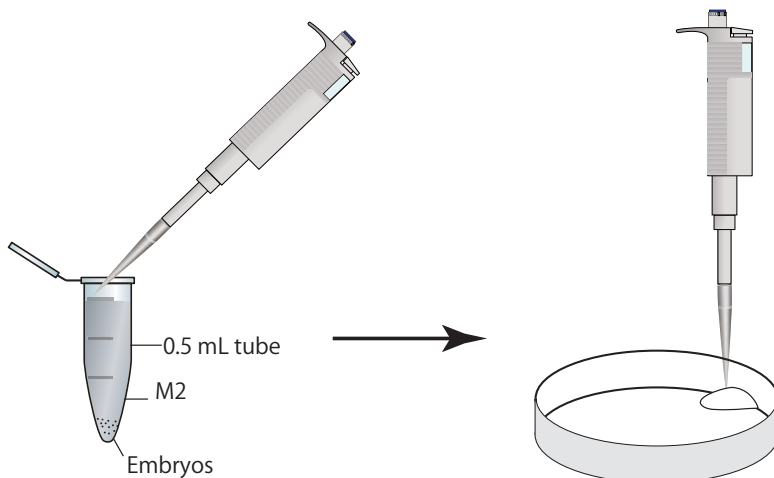


- Retrieve the paper box containing the samples from the thermos bottle.
- Leave the paper box at room temperature for 30 minutes.

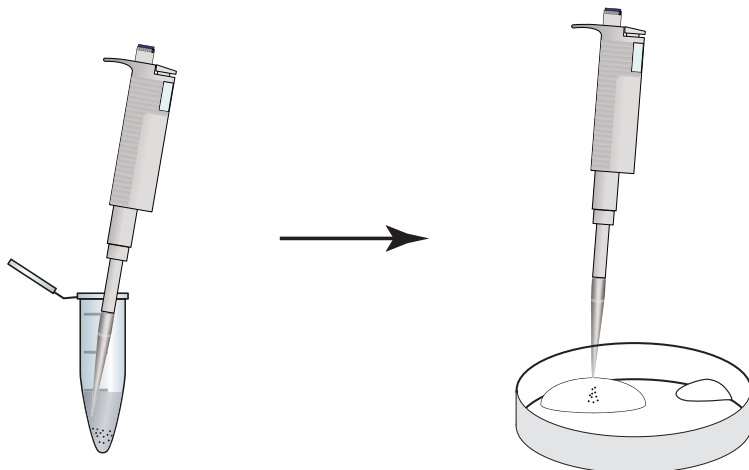
**[Removing the sample]** No. 11-01



- Open the paper box and gently remove the cotton. Once removed, pick up the tube containing the embryos and open it.
- Collect an upper layer of 200  $\mu\text{L}$  M2 from the tube using a gel-loading tip, then transfer the aliquot to the edge of the plastic dish.



- Carefully retrieve all M2 containing the embryos from the bottom of the tube using a gel-loading tip, then transfer the aliquot to the center of the plastic dish.

**Note**

The sample must be transferred at a refrigerated temperature. Please ask the courier service directly about conditions during transport.

**Comment**

The embryos will maintain developmental ability for up to 72 hours.

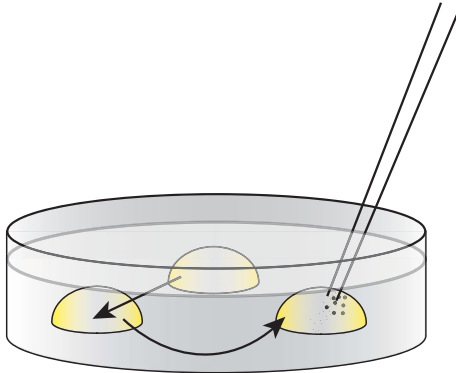
**Comment**

The embryos will sink to the bottom of the tube during those 30 minutes.

**Note**

For easy manipulation, take care to avoid aspirating air bubbles in the gel-loading tip.

7. Collect the embryos from the M2, then transfer and wash them in each of the three drops of 100  $\mu$ L KSOM/AA (washing dish).



8. Transfer the embryos into the oviducts of a pseudo-pregnant mouse.

## References

1. Takeo T., Kaneko T., Haruguchi Y., Fukumoto K., Machida H., Koga M., Nakagawa Y., Takeshita Y., Matsuguma T., Tsuchiyama S., Shimizu N., Hasegawa T., Goto M., Miyachi H., Anzai M., Nakatsukasa E., Nomaru K., and Nakagata N. 2009. Birth of mice from vitrified/warmed 2-cell embryos transported at a cold temperature. *Cryobiology*. **58**(2): 196-202.
2. Takeo T., Kondo T., Haruguchi Y., Fukumoto K., Nakagawa Y., Takeshita Y., Nakamuta Y., Tsuchiyama S., Shimizu N., Hasegawa T., Goto M., Miyachi H., Anzai M., Fujikawa R., Nomaru K., Kaneko T., Itagaki Y., and Nakagata N. 2010. Short-term storage and transport at cold temperatures of 2-cell mouse embryos produced by cryopreserved sperm. *J Am Assoc Lab Anim Sci*. **49**(4): 415-419.

### Note

If you cannot retrieve all stored embryos, rinse the inside of the tube using the 200  $\mu$ L M2 at the edge of the plastic dish.

### Comment

Ideally, embryo transfer to pseudo-pregnant mice should be performed immediately upon the arrival of embryos.

(Please refer to the chapter of Embryo Transfer into the Oviduct on page 66.)

## 5-2 Transportation of Mouse Oviducts Containing 2-Cell Embryos at Low Temperature (0°C)

### Materials and Equipment

1. 0.8 M Sucrose
2. PB1
3. KSOM/AA
4. Plastic bag
5. Thermos bottle
6. Crushed ice
7. Conical bottom cryotubes (Cat. No. 366656; NUNC)
8. Plastic dish (35 mm X 10 mm Cat. No. 430588; CORNING)

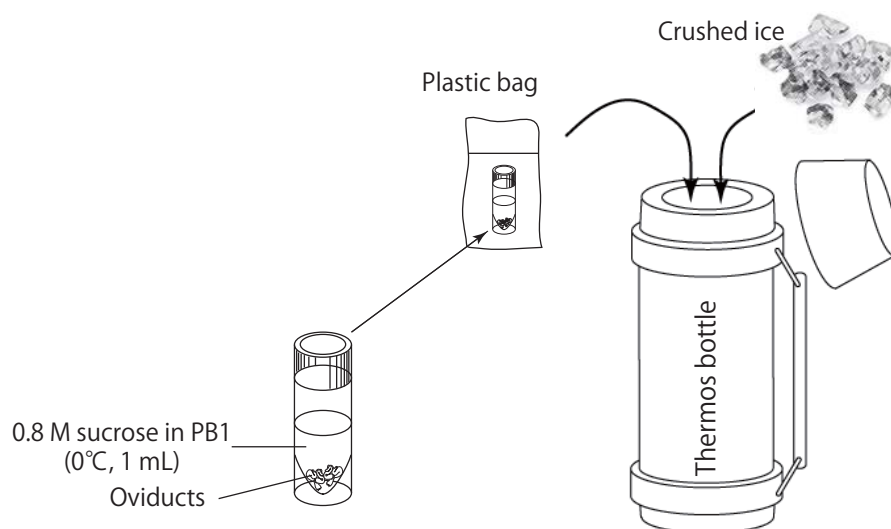
### Procedures

#### Dissection of Oviducts from Superovulated, Plugged Females

1. Inject females i.p. (8-12 weeks old) with 7.5 IU of PMSG (14:00-18:00).
2. Inject females i.p. with 7.5 IU of hCG 48-52 hours after giving them the PMSG injection, and allow the male and female mice to mate overnight.
3. Check the females for vaginal plugs from early the next morning until noon. (Please refer to the chapter of Embryo Transfer into the Oviduct on page 66.)
4. At 44-46 hours after administering the hCG, sacrifice the plugged females.
5. Remove the oviducts of the females, and put them into a 100-200  $\mu$ L drop of 0.8 M sucrose (0°C). (Please refer to the chapter of *In Vitro* Fertilization on page 9.)

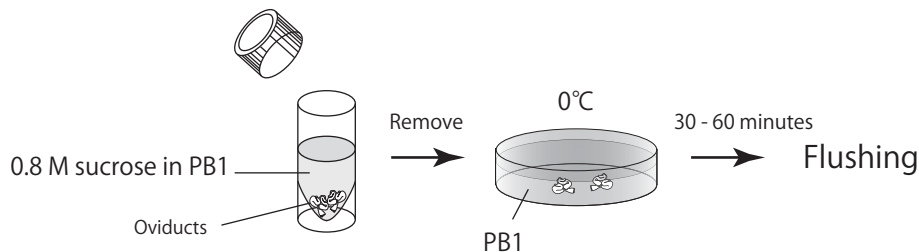
#### Transportation of Mouse Oviducts

1. Transfer the oviducts into a cryotube containing 1 mL of 0.8 M sucrose (0°C).
2. Enclose the tube in a plastic bag and seal it using a sealer.
3. Transfer the plastic bag to a thermos bottle containing crushed ice and transport it using a door-to-door delivery service.



### Collection of Embryos

1. Remove the tube from the thermos bottle.
2. Remove the oviducts from the tube and keep them in PB1 (0°C) for 30-60 minutes.
3. Flush the oviducts with PB1 (0°C). (Please refer to the chapter of Collecting 2-Cell Stage Embryos on page 42.)
4. Wash the embryos with 3 cycles of fresh KSOM/AA (37°C).



### References

1. Kamimura E., Nakashima T., Ogawa M., Ohwada K., and Nakagata N. 2003. Study of low-temperature (4°C) transport of mouse two-cell embryos enclosed in oviducts. *Comp. Med.* 53: 393-396.
2. Ogawa M., Fuchiwaki M., Valdez Jr. Delgado M., Yanagita T., Ide Y., Fukumoto K., Machida H., Kawabe T., Kaneko T., Kasai M., and Nakagata N. 2005. Development after freeze-thawing of mouse embryos collected from oviducts transported at 0°C. *Exp. Anim.* 54(3) Suppl: 242.

#### Note

Embryos will rapidly degenerate if step 2 is skipped.

#### Note

The oviducts should not remain in the tube longer than 48 hours or the embryos will degenerate. Embryos should be frozen if they will not be used immediately. (Please refer to the chapter of Simple Vitrification of Mouse Embryos on page 54.)



## 6-1 Simple Vitrification of Mouse Embryos

### Materials and Equipment

1. 1 M DMSO
2. DAP213
3. Plastic dish (35 mm X 10 mm Cat. No. 430588; CORNING)
4. Filter unit (Millex-GV 0.22  $\mu\text{m}$  Cat. No. SLGV013SL; MILLIPORE)
5. Gel loading tip (MBP Gel 200, Cat. No. 3621; Molecular BioProducts)
6. Transfer pipettes
7. Cryotubes (Cryogenic Vials Cat. No. MS-4501W; Sumitomo Bakelite, Japan is recommended. If you cannot get it, use Cat. No. 366656; NUNC.)
8. Micropipette
9. Vial canes
10. Nalgene Labtop Cooler (Cat. No. 5115-0012; NALGENE, USA)
11. Liquid nitrogen
12. Microscope
13. 0.25 M sucrose
14. KSOM/AA
15. Liquid paraffin

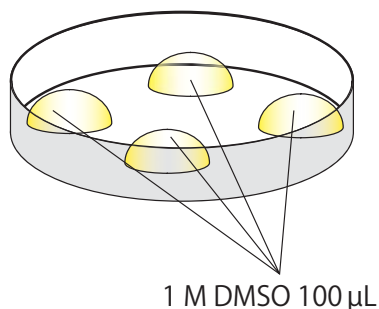
### Procedures

#### Preparation of a Block Cooler and Cryotubes

1. A day before use, place a block cooler (Cat. No. 5115-0012; NALGENE, USA) in a freezer at  $-20^{\circ}\text{C}$ .
2. About 10 minutes before commencing the vitrifying procedure, take the block cooler out of the freezer.
3. Stand some cryotubes in the block cooler. About 40 embryos / cryotube are easy to handle; in other words, when you want to vitrify 120 embryos, you need to stand 3 cryotubes in the block cooler.
4. Just before starting the procedure, check the temperature inside of the tubes is at  $0^{\circ}\text{C}$ .

#### Vitrification

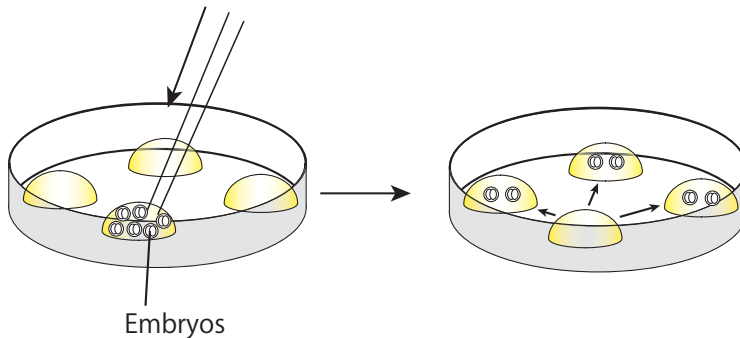
1. Filter the 1 M DMSO and put 4 drops of it ( $\sim 100\ \mu\text{L}$  / drop) into a dish. One drop is to wash the embryos taken from the collection medium, while the others are to hold the washed embryos.



