

SEP 01, 2023

OPEN ACCESS



DOI: dx.doi.org/10.17504/protocol s.io.kqdg3x93pg25/v1

Protocol Citation: Callum Howard, John B. Taggart, Caroline R. Bradley, Alejandro P. Gutierrez, John F. Taylor, Paulo A. Prodöhl, Herve Migaud, Michaël Bekaert 2023. DNA extraction from recently fertilised Atlantic salmon embryos for use in microsatellite validation of triploidy. **protocols.io** https://dx.doi.org/10.17504/p rotocols.io.kqdg3x93pg25/v1

License: This is an open access protocol distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited

## ONA extraction from recently fertilised Atlantic salmon embryos for use in microsatellite validation of triploidy

Caroline R. Callum Howard<sup>1</sup>, John B. Taggart<sup>1</sup>, Bradley<sup>2</sup>, Alejandro P. John F. Gutierrez<sup>1</sup>, Taylor<sup>1</sup>, Paulo A. Prodöhl<sup>2</sup>, Michaël

## Herve Migaud<sup>1</sup>, Bekaert<sup>1</sup>

<sup>1</sup>Institute of Aquaculture, University of Stirling, Stirling, United Kingdom; <sup>2</sup>School of Biological Sciences, Queen's University Belfast, Belfast, United Kingdom

Callum Howard: Current Address: AquaBioTech Group, Mosta, Malta; John F. Taylor: Current Address: AquaMaof Aquaculture Technologies Ltd., Rosh Ha'ayin, Israel Herve Migaud: Current Address: Mowi Scotland, Glen Nevis Business Park, Fort William

Herve Migaud: Current Address: Mowi Scotland, Glen Nevis Business Park, Fort William, United Kingdom

Michaël Bekaert: michael.bekaert@stir.ac.uk

# Michaël Bekaert

### ABSTRACT

The current methods used for producing triploid Atlantic salmon are generally reliable but not infallible, and each batch of triploids must be validated to ensure consumer trust and licensing compliance. Microsatellites have recently been shown to offer a cheaper and more convenient alternative to traditional flow cytometry for triploidy validation in a commercial setting. However, incubating eggs to at least the eyed stage for microsatellite validation poses challenges, such as reduced quality and performance of triploids produced from later eggs in the stripping season. To address these issues, we propose another option: extracting DNA from recently fertilised eggs for use in conjunction with microsatellite validation. To achieve this, we have developed an optimized protocol for HotSHOT extraction that can rapidly and cheaply extract DNA from Atlantic salmon embryos, which can then be used for triploidy validation through microsatellites. Our approach offers a simpler and more cost-effective way to validate triploidy, without the need for skilled dissection or expensive kits.

#### Protocol status: Working

We use this protocol and it's working

Created: Aug 08, 2023

Last Modified: Sep 01, 2023

**PROTOCOL integer ID:** 86092

**Keywords:** DNA extraction, DNA quality and quantity assessments, Microsatellites validation assessment, Egg, Tiploid, Salmon

#### MATERIALS

#### Consumables

Low throughput:

1.5 mL Screw cap tube

High throughput:

- 96-well Clear Round Bottom 2 mL Polypropylene Deep Well Plate
- 96-well Deep well plate seals

#### Reagents

- NaOH
- EDTA
- Tris-HCl 5 mM pH 8
- Tris-HCl dry
- TAE buffer
- MyTaq HS mix (Bioline, USA)
- Loading dye (ThermoFisher Scientific, UK)
- WellRED size standard (Eurofins, Germany)
- Gel electrophoresis reagents
- 100% ethanol
- ddH<sub>2</sub>O

#### Lab Equipment

- Forceps
- Beakers
- Heat block or laboratory oven
- Centrifuge (capable of 20,000 g)
- Gel electrophoresis machine
- PCR machine

#### **Reagent preparation**

For 200 mL each alkaline lysis reagent and neutralisation buffer (enough for 500 samples).

