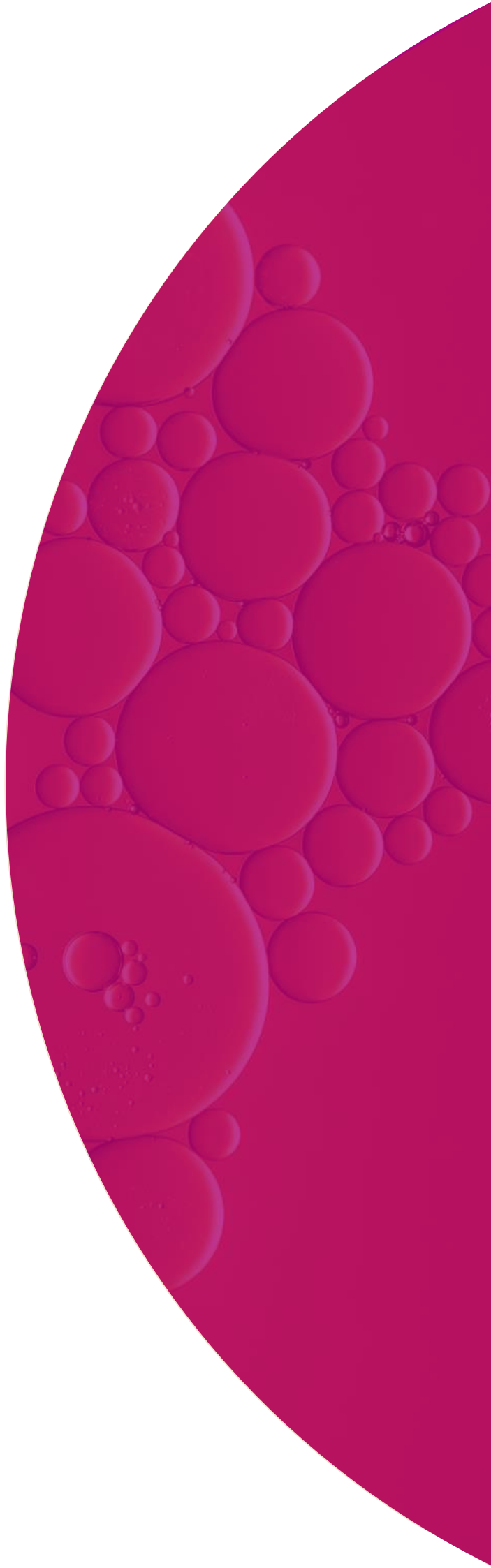


# Extracting HMW DNA from crab muscle using Nanobind<sup>®</sup> kits

Procedure & checklist



# User supplied equipment and reagent list

Equipment	Model
Nanobind® tissue kit	PacBio® (102-302-100)
Magnetic tube rack	Thermo Fisher DynaMag-2 (12321D)
Wheaton 1 mL Dounce tissue grinder with tight and loose pestles	Fisher Scientific (06-434)
Surgical scalpel	Fisher Scientific (22-079-712)
ThermoMixer	Eppendorf (5382000023)
Platform rocker	Thermo Scientific (M48725Q)
Mini-centrifuge	Ohaus Mini-Centrifuge (FC5306)
1.5 mL Protein LoBind microcentrifuge tubes*	Eppendorf (022431081)
2.0 mL Protein LoBind microcentrifuge tubes*	Eppendorf (022431102)
Wide bore 200 µL pipette tips	USA Scientific (1011-8410)
Ethanol (96–100%)	
Isopropanol (100%)	
UV/Vis	Thermo Fisher Scientific NanoDrop 2000
Fluorescent DNA quantification	Thermo Qubit 3.0, dsDNA BR and RNA BR Assay Kits

\*Eppendorf Protein LoBind tubes (Eppendorf #022431081 and #022431102) are highly recommended for all extractions to reduce protein contamination from tube carryover. Protein LoBind tubes are more effective in reducing carryover contamination than DNA LoBind tubes or other tubes and will result in improved UV purity.

## Kit storage

RNase A and Buffer CT should be stored at 4°C upon arrival.

Nanobind Disks and all other buffers should be stored at room temperature (15–30°C).

## Safety precautions

Buffer BL3 and Buffer CW1 contain guanidine hydrochloride. Warning! Guanidine hydrochloride is harmful if swallowed or inhaled and causes skin and eye irritation. DO NOT mix with bleach or acidic solutions.

## Product use

Nanobind tissue kits are intended for research use only.