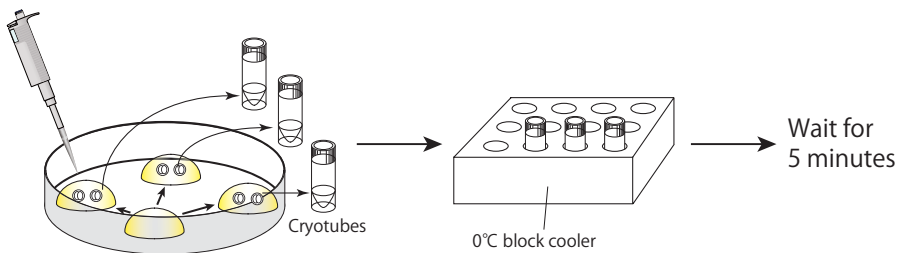
#### Comment

Crushed ice can be used instead of a block cooler.

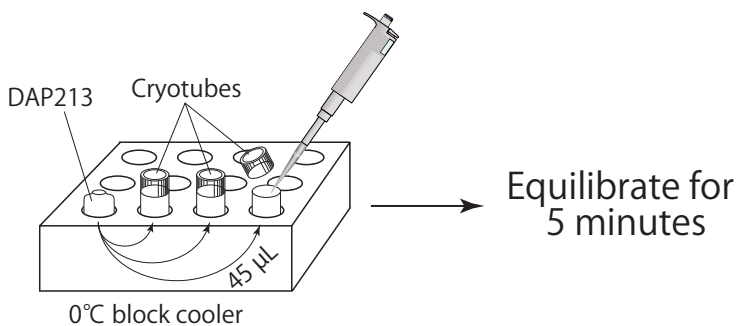
- Place a group of embryos into one of the 4 drops to rinse them of the collection medium. Divide the rinsed embryos equally between the other drops. These aliquots will eventually be transferred to a storage vial. For example, if one were to collect 120 embryos and vitrify them in 40-embryo aliquots, the embryos would first be placed together in the rinse drop and then divided equally among the three drops.



- Using a 20  $\mu\text{L}$  pipette and a gel-loading tip, transfer the embryos contained within 5  $\mu\text{L}$  of 1 M DMSO solution into a cryotube. Once transferred, put the cryotube into the block cooler at 0°C and wait for 5 minutes.



- Add 45  $\mu\text{L}$  of cryoprotective solution (DAP213) at 0°C into the cryotube and equilibrate for 5 minutes in the 0°C block cooler.

**Note**

It is possible to keep the cryotubes in the block cooler at 0°C for longer than 5 minutes (<20 minutes).

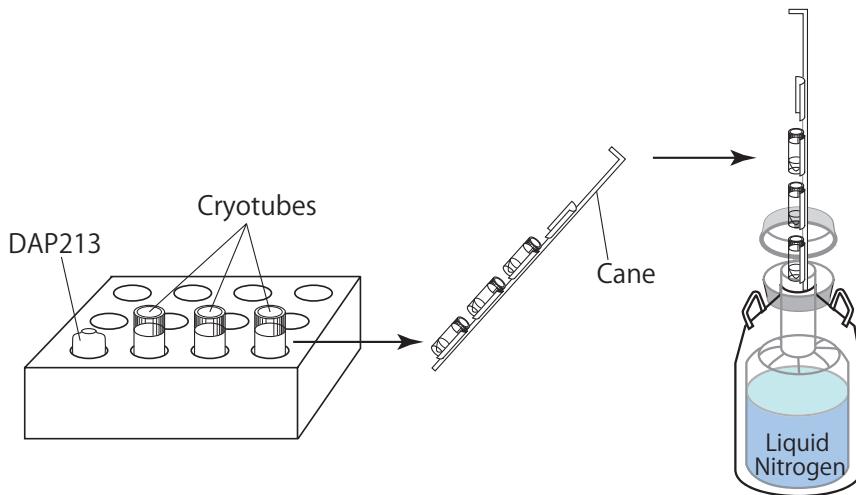
**Note**

If the embryos are pushed together in the center of the drop, it is easy to suck them all up in 5  $\mu\text{L}$  of the 1 M DMSO solution.

**Note**

Do not fasten the caps too tightly after adding the DAP213, or they will be too difficult to remove quickly when samples are recovered from the freezer.

- Quickly set the cryotubes on a cane and plunge the samples directly into liquid nitrogen.



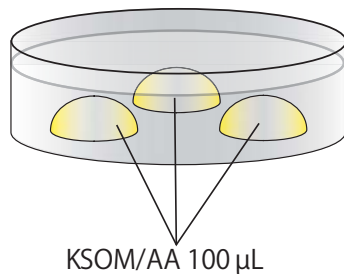
[Vitrifying Embryos] No. 13-01



### Preparation for Thawing

- Put 3 drops (100  $\mu$ L/drop) of KSOM/AA into a dish and cover them with liquid paraffin. Place the dish in an incubator (37°C, 5% CO<sub>2</sub> in air) for at least 30 minutes.

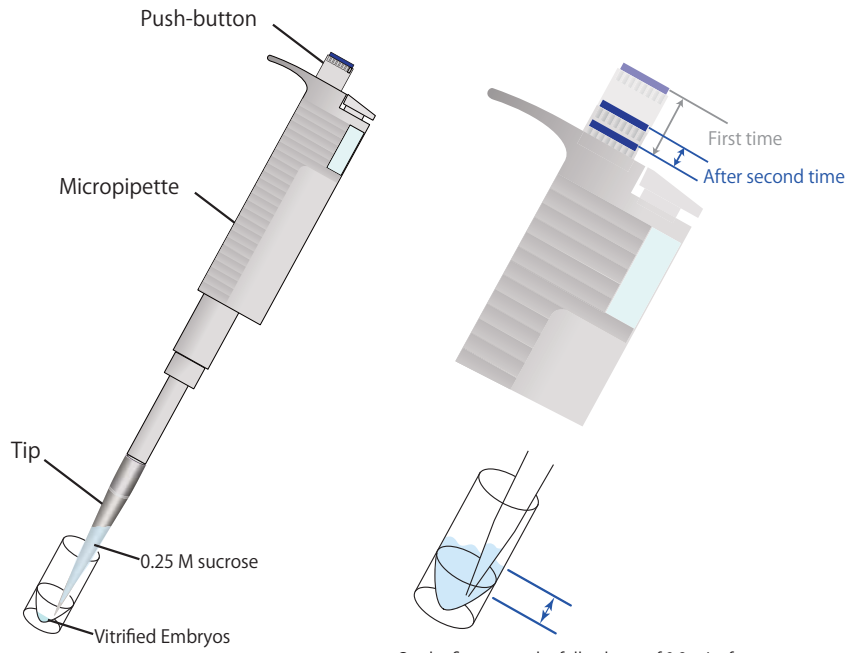
#### 【Washing dish】



- Warm 0.25 M sucrose in an incubator (37°C, 5% CO<sub>2</sub> in air) before use.

### Recovering Vitrified Embryos

- Remove the required sample from the liquid nitrogen and open the cryotube cap. Discard any liquid nitrogen in the tube and allow it to stand at room temperature for 30 seconds.
- Add 0.9 mL of 0.25 M sucrose (preheated to 37°C) to the cryotube and warm the sample quickly via pipetting. When pipetting, take care not to generate large amounts of bubbles and to not physically damage embryos by pipetting too quickly. Once warmed, transfer the contents of the cryotube into a culture dish.




The tip must not touch the bottom of the cryotube. If it does so, the 0.25 M sucrose in the tip will freeze and you will not be able to inject the 0.25 M sucrose into the cryotube.


On the first press, the full volume of 0.9 mL of 0.25 M sucrose will be expelled. On subsequent presses, around 10 amounts of 0.25 M sucrose which you have sucked up will be expelled. Pipetting smaller volumes in this way prevents bubbles from generating in the 0.25 M sucrose.

- Place 0.4-0.5 mL of 0.25 M sucrose into the cryotube, and transfer the contents into the plastic dish. This further dilutes the cryoprotectant and ensures that all of the embryos have been transferred.

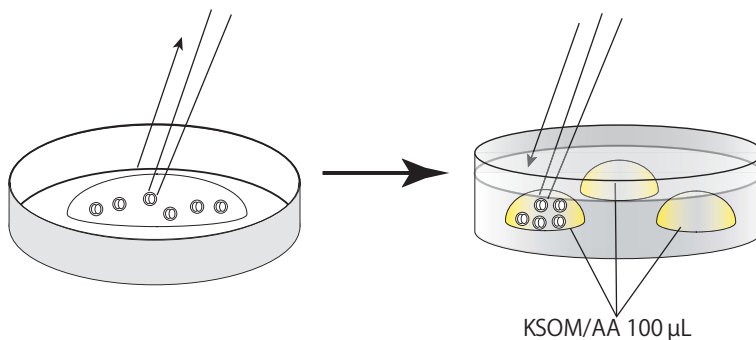
**[Recovering Vitrified Embryos]**

No. 13-02 

**[Pipetting to Recover Vitrified Embryos]**

No. 13-03 

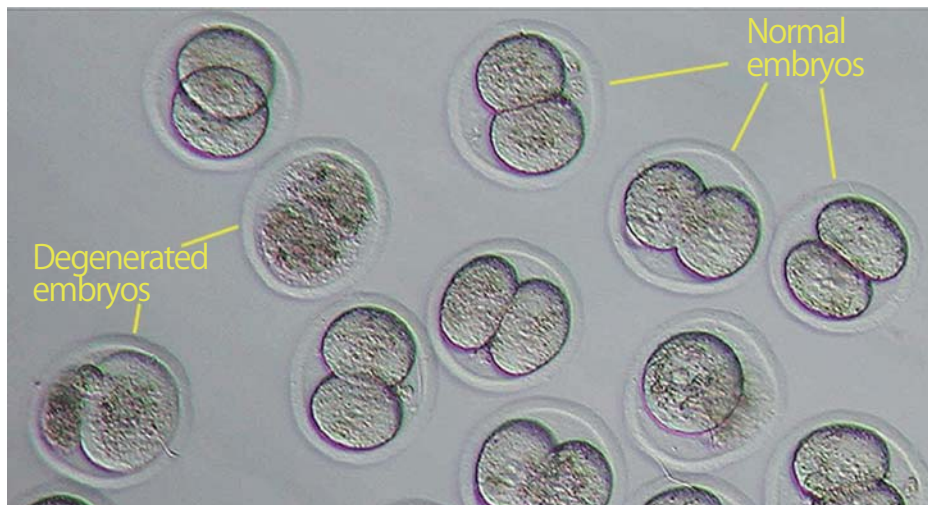
- Aspirate the embryos from the liquid and carefully transfer them into a drop of KSOM/AA (washing dish), then keep them in an incubator (37°C, 5% CO<sub>2</sub> in air).



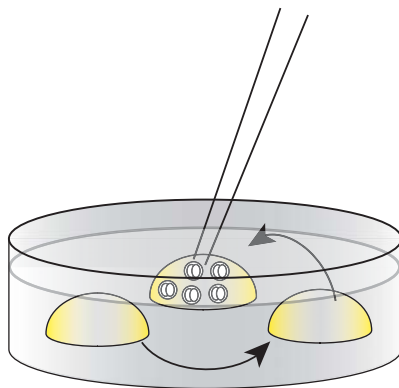
### Note

It is very important to warm the sample quickly to avoid damaging the embryos due to the toxicity of the cryoprotective solution (DAP213).

## 1. [Micrograph : Embryos Recovered after Vitrification]



5. After 10 minutes, wash the embryos with 2 cycles of fresh KSOM/AA (washing dish).



## References

1. Nakagata N. 1989. High survival rate of unfertilized mouse oocytes after vitrification. *J. Reprod. Fert.* **87**: 479-483.
2. Nakagata N. 1993. Production of normal young following transfer of mouse embryos obtained by *in vitro* fertilization between cryopreserved gametes. *J. Reprod. Fert.* **99**: 77-80.
3. Nakagata N. 1995. Studies on cryopreservation of embryos and gametes in mice. *Exp. Anim.* **44**: 1-8.
4. Nakao K., Nakagata N., and Katsuki M. 1997. Simple and efficient procedure for cryopreservation of mouse embryos by simple vitrification. *Exp. Anim.* **46**: 231-234.

## 6-2 Simple Vitrification of Mouse Oocytes

### Materials and Equipment

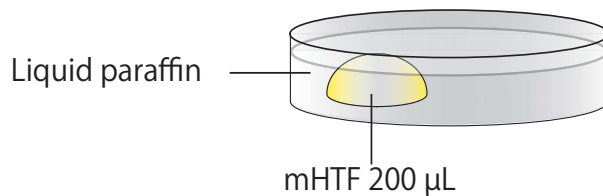
1. Female mice superovulated with PMSG and hCG
2. Plastic dishes (35 mm X 10 mm Cat. No. 430588; CORNING)
3. Liquid paraffin
4. Micropipette
5. Pipette tips
6. mHTF
7. 1% Hyaluronidase in mHTF
8. Fetal bovine serum (FBS Cat. No. 26140-087; Gibco)
9. Filter unit (Millex-GV 0.22  $\mu\text{m}$  Cat. No. SLGV013SL; MILLIPORE)
10. Glass capillaries for embryo handling
11. Humidified incubator (37°C 5% CO<sub>2</sub> in air)
12. Materials and equipment used for vitrification and warming of embryos (For washing warmed oocytes, mHTF drops are used.)  
(Please refer to the chapter of Simple Vitrification of Mouse Embryos on page 54.)

