





# **AQUA-FAANG - Standard Operating Protocol - Dechorionation of carp embryos and embryo preservation**

#### **Overview**

This protocol describes a method to remove chorions from carp embryo and to preserve embryos at the desired stages. Chorions can be a source of contamination for downstream applications such as ChIP-seq and ATAC-seq, hence their removal is adviced. Carp chorions are sticky and hard and difficult to remove manually, so this step should be performed right after fertilization.

#### **Materials**

- Petri dishes (agarose-coated)
- 500 ml Beakers
- 3ml Paster pipettes

## Reagents and solutions:

- 100X E3 medium stock solution: 28,667g NaCl, 1,267g KCl, 4,833g CaCl2 2H2O, 8,167g MgSO4 7H2O in 1l of H2O. Dilute to 1X for working solution and add 0.00001% of methylene blue
- Pronase: 10 mg/ml pronase solution
- Agarose

# Preparation of agarose coated petri dishes

- 1. Prepare and dissolve 1% agarose solution in 1X E3 with 0.00001% methylene blue.
- 2. Aliquot the agarose solution into petri dishes adding enough volume to cover one-third of the heigh of the petri dishes.
- 3. Allow the agarose to polymerize. Store agarose-coated plates at 4 degrees and return to room temperature before using with embryos.

# **Dechorionation procedure**

- 1. After fertilization wait some minutes for chorions to swell up. Carp embryos are sticky so they need to be scraped out of the surface where they are deposited and collected in a petri dish. Alternatively, perform *in-vitro* fertilization directly on a petri dish and wait for the chorions to swell up.
- 2. Remove the embryo water and replace it entirely with a 10mg/ml pronase solution.
- 3. Incubate the embryos at room temperature.

- 4. **KEY STEP:** During pronase incubation check the embryos under the microscope to observe chorion softening. When some chorions have started to break, the pronase will be ready to be washed.
- 5. Fill the beakers with E3 medium and immerse the embryos <u>very gently</u> in it without letting the embryos touch the air.
- 6. Wait for the embryos to fall into the bottom and remove as much medium as possible and repeat the wash in step 5 by gently pouring the liquid through the walls of the beaker.
- 7. Repeat washing steps (5 and 6) 5 times until the pronase has been completely washed out.
- 8. Cut the tip of a pasteur pipette until the diameter is big enough to pass one embryo easily.
- 9. Fill the room-temperature agarose-coated petri dishes with E3 medium
- 10. Using the pasteur pipetter prepared in step 8, transfer the embryos to the agarose coated petri dishes making sure that just the intact embryos are selected.
- 11. Wait 10 minutes for the embryos to sit on the plate and remove any possible debreeze or spoilt embryo.
- 12. Take the dish to an incubator.

### Notes:

- The general advice is not to wait until many chorions show cracks or softening as this will lead to overdigestion and embryo death.
- ➤ If some chorions remain after this procedure, it is now easy to remove them by gently pulling them apart with dechorionation tweezers. These chorions have to be cleant out of the fish medium as they contain pronase which will affect the embryo development.
- Embryos dechorionated at the 1-cell stage should be re-checked after 5-7 hours when any dead embryos have to be removed to improve the survival of the batch.
- > E3 medium has to be changed on a daily basis.

# Recommended embryo number per replicate and downstream application

	Late	Gastrula	Early	Mid	Late	Pharyngula
	blastula		somitogenesis	somitogenesis	somitogenesis	Filalyligula
RNA-seq	20	20	10	10	5	5
ATAC-seq	20	10	10	10	5	5
ChIP-seq	40	40	20	20	15	10

# **Embryo preservation**

Techni	Preparation				
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RNA-	Embryos at the desired stages were collected in 1.5ml tubes, frozen in 500 μl of trizol and				
seq	preserved at -80°C				
ATAC-	Embryo material was prepared following the nuclear prep protocol described in				
seq	https://data.faang.org/api/fire_api/samples/UOB_SOP_Nuclear_isolation_carp_embryos_202109				
	<u>15.pdf</u>				
ChIP-	Embryos were collected in 1.5 ml tubes and prepared following the crosslinking protocol				
seq	described in				
	https://data.faang.org/api/fire_api/samples/ROSLIN_SOP_uChIPmentation_ChIPseq_of_histone_				
	marks_using_fish_embryos_20210526.pdf				