# Crab muscle

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This Procedure & checklist describes an example of HMW DNA from crab muscle. Extraction data from Dungeness crab muscle is provided. This example provides sample specific details along with any modifications to the standard protocol.

For the standard Nanobind HMW tissue kit protocols, see either the [Procedure & checklist – Extracting HMW DNA from standard Dounce homogenizer tissue using Nanobind kits](#) or [Procedure & checklist – Extracting HMW DNA from animal tissue using TissueRuptor](#). Please use the [Guide & overview – Nanobind tissue kit](#) to determine which protocol is suitable for a given sample type and to find general information regarding the kit.

## Sample notes

- Crab muscle results in low to moderate yields, depending on species.

## Protocol notes

- This Procedure & checklist uses the **Nanobind tissue kit** (102-302-100).
- This Procedure & checklist describes DNA extraction from 100 mg Dungeness crab muscle using a Dounce homogenizer for disruption. This is much larger than recommend in our standard [Nanobind Tissue kit protocols](#) and is because of the low density of nuclei in crab muscle.
- TissueRuptor can also be used.

# Protocol

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1. Place the Dounce homogenizer and tight pestle on ice and chill the centrifuge to 4°C.
2. Place 100 mg Dungeness crab muscle on a clean, chilled surface, and finely mince to  $\leq 1 \text{ mm}^3$  pieces using a scalpel.
  - A plastic weigh boat cleaned with 70% EtOH can be placed on an upside-down, aluminum dry bath incubator heat block sitting in ice.
3. Transfer minced tissue to the chilled Dounce homogenizer. Keep the Dounce homogenizer on ice during the entire disruption process.
4. Add 750  $\mu\text{L}$  of cold Buffer CT.
  - Buffer CT should be placed on ice after removing from the refrigerator.
5. Gently homogenize the tissue with the pestle 10X.
  - Push the tissue with the pestle firmly into the bottom of the Dounce chamber with each stroke (Down + Up = 1X).
  - Keep the tissue between the tip of the pestle and bottom of the Dounce chamber for thorough homogenization.
  - It is not necessary to twist the pestle at the bottom of the Dounce chamber to disrupt the tissue.
  - The crab muscle homogenate became foamy. All the foam was transferred in the next step.
6. Transfer homogenate and any foam to a 2 mL Protein LoBind microcentrifuge tube.
7. Pellet homogenate by centrifuging at 1,500 x *g* and 4°C for 5 min. Discard supernatant.
8. Add 1 mL of cold Buffer CT. Pulse vortex 1 s x 10 times (max setting) to resuspend tissue.
9. Pellet homogenate by centrifuging at 1,500 x *g* and 4°C for 5 min. Discard supernatant.
10. Pulse vortex for 1 s x 2 times (max setting) to dislodge pellet.
11. Add 20  $\mu\text{L}$  of Proteinase K to the previous pellet.
12. Add 150  $\mu\text{L}$  of Buffer CLE3 and pipette mix 10X with a wide bore P200 pipette.
13. Incubate on a ThermoMixer at 55°C and 900 rpm for 30 min.
14. Spin the tube on a mini-centrifuge for 2 s to remove liquid from the cap.
15. Add 20  $\mu\text{L}$  of RNaseA.
16. Incubate on a ThermoMixer at 55°C and 900 rpm for 30 min.
17. Spin the tube on a mini-centrifuge for 2 s to remove liquid from the cap.
18. Add 60  $\mu\text{L}$  of Buffer SB and pulse vortex for 1 s x 5 times (max setting) to mix.
19. Centrifuge at 10,000 x *g* and RT (15–30°C) for 5 min.
20. Transfer up to 250  $\mu\text{L}$  of the supernatant to a new 1.5 mL Protein LoBind microcentrifuge tube using a wide bore P200 pipette. (Discard the 2 mL Protein LoBind microcentrifuge tube containing the precipitated pellet.)
  - Typical supernatant volumes will be 225–275  $\mu\text{L}$ .

## Quick tip

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Thorough tissue disruption is key to efficient lysis. It is also important to keep the tissue cold during the entire disruption process.

## Quick tip

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See Tissue Disruption Strategies section for Dounce Homogenization Tips and Tricks.

## Quick tip

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The 2 mL tube is essential for efficient lysis because of its shape; the narrow taper of a 1.5 mL tube prevents proper mixing of the lysate during thermomixing.