### Procedures

#### Preparation of Dishes

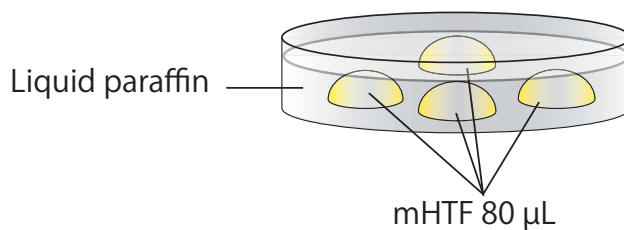
1. Put a 200  $\mu\text{L}$  drop of mHTF into a dish. Cover it with liquid paraffin and place it in an incubator (37°C, 5% CO<sub>2</sub> in air) for at least 30 minutes.

【Oocyte dish】

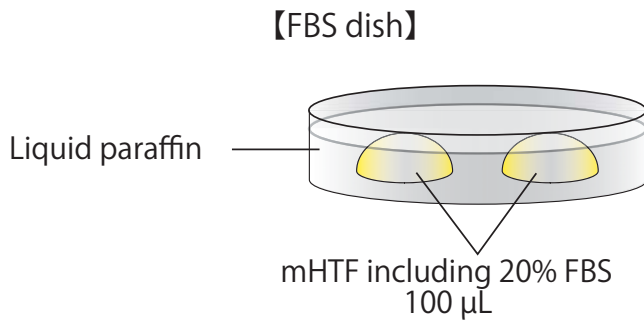


2. Put 4 drops (80  $\mu\text{L}$ /drop) of mHTF into a dish. Cover them with liquid paraffin and place the dish in an incubator (37°C, 5% CO<sub>2</sub> in air) for at least 30 minutes.

【Washing dish】

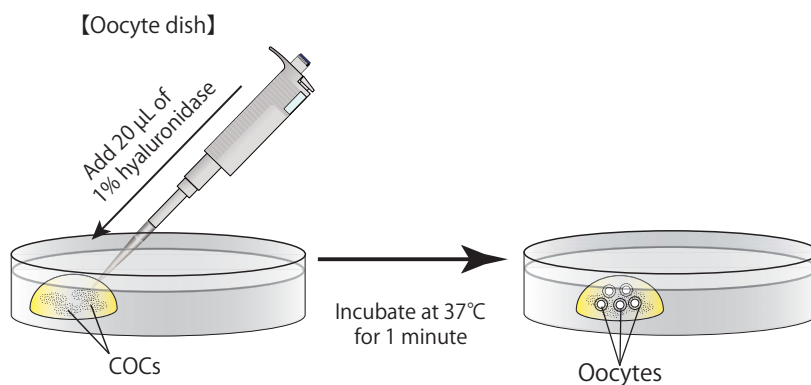


3. Prepare some mHTF including 20% FBS, and sterilize it using a filter. Put 2 drops (100  $\mu\text{L}$ / drop) of the medium into a dish. Cover them with liquid paraffin and place the dish in the incubator (37°C, 5%  $\text{CO}_2$  in air) for at least 30 minutes.

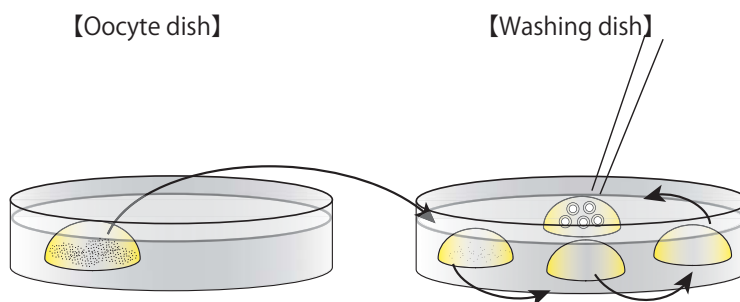


### Preparation of Denuded Oocytes

1. Collect cumulus-oocyte complexes (COCs) from superovulated female mice and introduce them into a 200  $\mu\text{L}$  drop of mHTF (Oocyte dish). (Please refer to the chapter of *In Vitro* Fertilization on page 9.)
2. Add 20  $\mu\text{L}$  of 1% hyaluronidase to the drop of mHTF containing the COCs, and keep the dish in an incubator (37°C, 5%  $\text{CO}_2$  in air) for 1 minute.



3. Promptly collect and transfer the oocytes into a 80  $\mu\text{L}$  drop of mHTF (Washing dish), and in turn wash them in the drops in the washing dish.



### Note

Be sure to carry out all operations, from sacrificing the female and removing her oviducts to introducing the COCs into a drop of mHTF (Oocyte dish), in the shortest time possible (within 30 seconds).

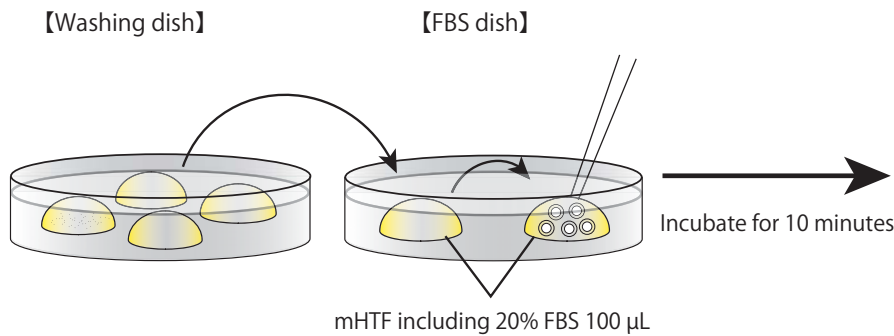
Moreover, when carrying out this process alone, do not sacrifice multiple mice at once; instead, sacrifice one mouse and swiftly remove its oviducts before moving on to the next mouse.

### Comment

If some cumulus cells become attached to the zona pellucida of the oocytes, they can be removed by manipulating the glass capillary.

### Culturing Oocytes in a Drop Containing FBS

1. Transfer the oocytes into the first drop in the FBS dish to rinse. Then, transfer them into the second drop to incubate (37°C, 5% CO<sub>2</sub> in air) for 10 minutes.



#### Comment

FBS can prevent zona hardening in the oocyte during vitrifying and warming.

### Simple Vitrification of Mouse Oocytes

1. The oocytes can be vitrified using the simple vitrification method for embryos, after removing cumulus cells and culturing them in a drop containing FBS. Moreover, the warming method is the same as for embryos. (Please refer to the chapter of Simple Vitrification of Mouse Embryos on page 54.)

### *In vitro* Fertilization using Vitrified-Warmed Oocytes

1. The vitrified-warmed oocytes can be used for *in vitro* fertilization using fresh, cold temperature transported and frozen-thawed spermatozoa. Please refer to the chapters of *In Vitro* Fertilization on page 6, *In Vitro* Fertilization using Epididymal Sperm Transported at Cold Temperature on page 18 and *In Vitro* Fertilization using Cryopreserved Spermatozoa on page 26.

## References

1. Nakagata N., Takeo T., Fukumoto K., Kondo T., Haruguchi Y., Takeshita Y., Nakamuta Y., Matsunaga H., Tsuchiyama S., Ishizuka Y., Araki K. 2013. Applications of cryopreserved unfertilized mouse oocytes for *in vitro* fertilization. *Cryobiology*. 67(2):188-92.

#### Note

There are three different methods of preparing CARD MEDIUM®, depending on whether *in vitro* fertilization will be carried out using fresh, frozen-thawed or cold-temperature transported spermatozoa. Please refer to the CARD MEDIUM® instruction manual.



## 6-3 Vitrification and Transplantation of Mouse Ovaries

### Materials and Equipment

1. 35-mm sterile plastic tissue culture dishes
2. mWM
3. Donor: Female mouse (1 day to 30 weeks old)
4. Recipient: Four-week-old female mouse (a strain that is histocompatible with the transplanted ovary)
5. Anesthetic
6. Micro-spring scissors (5 mm blade)
7. Pair of watchmaker's #5 forceps
8. Wound clip (Autoclip 9 mm; Clay Adams Cat. No. 427631) and clip applicator (Mik-Ron Autoclip Applier; Clay Adams Cat. No. 427630)
9. Hot plate (37°C)

### Procedures

#### Collection of Ovaries

1. Sacrifice a donor female and remove her ovaries.  
(Please refer to the chapter of *In Vitro* Fertilization on page 9.)
2. Place the ovaries in a dish containing an adequate amount of mWM.

#### Vitrification

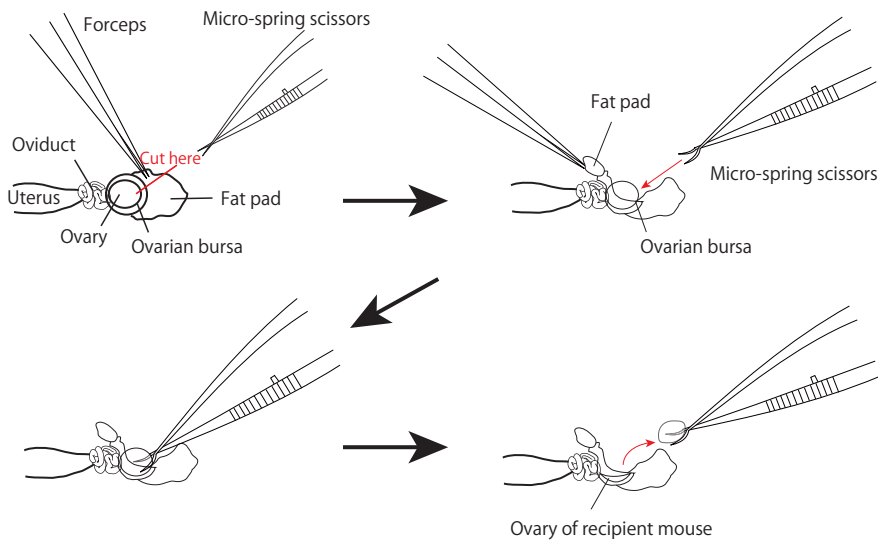
1. Ovaries can be cryopreserved following the same method used for embryos.  
(Please refer to the chapter of Simple Vitrification of Mouse Embryos on page 54.)

#### Transplantation

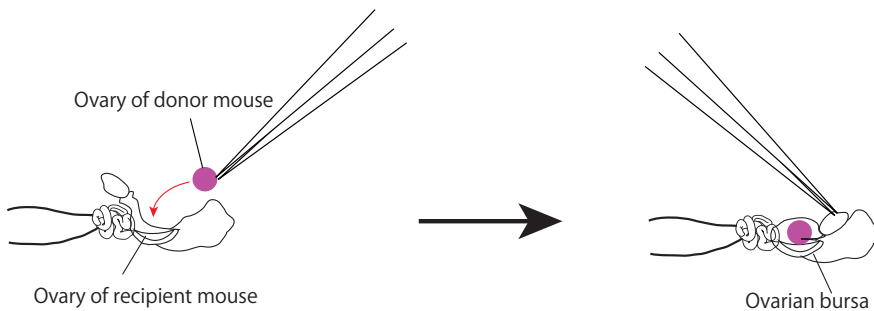
1. Anesthetize a recipient mouse.
2. Pull out the ovary, oviduct and part of the uterine horn as per the conventional procedure.  
(Please refer to the chapter of Embryo Transfer into the Oviduct on page 67.)
3. Using the micro-spring scissors, cut open approximately 1/4 of the recipient's ovarian bursa and some of the surrounding fat pad. Lift up the fat to reveal the ovary.
4. Cut off 1/2 - 2/3 of the ovary using the micro-spring scissors (curved blade).

#### Comment

By using micro-spring scissors with a curved blade, it becomes easier to place the donor ovary on the residual recipient ovary.



5. Insert the donor ovary into the residual ovary of the recipient, and cover it with the ovarian bursa.



**[Transplantation of Mouse Ovary]** No. 15-01



6. Push the ovary, oviduct, and part of the uterine horn back into the abdomen, and close the wound using wound clips.
7. Repeat the process for the ovary on the other side of the recipient mouse as described above.
8. Keep the mouse warm on a 37°C warming plate until the mouse recovers from the effects of the anesthesia.

## References

1. Migishima F., Suzuki-Migishima R., Song S.Y., Kuramochi T., Azuma S., Nishijima M., and Yokoyama M. 2003. Successful cryopreservation of mouse ovaries by vitrification. *Biol. Reprod.* **68**: 881-887.
2. Tsuchiyama S., and Nakagata N. 2009. Cryopreservation of ovaries from elderly female mice. *Exp. Anim.* **58**(3) Suppl: 248.