Alkaline Lysis Reagent

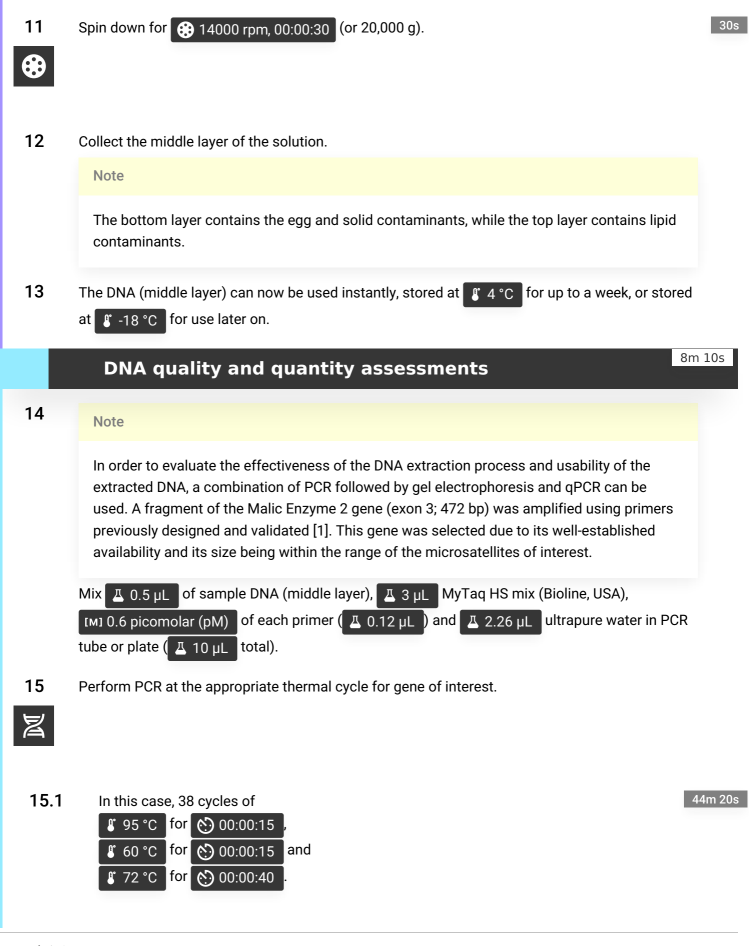
Reagent	Final conc.	Amount for 200 mL
Na0H	25 mM	200 mg
EDTA	0.2 mM	14.88 mg

*Add ddH20 for final volume of 200 mL. pH will be 12.* Neutralisation Buffer

ReagentFinal conc.Amount for 200 mLTris-HCl40 mM1.3 g

Add ddH20 for final volume of 200 mL. pH will be 5.

1	If eggs stored in ethanol, remove using forceps and place on clean tissue to remove excess ethanol.	
2 	Place embryos in a beaker of Tris-HCl (тмз 5 millimolar (mM) , Эн 8) for 👏 00:15:00.	15m
3	Remove the eggs and remove excess liquid with clean tissue.	
4	For low throughput needs the eggs can then be placed into individual 1.5 mL screw cap tubes, for high throughput needs the eggs can be places, one per well, into a 2 mL deep 96-well plate.	
5	Pierce the chorion by applying pressure using the end of the forceps.	
	Note	
	Between eggs, the forceps must be wiped clean before being sterilised using 100% ethanol and ddH <sub>2</sub> O.	
6 /	Add $\underline{\mathbf{Z}}$ 400 $\mu$ L alkaline lysis buffer to each tube/well and seal.	
7	Invert 5 times, and placed into either a heat block or a laboratory oven running at 90 °C for 00:30:00	30m
8	Remove and place SON ice for OO:05:00.	5m
9 /	Unseal and add an equal amount ( Δ 400 μL ) of neutralisation buffer.	
10 X	Reseal and rapidly invert 10 times and then spin down briefly using a centrifuge.	



16	Load $\blacksquare$ 2.5 µL of the PCR product into a 1.25% agarose gel with $\blacksquare$ 5 µL of 1.5× loading dye (ThermoFisher Scientific, UK) in 0.5× TAE buffer.	
17	Migrate the gel with ethidium bromide and visualised under UV in a transilluminator for the quality of bands and the presence of smear or primer dimer.	
18	Note	
X	The qPCR reactions were run on a QTower 3 (Analytik Jena, Germany) in accordance with the manufacturer's instructions:	
	Mix 🕹 1 µL of sample DNA (middle layer), 🗳 5 µL Sensifast SYBR No-ROX kit (Bioline, USA), Implied the sense of the sense	
19 图	Perform qPCR starting by 95 °C for 00:03:00 followed by the appropriate thermal cycle for gene of interest.	3m
19.1	In this case, 40 cycles of   95 °C for   60 °C for   72 °C for   00:00:30	40m
	Microsatellites validation assessment	
20	Note	
X	A qualitative assessment of the strength of the band was used to determine the amount of PCR product to be added to the capillary electrophoresis (between 0.5 $\mu$ L and 1 $\mu$ L).	
	Mix required quantity of PCR product with $\boxed{I}_{30 \ \mu L}$ of sample loading solution (SLS), and	
	Δ 0.35 μL of size standard (WellRED size standard, Eurofins, Germany) and add to well of capillary electrophoresis plate.	
21	Top each well off with one drop of mineral oil.	
22	Run capillary electrophoresis machine (Beckman Coulter CEQ 8000, Beckman Coulter, USA) according to the manufacturer's instructions.	