

VERSION 2

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HotSHOT genomic DNA extraction V.2

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ABSTRACT

How to extract genomic DNA from larvae or finclips using the HotSHOT method.

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Materials

- **1** Prepare the BASE stock solution (50X):
 - 🛚 14.03 g KOH crystals (1.25M final concentration)
 - ☐ 4 mL of 0.5M EDTA (10 mM final concentration)
 - ddH20 to 🗸 200 mL total volume

Can be stored at room temperature for up to four years

- 2 Prepare the NEUTRALISATION stock solution (50X)
 - 🗸 63.04 g Tris-HCL (2M final concentration), also called Trizma HCl
 - ddH20 to 🗓 200 mL total volume

Can be stored at room temperature for up to four years

Procedure

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- 3 Prepare fresh 1X BASE and 1X NEUTRALISATION solution in nuclease-free H2O.
- 4 Transfer finclips or culled larvae to individual wells of a 96-well plate.

Note

(Finclips) Transfer directly to a 96-well plate during fin-clipping (Larvae) After culling:

- ullet Pipette technique: Use a pipette set to 5µL to transfer larvae with as little fish water as possible
- Forceps technique: Transfer individual larvae using blunt round-ended forceps. This technique is only possible if you have already loaded the plate with 50µL 1X BASE solution (step 5)
- 5 Add \coprod 50 μ L of 1X BASE solution into each well of the plate

7 Cool at § Room temperature

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- 8 Add \coprod 50 μ L of 1x NEUTRALISATION solution into each well of the plate.
- **9** Note, the extracted DNA concentration is often *too high* for downstream applications like PCR or KASP.

Storage

Store at 4°C if you will use the DNA in the next few weeks. Beware, the samples will slowly evaporate from a sealed plate. Alternatively, 4°C for long-term storage. This will also prevent the samples from evaporating.