## 7-1 Vasectomy for the Creation of Sterile Males

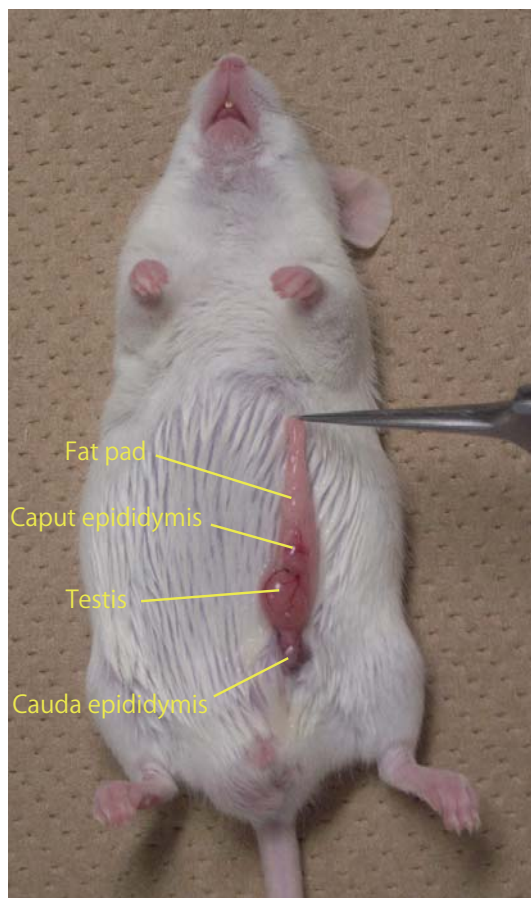
### Materials and Equipment

1. Male mice (5 weeks old)
2. Anesthetic
3. Fine scissors
4. Pair of watchmaker's #5 forceps
5. Wound clip (Autoclip 9 mm; Clay Adams Cat. No. 427631) and clip applicator (Mik-Ron Autoclip Applier; Clay Adams Cat. No. 427630)
6. Hot plate (37°C)

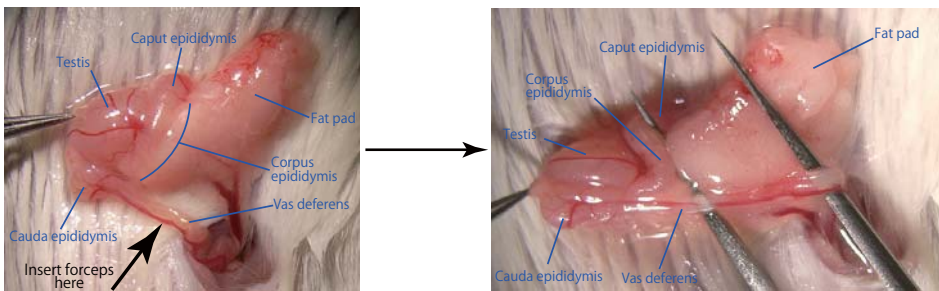
### Procedures

#### Vasectomy

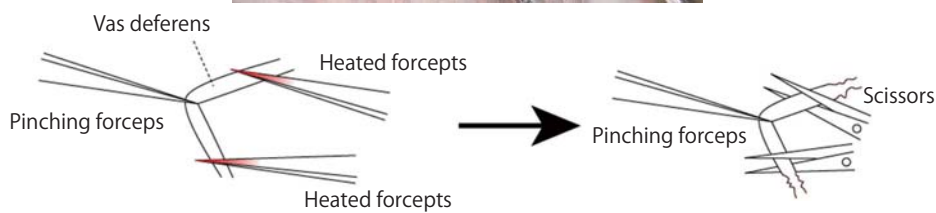
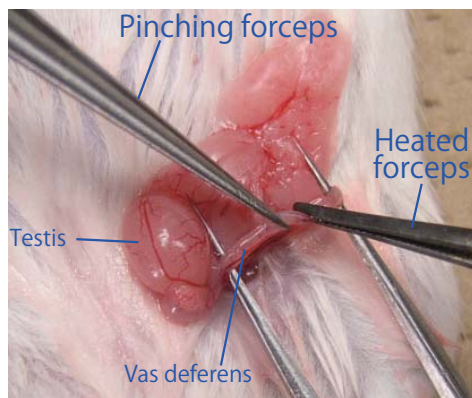
1. Anesthetize a male mouse.
2. In accordance with the conventional procedure, make a median incision. The incision should start at a point level with the top of the hind leg, and extend approximately 1 cm from this point towards the head of the mouse. After making the incision, pull out the testis, the epididymis and part of the vas deferens from the abdominal cavity.



- Pick up the vas deferens with one pair of forceps, and insert a second pair of forceps under the vas deferens to separate it from the connecting tissue.



- Hold the vas deferens with a pair of pinching forceps, then cauterize the vas deferens at two locations using a second pair of heated forceps, as shown in the diagram below.
- Cut the part of the vas deferens that lies between the two cauterized regions.



- Push the testis, epididymis and part of the vas deferens back into the abdominal cavity, then close the wound using wound clips.
- Repeat steps 2-6 for the other testis.

## References

- Nagy A., Gertsenstein M., Vintersten K., and Behringer R. 2003. Manipulating the Mouse Embryo, A Laboratory Manual (Third edition). *Cold Spring Harbor Laboratory Press*. ISBN 0-87969-591-9.

## Comment

After completing the procedure, house each mouse individually. If the mice are housed in a group, they are likely to fight and some of them may be killed. The vasectomized males will be ready for use from as early as 8 weeks of age.

## 7-2 Embryo Transfer into the Oviduct

In our laboratory, we transfer 2-cell embryos through the wall of the oviduct of pseudopregnant recipients. This procedure is much easier and simpler to conduct than the conventional embryo transfer procedure, and is therefore suitable for inexperienced users.

### Materials and Equipment

1. Female mice on Day 1 of pseudopregnancy (the day on which a vaginal plug is observed).

[Appearance of the vagina in proestrus]



Mate with  
vasectomized male

[Vaginal plug]

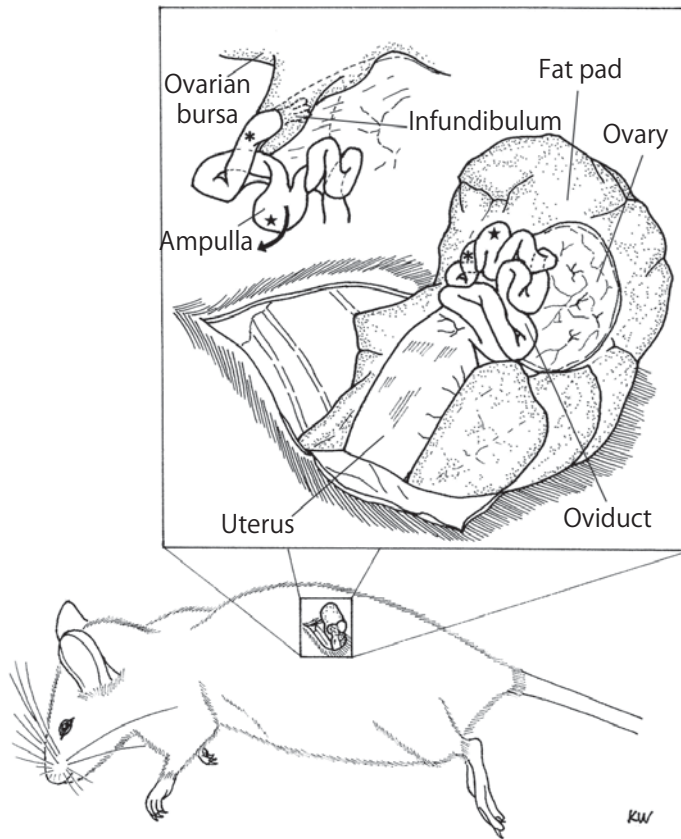


2. Anesthetic
3. Micro-spring scissors (5 mm blade)
4. Pair of watchmaker's #5 forceps
5. Serrefine clamp
6. Wound clip (Autoclip 9 mm; Clay Adams Cat. No. 427631) and clip applicator (Mik-Ron Autoclip Applier; Clay Adams Cat. No. 427630)
7. Plastic dish (35 mm X 10 mm Cat. No. 430588; CORNING)
8. Glass capillaries for embryo transfer and handling
9. Hot plate (37°C)

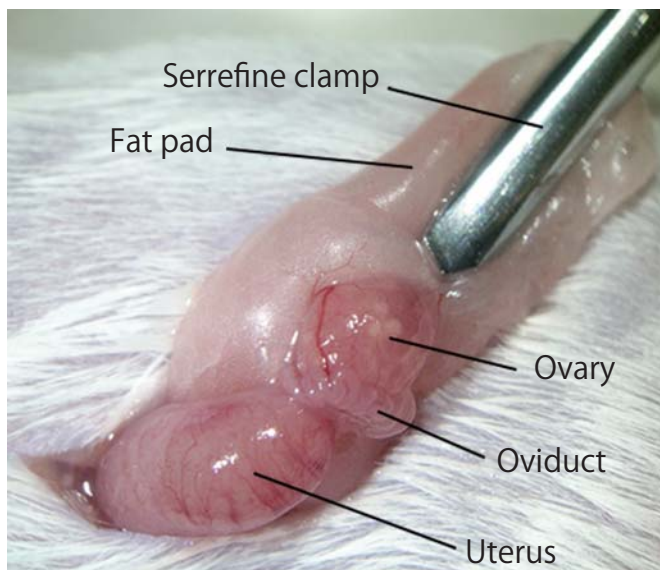
## Procedures

### Preparation of Mice

1. Anesthetize a female mouse.
2. Pull out the ovary, oviduct, and part of the uterine horn.



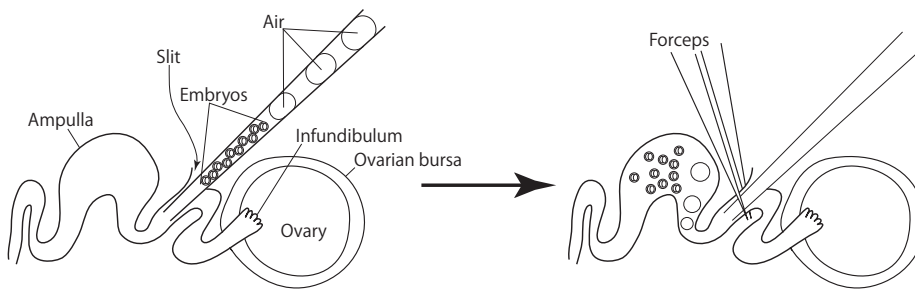
3. Clip a serrefine clamp onto the fat pad which is attached to the ovarian bursa.



## Positioning of Oviduct

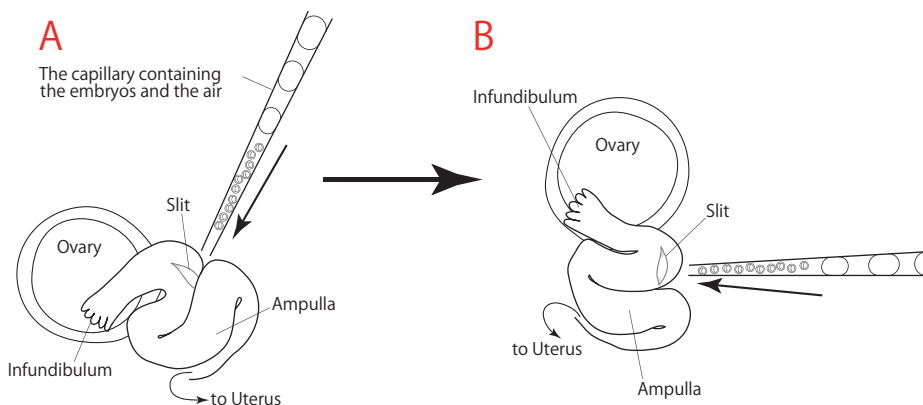
As indicated in the diagram below, embryo transfer into the oviduct is carried out by cutting the oviduct, inserting a capillary thereinto and expelling embryos towards the ampulla.

[Schematic Diagram]



Unfortunately, the oviducts of mice are small and the ducts are folded in a complicated manner, as shown in the schematic diagram of an exteriorized oviduct below (A). This makes it very difficult to insert the capillary into the oviduct towards the ampulla, because the insertion is made from above.

To make this procedure easier, position the oviduct by changing the position of the serrefine clamp and the mouse before starting the operation (B).



1. Observe the oviduct under a stereomicroscope and confirm the position of the infundibulum and ampulla using the tip of a set of forceps, or by changing the position of the serrefine clamp.
2. Position the oviduct by changing the position of the serrefine clamp and the mouse.

### Note

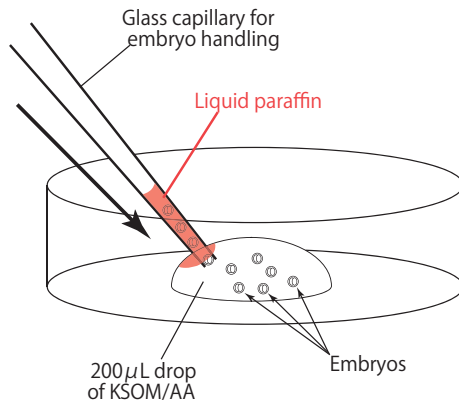
Because the folds in the oviduct vary between each mouse, look closely and adjust the position of the oviduct to make easier to work on.

### Comment

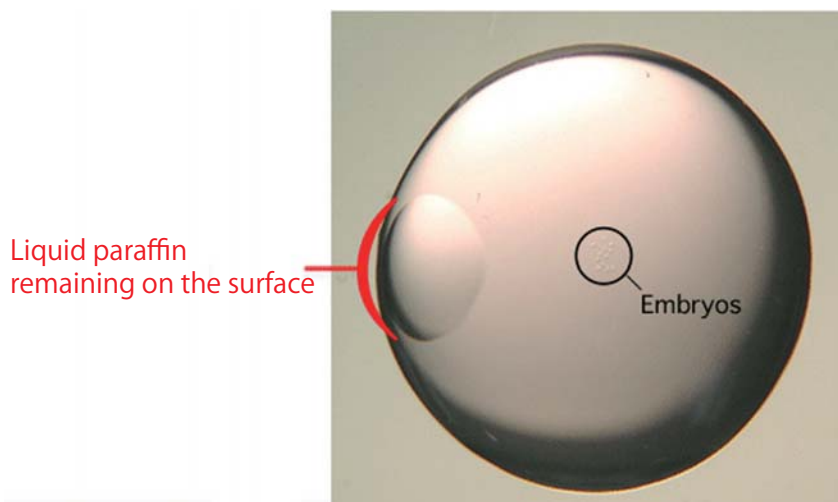
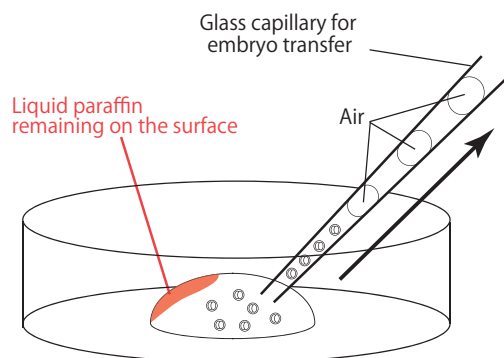
If you are left-handed, position the oviduct so that you can easily carry out the procedure with your left hand.

