



Apr 23, 2024

DNA-extraction of Daphnia and symbionts

DOI

<https://dx.doi.org/10.17504/protocols.io.5jyl82n96l2w/v1>

Pascal Angst¹, Peter D. Fields¹

¹University of Basel



Pascal Angst

University of Basel

Create & collaborate more with a free account

Edit and publish protocols, collaborate in communities, share insights through comments, and track progress with run records.

Create free account

OPEN  ACCESS



DOI: <https://dx.doi.org/10.17504/protocols.io.5jyl82n96l2w/v1>

Protocol Citation: Pascal Angst, Peter D. Fields 2024. DNA-extraction of Daphnia and symbionts. **protocols.io**
<https://dx.doi.org/10.17504/protocols.io.5jyl82n96l2w/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

Protocol status: Working

We use this protocol and it's working

Created: March 27, 2024

Last Modified: April 23, 2024

Protocol Integer ID: 97452

Keywords: DNA Extraction, Daphnia, Crustacean, extraction of daphnia, dna extraction, adult female daphnia magna, achieving hmw dna, archiving hmw dna, hmw dna, daphnia, extraction, dna, microbe

Abstract

This protocol was designed for DNA extraction of about 50 adult female *Daphnia magna*, it should also work for 1-150 animals but with adjusted reagent volumes. For achieving HMW DNA or maximizing yield, some modifications are indicated as substeps. Before DNA extraction, animals can be freed from microbes using antibiotics (https://www.evolution.unibas.ch/ebert/lab/daphnia_dna.htm) and should be dehydrated and snap-frozen with liquid nitrogen for archiving HMW DNA.

Attachments



General_DNA_Extracti...

19KB

Protocol materials

- ✕ Cell Lysis Solution **Qiagen Catalog #1045696**
- ✕ Protein Precipitation Solution **Qiagen Catalog #1045697**
- ✕ DNA Hydration Solution **Qiagen Catalog #1045698**
- ✕ SRE Kit **PacBio Catalog #102-208-300**



Tissue lysis and digest

19h 31m 5s

- 1 Add animals and 200 μ L Cell Lysis Solution **Qiagen Catalog #1045696** to a 1.5 ml tube. (See abstract for how to prepare the animals.)
- 1.1 For HMW DNA, use snap-frozen animals and pre-cool lysis solution and tube with ice.
- 2 Grind animals with a clean, DNA(ase)-free plastic pestle, matching the shape of the 1.5 ml tube to maximize tissue maceration.
- 2.1 For HMW DNA, use cold a pestle and only move it up and down 10 times (no twisting).
- 3 Add 300 μ L Cell Lysis Solution **Qiagen Catalog #1045696** and vortex shortly.
- 4 Add 20 μ L ProtK 20 mg/mL and carefully invert 25 times.
- 5 Incubate at 55 $^{\circ}$ C while shaking at 400 rpm overnight (Overnight incubation increases yield dramatically).
- 6 Put sample On ice , add 20 μ L RNase A 20 mg/mL to the cooled sample, and carefully invert 25 times.
- 7 Incubate at 37 $^{\circ}$ C while shaking at 400 rpm for 01:00:00 .
- 8 Put sample on On ice for 00:01:00 .
- 9 Add 300 μ L Protein Precipitation Solution **Qiagen Catalog #1045697** and vortex for 00:00:15 .



10 Centrifuge for 00:04:00 at 16000 x g .

4m



10.1 If the pellet is not tight, put tube On ice for 00:05:00 and go to step #10 or pre-cool centrifuge at 4 °C .

5m



11 Pipette supernatant (800 µL - 1.000 µL) to a 2 ml tube. Discard tissue.



11.1 For HMW DNA, use a 70 µm mesh.



Equipment

pluriStrainer Mini 70 µm

NAME

Cell Strainer

TYPE

pluriSelect

BRAND

43-10070-40

SKU

12 Add the same amount of isopropanol (800 µL - 1.000 µL) to the supernatant and carefully invert 25 times.

30s



12.1 For HMW DNA, carefully invert 50 times.



12.2 For maximum yield (but not HMW), use cold isopropanol and add 2 µL glycogen. Then, put the sample in the freezer for 01:00:00 .

1h



13 Centrifuge for 00:03:00 at 16000 x g .

3m





14 Discard supernatant, add  500 μL 70 % ethanol, and carefully invert until the pellet dislodges.

10s



14.1 For maximum yield (but not HMW), use cold ethanol.



15 Centrifuge for  00:01:00 at  16000 x g .

1m



16 Discard supernatant.

16.1 Apply  SRE Kit **PacBio Catalog #102-208-300** for HMW DNA and repeat step 14.-16. twice for purification.



17 Put the open tube in the vacuum centrifuge for  00:15:00 .

15m



18 Add  80 μL  DNA Hydration Solution **Qiagen Catalog #1045698** and incubate in the dark overnight. If fewer animals are being used or less DNA yield is expected, add less ( 20 μL -  50 μL)