## Quick tip

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If there are still visible, undigested tissue pieces after step 13, the incubation may be extended up to 2 h. However, if tissue is appropriately disrupted in steps 1-5, then 30 min should be sufficient.

## Quick tip

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The narrow taper of the 1.5 mL tube is critical for proper removal of wash buffer in steps 32 & 33 and for thorough recovery of eluate in step 36.

- Crab muscle may not result in a visible pellet after this spin.
  - If the pellet is not visible, remove supernatant as if there were a pellet present and avoid pipetting from the very bottom of the tube.
21. Add 50  $\mu$ L of Buffer BL3 to the previous supernatant and inversion mix 10X.
    - Solution may become cloudy but should clear in step 24.
  22. Spin the tube on a mini-centrifuge for 2 s to remove liquid from the cap.
  23. Add Nanobind disk to lysate and add 300  $\mu$ L of isopropanol. Inversion mix 10X.
    - The Nanobind disk must be added before isopropanol.
    - A large, cloudy mass appeared upon addition of isopropanol and inversion mixing; this adhered to the Nanobind disk and became clear during the next step.
  24. Mix on a platform rocker at 20 rpm for 30 min at RT.
  25. Place tube rack on the magnetic base using the method described in the [Guide & overview – Nanobind tissue kit](#) Magnetic Rack Handling Procedure section.
  26. Discard the supernatant with a pipette using the method described in the [Guide & overview – Nanobind tissue kit](#) Pipetting section, taking care to avoid pipetting the DNA or contacting the Nanobind disk.
  27. Add 500  $\mu$ L of Buffer CW1, remove tube rack from magnetic base, inversion mix 4X, replace the tube rack on the magnetic base, and discard the supernatant.
  28. Repeat step 27.
  29. Add 500  $\mu$ L of Buffer CW2, remove tube rack from magnetic base, inversion mix 4X, replace the tube rack on the magnetic base and discard the supernatant.
  30. Repeat step 29.
  31. Pipette out any residual liquid from the tube cap.
  32. Spin the tube on a mini-centrifuge for 2 s. With the tube rack already on the magnetic base and right-side-up, place tube on tube rack and remove residual liquid.
    - If the Nanobind disk is blocking the bottom of the tube, gently push it aside with the tip of the pipette towards the magnet.
  33. Repeat step 32.
  34. Add 75  $\mu$ L of Buffer EB directly onto the Nanobind disk and incubate at RT for 10 min.
    - The Nanobind disk does not need to be fully immersed in Buffer EB – it need only be wetted and sitting atop the liquid.
  35. Collect DNA by transferring eluate to a new 1.5 mL microcentrifuge tube using a wide bore P200 pipette.
    - Either Protein LoBind or DNA LoBind tubes can be used in this step.
    - Avoid Axygen tubes as these have been shown to interfere with PacBio sequencing.
  36. Spin the tube containing the Nanobind disk on a mini-centrifuge for 5 s. Use a standard P200 pipette to combine any additional liquid that comes off the disk with the previous eluate. Repeat if necessary.
    - For crab muscle, step 36 typically has to be performed 2 times.
  37. Pipette mix 5X with a standard P200 pipette to homogenize.

### Quick tip

The Nanobind disk only needs to be wetted in the elution step: **THE DISK DOES NOT NEED TO BE FULLY SUBMERGED IN BUFFER EB.**

### Quick tip

This **5 s spin** is **CRITICAL** for recovering the DNA. We do not recommend a 2<sup>nd</sup> elution.

- Limited pipette mixing will not noticeably reduce DNA size or sequencing read lengths but is important for accurate quantitation and consistent sequencing performance.
- Take care to disrupt any regions that feel more viscous than other regions.

**38. Let eluate rest overnight at RT to allow DNA to solubilize.**

- Visible “jellies” should disperse after resting.
- The extracted HMW DNA can be heterogeneous. This is normal and is one of the challenges of working with HMW DNA. The bigger the DNA, the more this will be apparent.

**39. Following overnight rest, pipette mix 5X with a standard P200 pipette and perform triplicate NanoDrop measurements by sampling the top, middle, and bottom of the eluate.**

- If the concentration %CV exceeds 30%, 5X pipette mix with a standard P200 pipette and allow DNA to rest at RT for 1 hour to overnight. Take care to disrupt any regions that feel more viscous than other regions. Remeasure with NanoDrop.
- Pipette mixing with a standard P200 pipette will have only minor to unnoticeable effects on DNA size and sequencing read lengths.
- We routinely see A260/A280 in the range of 1.69–2.01 and A260/A230 in the range of 1.25–2.12 for Dungeness crab muscle.