### Preparation of Embryos and Glass Capillary

1. Make a 200  $\mu$ L drop of KSOM/AA in a dish (without liquid paraffin), and introduce 20 embryos into the drop.



2. Aspirate air and medium in alternate intervals of 2-3 mm into a glass capillary in preparation for embryo transfer. Draw ten embryos into the glass capillary.



#### Comment

When the glass capillary is first inserted into the drop, some liquid paraffin will remain on the outer surface of the drop as shown below.

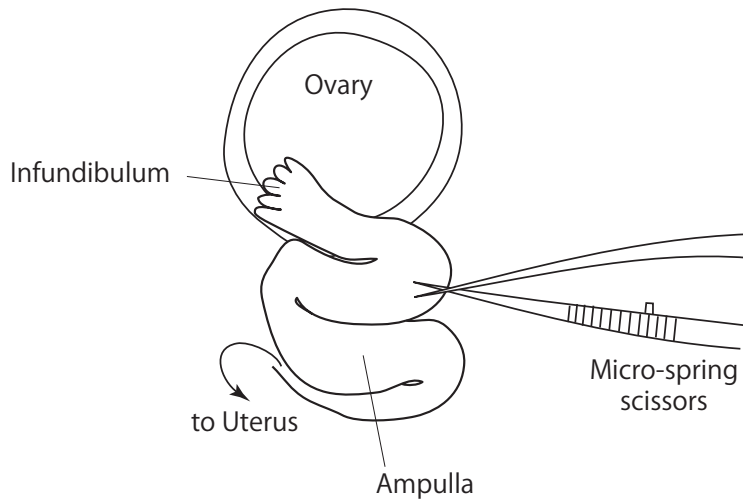
The embryos should be drawn into the glass capillary from the opposite side of the drop to avoid sucking up any liquid paraffin.

Evidence suggests that liquid paraffin which passes into the oviduct may have adverse effects on the development of the embryos into offspring.

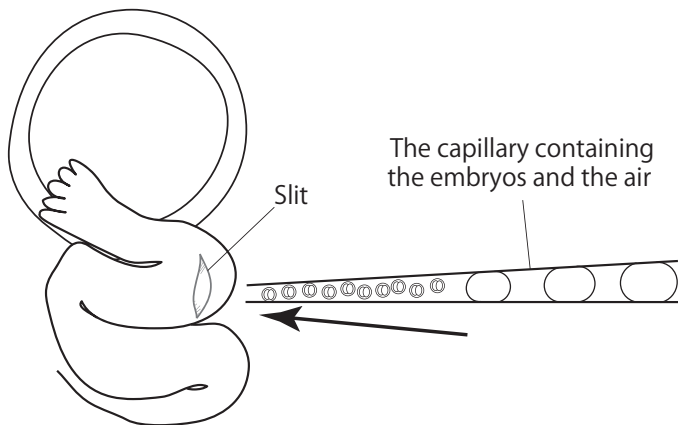


### Embryo Transfer

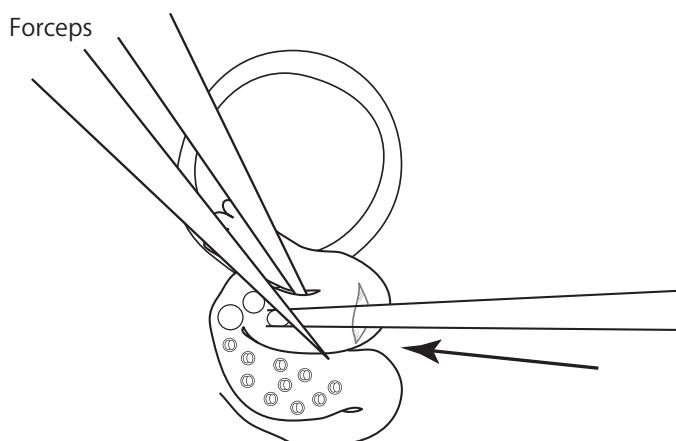
- Using a pair of watchmaker's #5 forceps and micro-spring scissors, dissect the wall of the oviduct between the infundibulum and ampulla.



- Insert the tip of the capillary containing the embryos into the slit, then push the capillary further into the slit towards the ampulla.



- Use the forceps to hold the portion of the oviduct into which the capillary was inserted.
- Expel the embryos and 2-3 of the air bubbles into the ampulla.



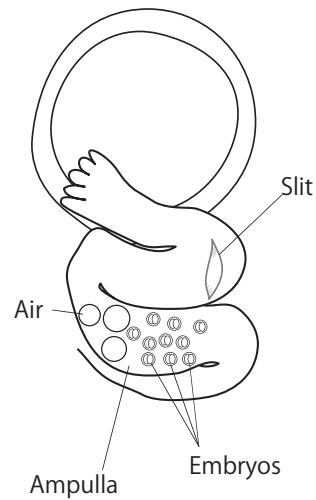
#### Comment

If performed successfully, you should be able to see air bubbles through the wall of the ampulla.

#### Note

If you cannot expel the embryos and the air bubbles into the oviduct, pull the capillary back out just a little from the slit and expel them again.

- Withdraw the capillary gently from the slit.



**[Transferring Embryos into the Oviduct]** No. 17-01 

- Push the ovary, oviduct and uterine horn back into the abdomen and close the wound using wound clips.



- Repeat the process to transfer the remaining 10 embryos into the other oviduct as described before.
- Keep the mice warm on a 37°C warming plate until they recover from the effects of anesthesia.

## References

- Nakagata N. 1992. Embryo transfer through the wall of the fallopian tube in mice. *Exp. Anim.* 41: 387-388.
- Nagy A., Gertsenstein M., Vintersten K., and Behringer R. 2003. *Manipulating the Mouse Embryo, A Laboratory Manual* (Third edition). Cold Spring Harbor Laboratory Press. ISBN 0-87969-591-9.

### Note

Transfer the embryos after adjusting the position and direction of the oviduct. If the oviduct is aligned parallel to the capillary, then it will be easier to insert the capillary into the oviduct.

## 7-3 Embryo Transfer into the Uterus

### Materials and Equipment

1. Female mice on Day 3 of pseudopregnancy (Day 1 is the day on which a vaginal plug is observed)
2. Anesthetic
3. Fine scissors
4. Pair of watchmaker's #5 forceps
5. Serrefine clamp
6. 27 gauge needle
7. Wound clip (Autoclip 9mm; Clay Adams Cat. No. 427631) and clip applicator (Mik-Ron Autoclip Applier; Clay Adams Cat. No. 427630)
8. Plastic dish (35 mm X 10 mm Cat. No. 430588; CORNING)
9. Glass capillary for embryo transfer
10. Hot plate (37°C)

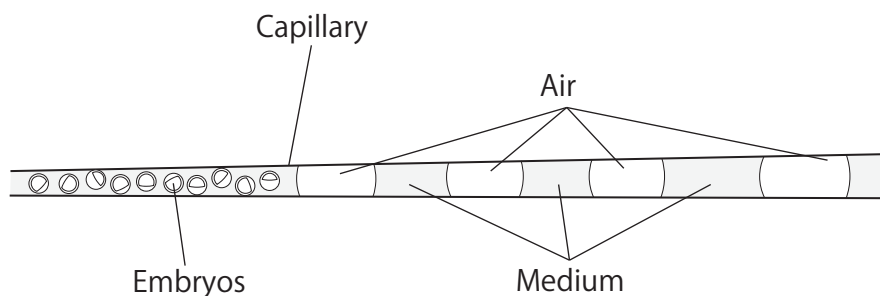
### Procedures

#### Embryo Transfer

Prepare recipient mice, embryos (8-cell to blastocyst stage) and a glass capillary as per the method used for embryo transfer into the oviduct.

(Please refer to the chapter of Embryo Transfer into the Oviduct on page 67.)

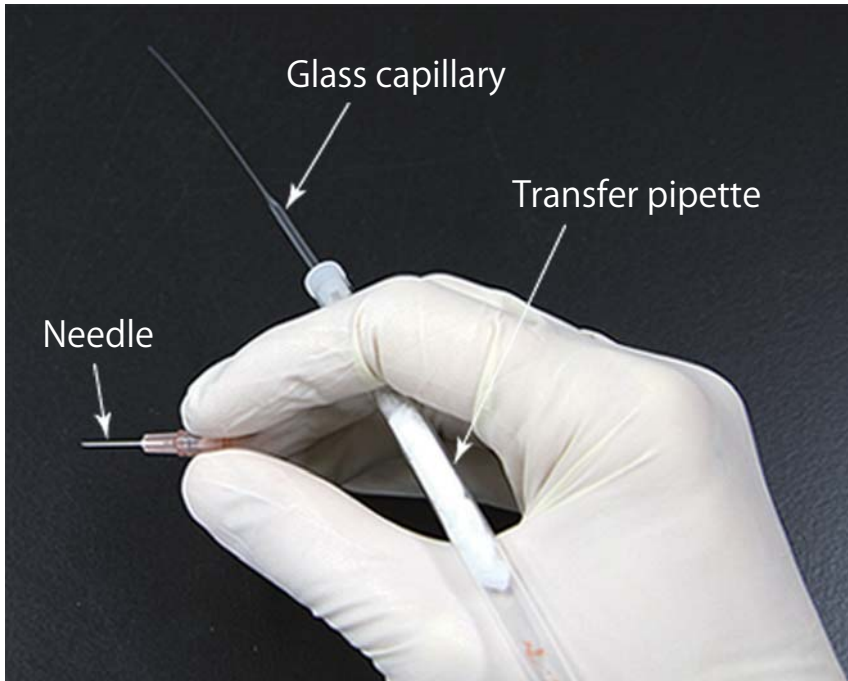
1. Pull out the ovary, oviduct and part of the uterine horn as per the conventional procedure.
2. Clip a serrefine clamp onto the fat pad which is attached to the ovarian bursa.
3. Aspirate air and medium in alternate intervals of 2-3 mm into a glass capillary in preparation for embryo transfer. Draw ten embryos into the glass capillary as indicated in the diagram below.



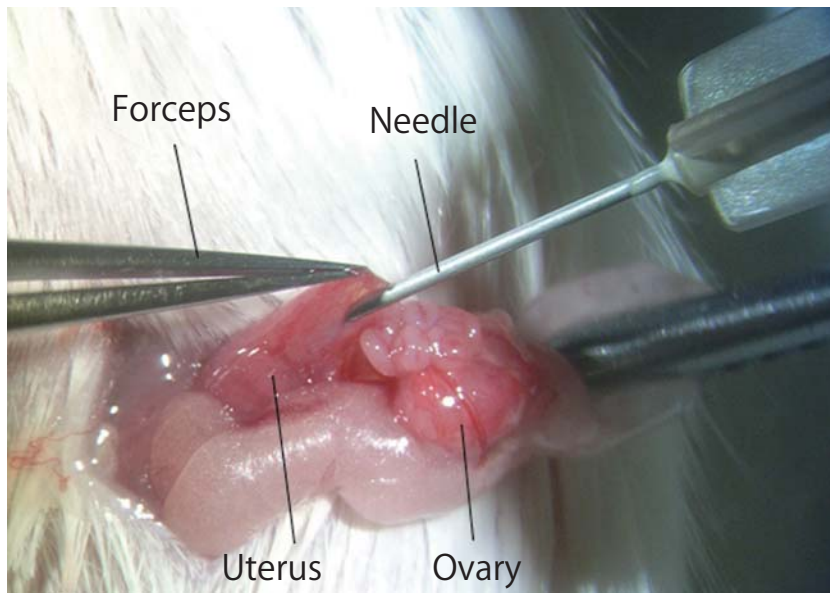
#### Note

When preparing the glass capillary, avoid placing the glass capillary into any liquid paraffin.

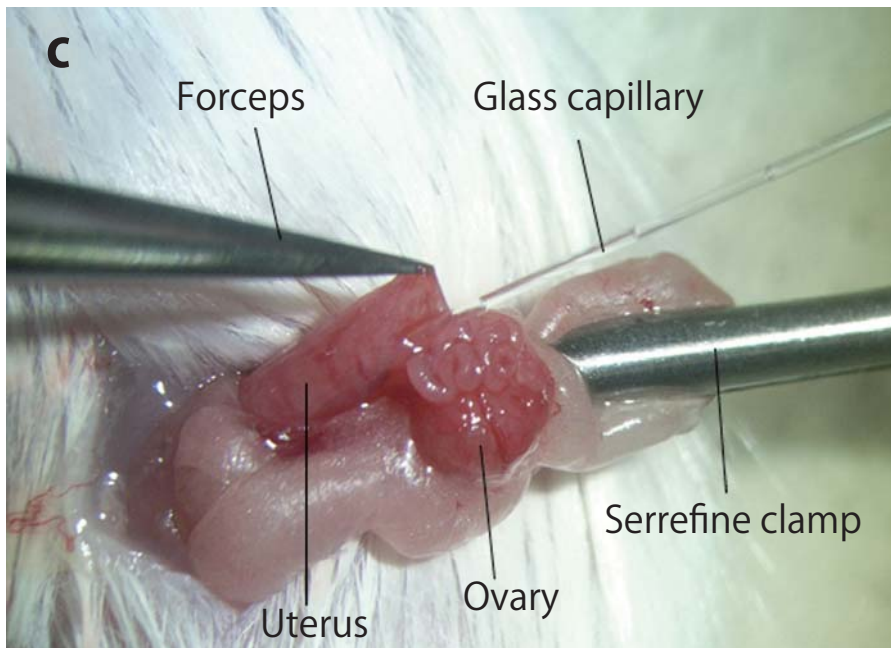
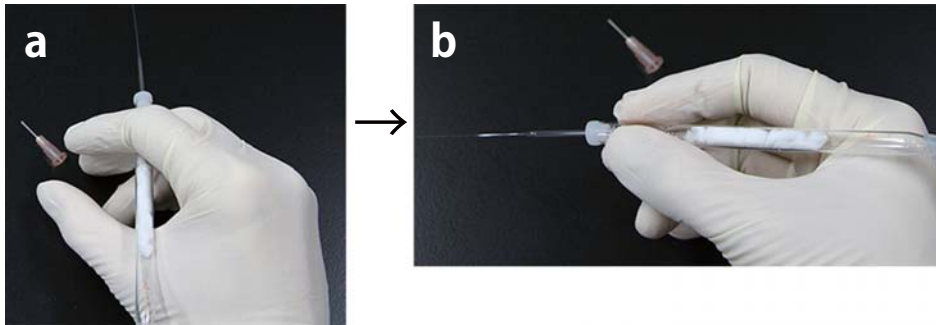
4. Hold a 27 gauge needle and the transfer pipette as shown in the picture below, and simultaneously look at both the tip of the needle and the uterus under a stereomicroscope. Be sure to look at the needle and the uterus simultaneously under the same stereomicroscope in order to confirm the position of the needle in relation to the uterus.



5. Gently hold the top of the uterine horn using fine forceps, and insert a 27 gauge needle into the wall of the uterus as far as the uterine cavity.



6. Release the needle and hold the transfer pipette as shown in diagrams a and b. Insert the tip of the capillary containing the embryos and air bubbles deep into the uterine cavity via the hole that you made with the needle, as shown in diagram c.



7. Expel the embryos into the uterine cavity along with 2-3 air bubbles.  
8. Gently withdraw the capillary from the hole.

[Embryo Transfer into the Uterus] No. 18-01 

[Performing the Operation] No. 18-02 

#### Note

You should hold onto the top of the uterine horn and keep watching the hole made by the needle until you complete embryo transfer. If you avert your eyes from the hole before completing this procedure, it may be difficult to find the hole again.

#### Note

If you cannot expel the embryos and the air bubbles into the uterine cavity, pull the capillary back out just a little from the hole and expel them again.

#### Note

To help you keep an eye on the hole made with the needle, you should hold both the needle and the transfer pipette in your dominant hand before you start the procedure.

9. Push the ovary, oviduct and uterine horn back into the abdomen and close the wound using wound clips.



10. Repeat the process to transfer 10 embryos into the other uterine cavity as described before.
11. Keep the mice warm on a 37°C warming plate until they recover from the effects of the anesthesia.

## References

1. Nagy A, Gertsenstein M., Vintersten K., and Behringer R. 2003. Manipulating the Mouse Embryo, A Laboratory Manual (Third edition). *Cold Spring Harbor Laboratory Press*. ISBN 0-87969-591-9.

## 7-4 Caesarean Section and Fostering

A Caesarean section should be performed if a pregnant recipient has not given birth to babies by the estimated date of delivery.

### Materials and Equipment

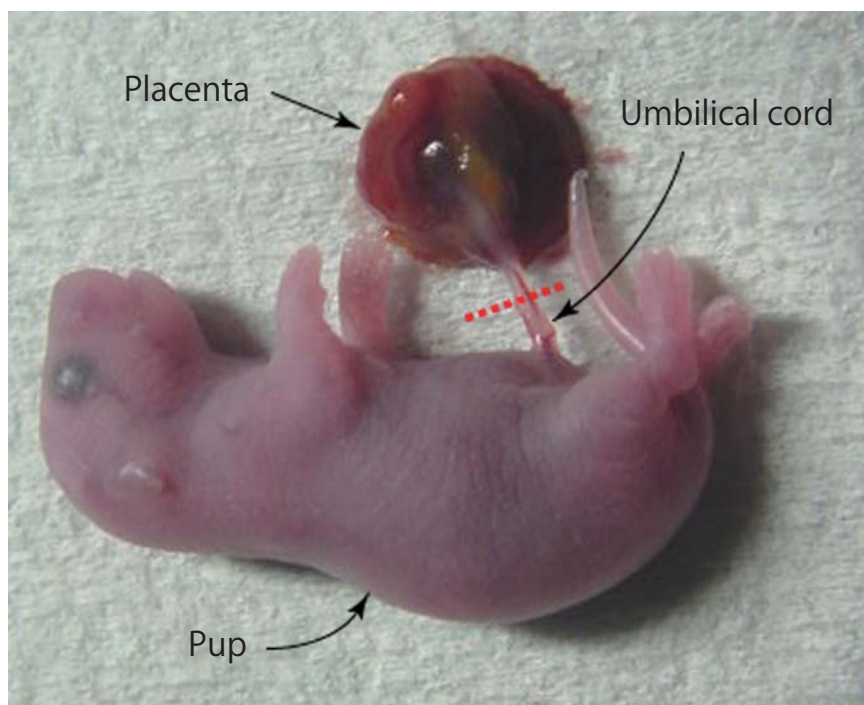
1. Foster mother (a foster mother is a female that has given birth on either the same day as, or the day preceding, the estimated date of delivery of the pregnant female)
2. Fine scissors
3. Pair of watchmaker's #5 forceps
4. Hot plate (37°C)
5. Pregnant female mouse

### Procedures

#### Caesarean Section

1. Sacrifice the pregnant female and wipe the abdomen with a piece of cotton that has been soaked thoroughly in 70% ethanol.
2. Immediately open the abdomen and remove the uteruses containing the pups with a pair of fine scissors.
3. Place the uteruses on a paper towel, and cut through the uterine wall.
4. Quickly remove the pups from the yolk sac and amnion and cut the umbilical cord.

[Cutting the Umbilical Cord]



5. Use fine tissues to wipe the amniotic fluid, secretion, and blood from the surface of the pups' bodies.
6. Place the pups on a warming plate at 37°C, and softly pinch the tail of each pup a number of times with a pair of forceps until they start to breathe and turn sufficiently pink.


**[From the Removal of the Uteruses through to the First Breaths of the Pups]**

No. 19-01 

## Fostering

Select a foster mother whose pups have a different coat color than the caesarean sectioned pups, so that you can distinguish between them later.

1. Remove the foster mother from the cage.
2. Reduce the foster mother's litter of pups by half (for example, if the litter size of the foster mother is 10, remove 4-5 pups).
3. Mix the caesarean sectioned pups to be fostered (the same number of pups as the number of pups taken away) with the bedding, then mingle them with the remaining pups of the foster mother.
4. Put the foster mother back with the pups in the original cage.

**[Fostering]** No. 19-02 

## References

1. Nagy A., Gertsenstein M., Vintersten K., and Behringer R. 2003. Manipulating the Mouse Embryo, A Laboratory Manual (Third edition). *Cold Spring Harbor Laboratory Press*. ISBN 0-87969-591-9.



## 8-1 Storage of Media and Solutions in Ampoules Under Nitrogen Gas

### Materials and Equipment

1. Twin jet ampoule sealer (Adelphi Manufacturing, West Sussex, UK)
2. Ampoule (sterilized via hot air sterilization (180°C, 3 hours))
3. Medium
4. Syringe and 18 gauge needle
5. Forceps
6. Nitrogen gas

### Procedures

#### Cleaning and Sterilizing Glass Ampoules

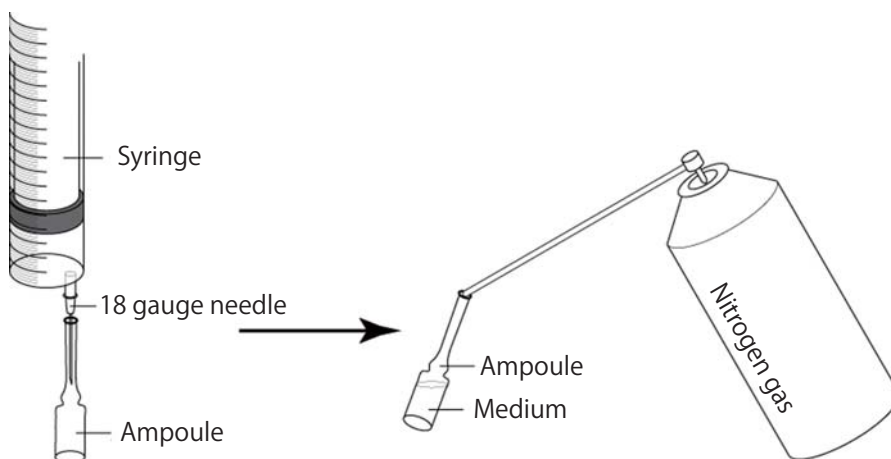
1. Rinse the glass ampoules one time using tap water.
2. Rinse the ampoules 2 times using distilled water.
3. Heat sterilize the ampoules at 180°C for at least 3 hours.

#### Ignition

1. Open the gas cock and ignite the twin jet ampoule sealer.
2. Adjust the flames of the twin jet ampoule sealer so that the flame burns blue.

#### Sealing Ampoules

1. Add an appropriate amount of medium to each ampoule.
2. Introduce nitrogen gas into one of the ampoules and immediately seal the tip of the ampoule using the flames of the twin jet ampoule sealer. Repeat for all the remaining ampoules.



[Filling Ampoules with Medium] No. 20-01 

## 8-2 Table of Media Composition

### mHTF

#### Composition of mHTF

Reagent	mg/100 mL*	Vendor	Catalog Number
NaCl	593.8	Sigma	S 5886
KCl	35.0	Sigma	P 5405
MgSO <sub>4</sub> · 7H <sub>2</sub> O	4.9	Sigma	M 2773
KH <sub>2</sub> PO <sub>4</sub>	5.4	Sigma	P 5655
CaCl <sub>2</sub>	57.0	Sigma	C 5670
NaHCO <sub>3</sub>	210.0	Sigma	S 5761
Glucose	50.0	Sigma	G 6152
Na-lactate**	0.34 mL	Sigma	L 7900
Na-Pyruvate	3.7	Sigma	P 4562
Penicillin G	7.5	Sigma	P 7794
Streptomycin	5.0	Sigma	S 1277
<b>BSA</b> (Albumin, Bovine serum Fraction V Fatty Acid-Free)	400	MERCK/ CALBIOCEM	126575
<b>0.5% phenol red</b>	<b>0.04 mL</b>	<b>Sigma</b>	<b>P 0290</b>

\*Water for embryo transfer; Sigma W1503

\*\*Assay; 70%

mHTF is enclosed in brown ampoules and stored at 4°C.

### References

1. Kito S., Hayao T., Noguchi-Kawasaki Y., Ohta Y., Hideki U., and Tateno S. 2004. Improved *in vitro* fertilization and development by use of modified human tubal fluid and applicability of pronucleate embryos for cryopreservation by rapid freezing in inbred mice. *Comp. Med.* 54(5): 564-570.

## Hyaluronidase

### Composition of Hyaluronidase

Prepare 1% solution stock as indicated below. Filter-sterilize and store in 100  $\mu\text{L}$  aliquots at  $-20^{\circ}\text{C}$ . Before use, dilute the stock by 10 times. ex.) Add 20  $\mu\text{L}$  stock solution (1%) to a 200  $\mu\text{L}$  drop of mHTF containing oocytes to make a diluted solution of about 0.1%.