**40. Use Qubit dsDNA BR Assay to determine DNA concentration.**

- We recommend making multiple measurements from the top, middle, and bottom of the eluate for an accurate DNA concentration reading.

**41. Run pulsed field gel electrophoresis (PFGE) or Agilent Femto Pulse to size the HMW DNA.**

### Quick tip

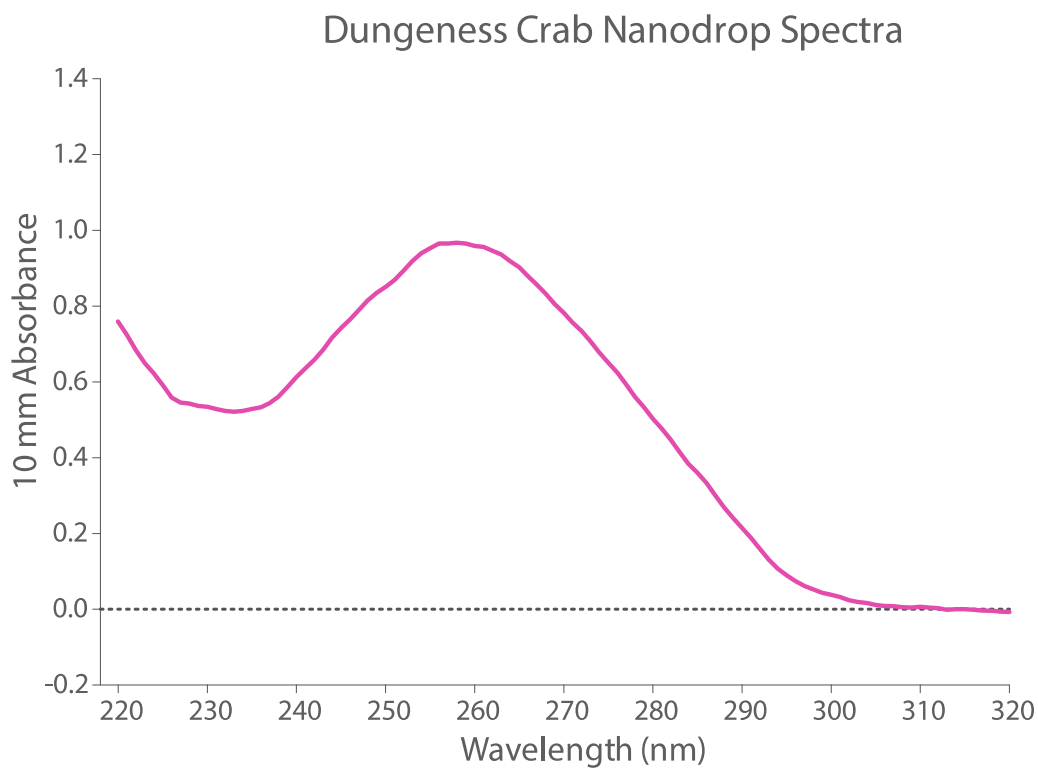
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The DNA will solubilize after resting at RT or by coaxing it into solution using standard P200 pipette mixing. For samples that need to be used immediately, we recommend needle shearing.

# DNA extraction yield and purity

- Crab muscle results in moderate extraction yields.
- Purities obtained from crab muscle are typically good.

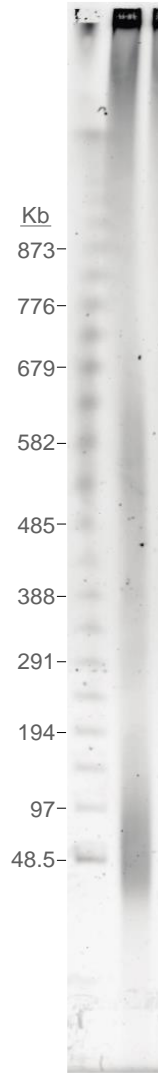
Sample	Sample input	dsDNA yield	% RNA	A260/A280	A260/A230
Dungeness crab muscle	100 mg	6.6 µg	9.8	1.85	1.79



# DNA size

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- Size of DNA extracted from crab muscle is <50–100+ kb



PFGE of DNA extracted from crab muscle.



<b>Revision history (description)</b>	<b>Version</b>	<b>Date</b>
Initial release	01	July 2022
Minor updates throughout	02	December 2022

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