Reagent	mg*	Vendor	Catalog Number
Hyaluronidase	10	Sigma	H 3506

\*mg/mL in mHTF

**0.3 M Sucrose (BSA -)****Composition of 0.3 M sucrose (BSA-)**

Reagent	mg*	Vendor	Catalog Number
Sucrose	2053.8	Sigma	S 1888

\*mg/20 mL in PB1

**Composition of PB1(BSA-)**

Reagent	mg/100 mL*	Vendor	Catalog Number
NaCl	800.0	Sigma	S 5886
KCl	20.0	Sigma	P 5405
CaCl <sub>2</sub>	12.0	Sigma	C 5670
KH <sub>2</sub> PO <sub>4</sub>	20.0	Sigma	P 5655
MgCl <sub>2</sub> · 6H <sub>2</sub> O	10.0	Sigma	M 2393
Na <sub>2</sub> HPO <sub>4</sub>	115.0	Sigma	S 5136
Na-Pyruvate	3.6	Sigma	P 4562
Glucose	100.0	Sigma	G 6152
Penicillin	7.5	Sigma	P 7794
Streptomycin	5.0	Sigma	S 1277

\*Water for embryo transfer; Sigma W1503

0.3 M Sucrose (BSA-) is enclosed in brown ampoules and stored at 4°C.

**0.3 M Sucrose (BSA +)****Composition of 0.3 M sucrose (BSA+)**

Reagent	mg*	Vendor	Catalog Number
Sucrose	2053.8	Sigma	S 1888

\*mg/20 mL in PB1

**Composition of PB1(BSA+)**

Reagent	mg/100 mL*	Vendor	Catalog Number
NaCl	800.0	Sigma	S 5886
KCl	20.0	Sigma	P 5405
CaCl <sub>2</sub>	12.0	Sigma	C 5670
KH <sub>2</sub> PO <sub>4</sub>	20.0	Sigma	P 5655
MgCl <sub>2</sub> · 6H <sub>2</sub> O	10.0	Sigma	M 2393
Na <sub>2</sub> HPO <sub>4</sub>	115.0	Sigma	S 5136
Na-Pyruvate	3.6	Sigma	P 4562
Glucose	100.0	Sigma	G 6152
Penicillin	7.5	Sigma	P 7794
Streptomycin	5.0	Sigma	S 1277
BSA	300.0	Sigma	A 4378

\*Water for embryo transfer; Sigma W1503

0.3 M Sucrose (BSA+) is enclosed in brown ampoules and stored at 4°C.

## KSOM/AA

## Composition of KSOM/AA

Reagent	mg/100 mL*	Vendor	Catalog Number
NaCl	555.0	Sigma	S 5886
KCl	18.5	Sigma	P 5405
KH <sub>2</sub> PO <sub>4</sub>	4.75	Sigma	P 5655
MgSO <sub>4</sub> · 7H <sub>2</sub> O	4.95	Sigma	M 2773
CaCl <sub>2</sub> · 2H <sub>2</sub> O	25.0	Sigma	C 7902
NaHCO <sub>3</sub>	210.0	Sigma	S 5761
Glucose	3.6	Sigma	G 6152
Na-Pyruvate	2.2	Sigma	P 4562
DL-Lactic Acid sodium salt	0.174 mL	Sigma	L 1375
10 mM EDTA	100 µL	Sigma	E 6635
Streptomycin	5.0	Sigma	S 9137
Penicillin	6.3	Sigma	P 7794
0.5% phenol red	0.1 mL	Sigma	P 0290
L-Glutamine	14.6	Sigma	G 8540
MEM Essential Amino Acids solution	1.0 mL	GIBCO	11130-051
MEM Non-essential Amino acid solution	0.5 mL	Sigma	M 7145
BSA	100.0	Sigma	A 4378

\*Water for embryo transfer; Sigma W1503  
KSOM/AA is enclosed in brown ampoules and stored at 4°C.

## References

1. Lawitts J. A., and Biggers J. D. 1993. Culture of preimplantation embryos. *Methods Enzymol.* 225:153-164.

## 0.8 M Sucrose

### Composition of 0.8 M sucrose

Reagent	mg*	Vendor	Catalog Number
Sucrose	5476.8	Sigma	S 1888

\*mg/20 mL in PB1

### Composition of PB1

Reagent	mg/100 mL*	Vendor	Catalog Number
NaCl	800.0	Sigma	S 5886
KCl	20.0	Sigma	P 5405
CaCl <sub>2</sub>	12.0	Sigma	C 5670
KH <sub>2</sub> PO <sub>4</sub>	20.0	Sigma	P 5655
MgCl <sub>2</sub> · 6H <sub>2</sub> O	10.0	Sigma	M 2393
Na <sub>2</sub> HPO <sub>4</sub>	115.0	Sigma	S 5136
Na-Pyruvate	3.6	Sigma	P 4562
Glucose	100.0	Sigma	G 6152
Penicillin	7.5	Sigma	P 7794
Streptomycin	5.0	Sigma	S 1277
BSA	300.0	Sigma	A 4378

\*Water for embryo transfer; Sigma W1503

0.8 M Sucrose is enclosed in brown ampoules and stored at 4°C.

**PB1****Composition of PB1**

Reagent	mg/100 mL*	Vendor	Catalog Number
NaCl	800.0	Sigma	S 5886
KCl	20.0	Sigma	P 5405
CaCl <sub>2</sub>	12.0	Sigma	C 5670
KH <sub>2</sub> PO <sub>4</sub>	20.0	Sigma	P 5655
MgCl <sub>2</sub> · 6H <sub>2</sub> O	10.0	Sigma	M 2393
Na <sub>2</sub> HPO <sub>4</sub>	115.0	Sigma	S 5136
Na-Pyruvate	3.6	Sigma	P 4562
Glucose	100.0	Sigma	G 6152
Penicillin	7.5	Sigma	P 7794
Streptomycin	5.0	Sigma	S 1277
BSA	300.0	Sigma	A 4378

\*Water for embryo transfer; Sigma W1503

PB1 is enclosed in brown ampoules and stored at 4°C.



## 1 M DMSO

## Composition of 1 M DMSO

Reagent	mL*	Vendor	Catalog Number
DMSO	1.56	Sigma	D 2650
PB1	18.44	-	-

\*Final volume: 20 mL

## Composition of PB1

Reagent	mg/100 mL*	Vendor	Catalog Number
NaCl	800.0	Sigma	S 5886
KCl	20.0	Sigma	P 5405
CaCl <sub>2</sub>	12.0	Sigma	C 5670
KH <sub>2</sub> PO <sub>4</sub>	20.0	Sigma	P 5655
MgCl <sub>2</sub> · 6H <sub>2</sub> O	10.0	Sigma	M 2393
Na <sub>2</sub> HPO <sub>4</sub>	115.0	Sigma	S 5136
Na-Pyruvate	3.6	Sigma	P 4562
Glucose	100.0	Sigma	G 6152
Penicillin	7.5	Sigma	P 7794
Streptomycin	5.0	Sigma	S 1277
BSA	300.0	Sigma	A 4378

\*Water for embryo transfer; Sigma W1503

1 M DMSO is enclosed in brown ampoules and stored at 4°C.

## DAP213

## Method of preparing DAP213

1. Solution A and Solution B are first prepared and are each fully dissolved.
2. Equal volumes of A and B are then mixed to form DAP213.

## Solution A

Reagent	mL*	Vendor	Catalog Number
PB1	2.3088	-	-
DMSO	3.1252	Sigma	D 2650
Propylene glycol (PG)	4.556	Sigma	134368

## Caution

The solution may become cloudy when DMSO is added.

## Solution B

Reagent	mg*	Vendor	Catalog Number
Acetamide (AA)	1181.4	Sigma	A 0500

\*mg/10 mL in PB1

## Composition of PB1

Reagent	mg/100 mL*	Vendor	Catalog Number
NaCl	800.0	Sigma	S 5886
KCl	20.0	Sigma	P 5405
CaCl <sub>2</sub>	12.0	Sigma	C 5670
KH <sub>2</sub> PO <sub>4</sub>	20.0	Sigma	P 5655
MgCl <sub>2</sub> · 6H <sub>2</sub> O	10.0	Sigma	M 2393
Na <sub>2</sub> HPO <sub>4</sub>	115.0	Sigma	S 5136
Na-Pyruvate	3.6	Sigma	P 4562
Glucose	100.0	Sigma	G 6152
Penicillin	7.5	Sigma	P 7794
Streptomycin	5.0	Sigma	S 1277
BSA	300.0	Sigma	A 4378

\*Water for embryo transfer; Sigma W1503

DAP213 is enclosed in brown ampoules and stored at 4°C.

**0.25 M Sucrose****Composition of 0.25 M sucrose**

Reagent	mg*	Vendor	Catalog Number
Sucrose	1711.5	Sigma	S 1888

\*mg/20 mL in PB1

**Composition of PB1**

Reagent	mg/100 mL*	Vendor	Catalog Number
NaCl	800.0	Sigma	S 5886
KCl	20.0	Sigma	P 5405
CaCl <sub>2</sub>	12.0	Sigma	C 5670
KH <sub>2</sub> PO <sub>4</sub>	20.0	Sigma	P 5655
MgCl <sub>2</sub> · 6H <sub>2</sub> O	10.0	Sigma	M 2393
Na <sub>2</sub> HPO <sub>4</sub>	115.0	Sigma	S 5136
Na-Pyruvate	3.6	Sigma	P 4562
Glucose	100.0	Sigma	G 6152
Penicillin	7.5	Sigma	P 7794
Streptomycin	5.0	Sigma	S 1277
BSA	300.0	Sigma	A 4378

\*Water for embryo transfer; Sigma W1503

0.25 M Sucrose is enclosed in brown ampoules and stored at 4°C.

**mWM****Composition of mWM**

Reagent	mg/100 mL*	Vendor	Catalog Number
NaCl	640.0	Sigma	S 5886
KCl	35.6	Sigma	P 5405
KH <sub>2</sub> PO <sub>4</sub>	16.2	Sigma	P 5655
MgSO <sub>4</sub> · 7H <sub>2</sub> O	29.4	Sigma	M 7774
NaHCO <sub>3</sub>	190.0	Sigma	S 5761
Glucose	100.0	Sigma	G 6152
Na-Pyruvate	2.5	Sigma	P 4562
Ca-lactate pentahydrate	46.0	Sigma	C 8356
Streptomycin	5.0	Sigma	S 1277
Penicillin G	7.5	Sigma	P 7794
0.5% phenol red	0.2 mL	Sigma	P 0290
20 mM 2-ME	10.0 µL	Sigma	M 7522
100 mM EDTA	50.0 µL	Sigma	E 6635
BSA	300.0	Sigma	A 4378

\*Water for embryo transfer; Sigma W1503  
mWM is enclosed in brown ampoules and stored at 4°C.



All video files shown in this manual are provided in the supplementary USB flash drive. If you need it or have questions, please feel free to contact us.

COSMO BIO flash drive



Front

Back

**Contact:**

Cosmo Bio Co., Ltd.

International Sales Dept.

Toyo-Ekimae Bldg., 2-20, Toyo 2-Chome, Koto-ku, Tokyo 135-0016, Japan

Tel: +81-3-56329617

Fax: +81-3-56329618

Email: [export@cosmobio.co.jp](mailto:export@cosmobio.co.jp)

Web: [www.cosmobio.com](http://www.cosmobio.com)

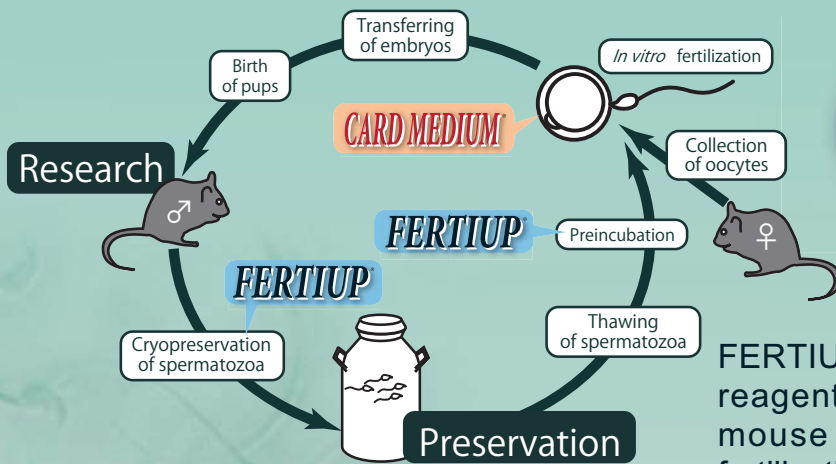
# FERTIUP®

Cryoprotectant  
Preincubation Medium

# CARD MEDIUM®

Mouse *in vitro* Fertilization Medium

**IVF Slump? FERTIUP® Will Get You Out!!**



FERTIUP® MS CPA  
KYD-001-EX



FERTIUP® MS PM  
KYD-002-EX

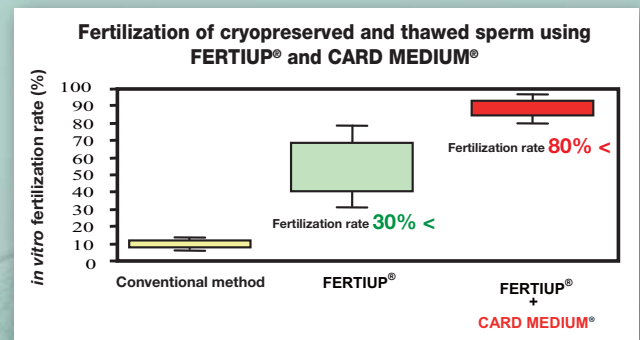


CARD MEDIUM®  
KYD-003-EX

FERTIUP® and CARD MEDIUM® are valuable reagents to improve the recovery of frozen mouse spermatozoa and improve *in vitro* fertilization efficiency of laboratory mice.

Combined usage of FERTIUP® Cryoprotectant, FERTIUP® Preincubation Medium and CARD MEDIUM® Mouse Fertilization Medium offers the following benefits:

- Fertilization rates over 80%
- Improved management of transgenic mouse
- Reduction of labor, facilities and breeding costs
- Reduction of colony expansion time
- Efficient production in difficult breeding

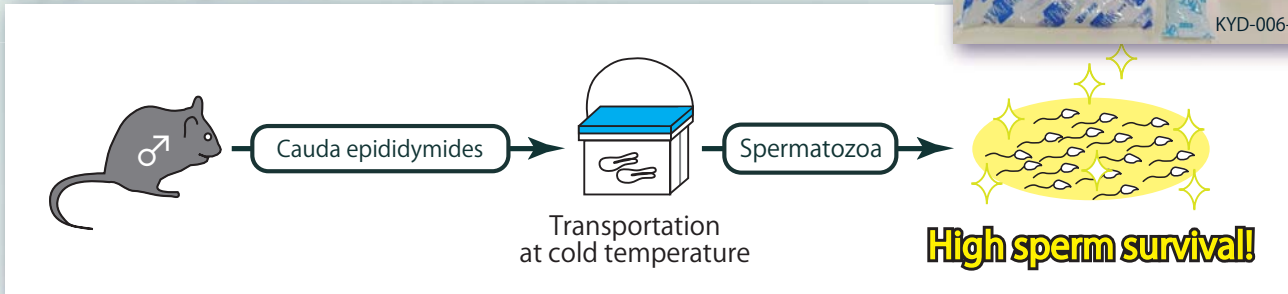


Description	Cat. No.	Quantity
<b>FERTIUP® Cryoprotectant: CPA</b>	KYD-001-EX	1 mL
	KYD-001-EX-X5	5 x 1 mL
	KYD-001-05-EX	0.5 mL
	KYD-001-05-EX-X5	5 x 0.5 mL
<b>FERTIUP® Preincubation Medium: PM</b>	KYD-002-EX	1 mL
	KYD-002-EX-X5	5 x 1 mL
	KYD-002-05-EX	0.5 mL
	KYD-002-05-EX-X5	5 x 0.5 mL
<b>CARD MEDIUM®</b> Kit includes 1 ampoule including medium (A), 1 vial including powder (B), a 1.5 mL plastic tube (C), a 1.5 mL plastic tube (D), a 2.5 mL disposable syringe, 1 needle, 1 filter unit (pore size: 0.22 µm)	KYD-003-EX	1 kit
<b>FERTIUP® PM 1ML-CARD MEDIUM® set</b> FERTIUP® Preincubation Medium: PM (1 mL) x 1 vial, CARD MEDIUM® x 1 kit	KYD-004-EX	1 set
<b>FERTIUP® PM 0.5ML-CARD MEDIUM® set</b> FERTIUP® Preincubation Medium: PM (0.5 mL) x 1 vial, CARD MEDIUM® x 1 kit	KYD-005-EX	1 set

## CARD COLD TRANSPORT KIT

Specially designed for inexpensive, safe transportation of mouse cauda epididymides and embryos at cold temperature.

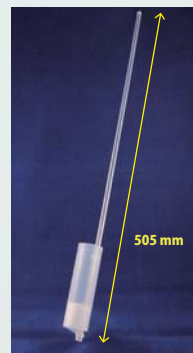
- Reduce cost for transportation of live mice
- Eliminate risk of mouse fatalities or escape during transportation
- Prevent transmission of pathogens
- Valuable for the Rescue *in vitro* Fertilization Method for Legacy Stock of Cryopreserved Spermatozoa



Description	Cat. No.	Quantity
<b>Card Cold Transport Kit</b>	KYD-006-EX	1 set
Foam transport box (1 box), Cold packs (4 large packs, 2 small packs), Thermos bottle (1 bottle), Paper box (1 box), Shock-absorbing material (1 piece)		

## FERTIUP® & CARD MEDIUM peripheral products

Description	Cat. No.	Quantity
<b>Sperm Straws (10 Pieces x 10 Units)</b>	KYD-S020X10	10 pc
<b>Freezing Canister</b>	KYD-S018	1 unit
<b>Straw Connector (5 parts included)</b>	KYD-S025	1PC
<b>Triangular Cassette Short (10units)</b>	KYD-S021	10 unit
<b>Triangular Cassette Long (10units)</b>	KYD-S035	10 unit
<b>Embryo Manipulation Instrument Set</b>	KYD-S036	1 set
<b>Glass capillary 20PC</b>	KYD-S037	1 set
<b>Touch Burner APT-3</b>	PHD-APT3-EX	10 unit

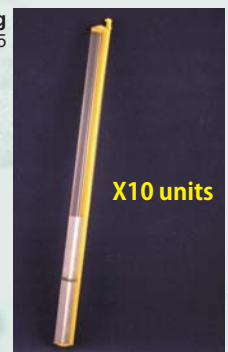


Freezing Canister  
KYD-S018



Straw Connector  
KYD-S025

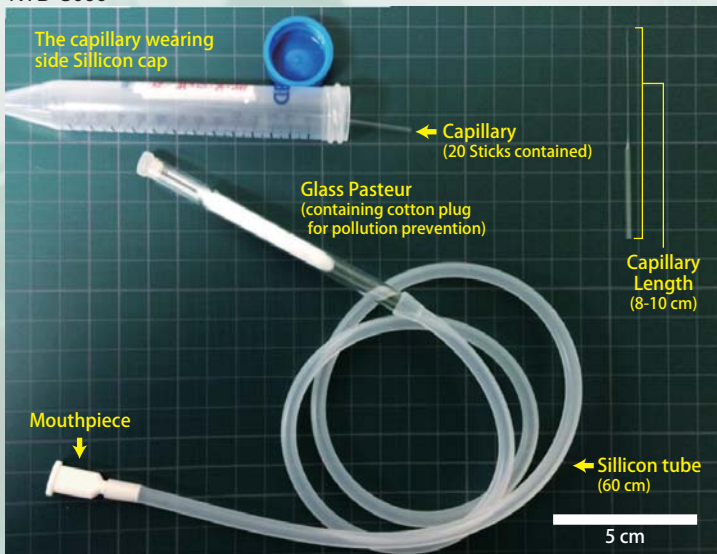
Triangular Cassette Long  
KYD-S035



Triangular Cassette Short  
KYD-S021



Embryo Manipulation Instrument Set  
KYD-S036



Sperm Straws  
KYD-S020X10



Touch Burner APT-3  
PHD-APT3-EX



# CARD HyperOva<sup>®</sup>

Enhanced Superovulation Reagent for Mouse



Live IVF pups from **A SINGLE** C57BL/6J female following superovulation with CARD HyperOva<sup>®</sup>

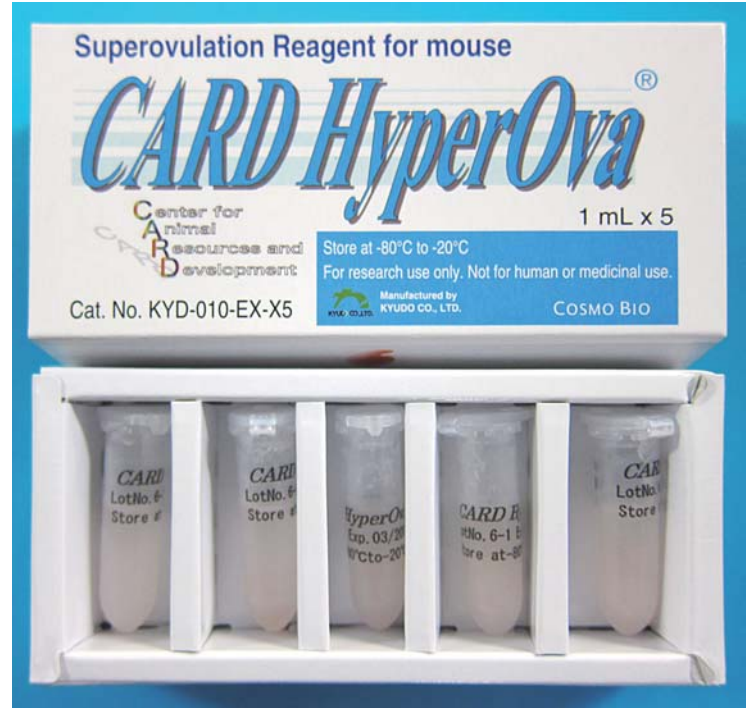
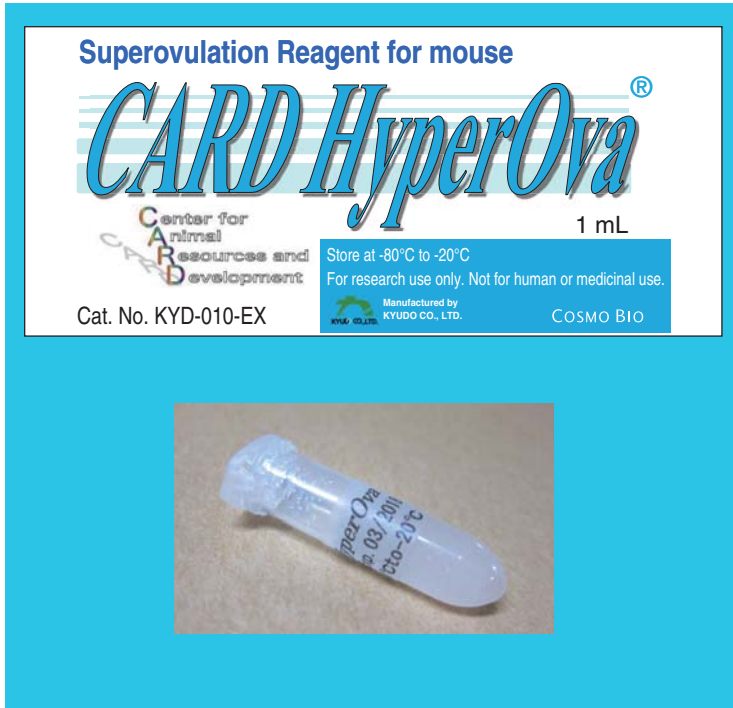
**Great for IVF.**

**Use HyperOva<sup>®</sup> to obtain more ovulated oocytes.**

Center for  
Animal  
Resources and  
Development

- Approximately 100 oocytes can be collected from a single C57BL/6J mouse.
- Maximize IVF efficiency by using HyperOva<sup>®</sup> in conjunction with FERTIUP<sup>®</sup> Mouse Sperm Cryopreservation Media, FERTIUP<sup>®</sup> Mouse Sperm Preincubation Media, and CARD MEDIUM<sup>®</sup> mouse fertilization Medium.





### Composition:

An optimized blend of anti-inhibin antibody and purified equine chorionic gonadotropin (eCG)

### Superovulation Procedure:

1. Inject 0.1-0.2 mL CARD HyperOva<sup>®</sup> i.p. into a 26-30 day old female mouse (birthdate = 0). Injections are usually performed during the light cycle, between 17:00 and 18:00.
2. At 48 hours after CARD HyperOva<sup>®</sup> recipients are injected i.p. with 7.5 IU human chorionic gonadotropin (hCG) (not included).

### References:

1. Takeo T., Nakagata N. 2015. Superovulation using the combined administration of inhibin antiserum and equine chorionic gonadotropin increases the number of ovulated oocytes in C57BL/6 female mice. *PLoS ONE* **10**(5): e0128330. doi:10.1371/journal.pone.0128330
2. Takeo T., Nakagata N. 2016. Immunotherapy using inhibin antiserum enhanced the efficacy of equine chorionic gonadotropin on superovulation in major inbred and outbred mice strains. *Theriogenol.* doi:10.1016/j.theriogenology.2016.04.076

Description	Cat. No.	Quantity	Storage
CARD HyperOva <sup>®</sup>	KYD-010-EX	1 mL	-20°C
	KYD-010-EX-X5	5x1 mL	-20°C

Shipping: Dry Ice

# Vitrification / thaw / liquids and Medium For Mouse, Rat and Embryo Manipulation

## For Mouse

Description	Application	Cat. No.	Quantity	Storage
HTF	<i>in vitro</i> fertilization	CSR-R-B071	5 mL x 10 vials	4°C
HTF	<i>in vitro</i> fertilization	CSR-R-B070	2 mL x 10 vials	4°C
mHTF	<i>in vitro</i> fertilization	KYD-008-02-EX	2 mL x 1 vial	4°C
mHTF	<i>in vitro</i> fertilization	KYD-008-02-EX-X5	2 mL x 10 vials	4°C
mHTF	<i>in vitro</i> fertilization	KYD-008-05-EX	5 mL x 1 vial	4°C
mHTF	<i>in vitro</i> fertilization	KYD-008-05-EX-X3	5 mL x 10 vials	4°C
KSOM	<i>in vitro</i> culture	CSR-R-B075	5 mL x 10 vials	4°C
KSOM	<i>in vitro</i> culture	CSR-R-B074	2 mL x 10 vials	4°C
mWM	<i>in vitro</i> culture	CSR-R-B081	5 mL x 10 vials	4°C
mWM	<i>in vitro</i> culture	CSR-R-B080	2 mL x 10 vials	4°C
0.25M sucrose	freeze-thawing	CSR-R-Y078	5 mL x 10 vials	4°C
0.25M sucrose	freeze-thawing	CSR-R-Y077	2 mL x 10 vials	4°C
1M DMSO	cryopreservation	CSR-R-T072	2 mL x 10 vials	4°C
DAP213	cryopreservation	CSR-R-T073	1 mL x 10 vials	4°C

## For Rat

Description	Application	Cat. No.	Quantity	Storage
mR1ECM	<i>in vitro</i> culture	CSR-R-M174	5 mL x 10 vials	4°C
mR1ECM	<i>in vitro</i> culture	CSR-R-M191	2 mL x 10 vials	4°C

## For Embryo Manipulation

Description	Application	Cat. No.	Quantity	Storage
M2	<i>in vitro</i> manipulation	CSR-R-M084	5 mL x 10 vials	4°C
M2	<i>in vitro</i> manipulation	CSR-R-M083	2 mL x 10 vials	4°C
PB1	<i>in vitro</i> manipulation	CSR-R-P183	5 mL x 10 vials	4°C
PB1	<i>in vitro</i> manipulation	CSR-R-P185	2 mL x 10 vials	4°C
PEPeS	cryopreservation	CSR-R-P187	1 mL x 10 vials	4°C
P10	cryopreservation	CSR-R-P186	2 mL x 10 vials	4°C

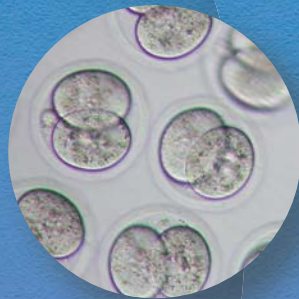


# Reproductive Engineering Techniques in Mice

Technical Manual

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Center for Animal Resources & Development (CARD)  
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