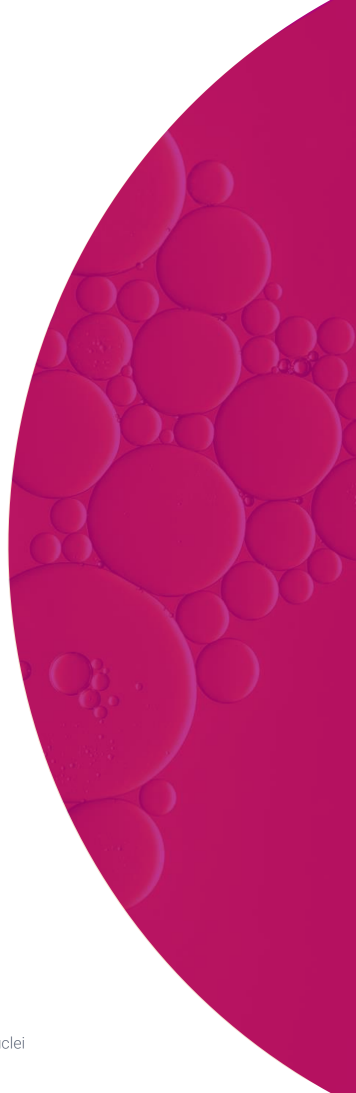


# Nanobind® HMW DNA extraction – plant nuclei

Procedure & checklist



For HMW (50 kb – 300+ kb) DNA extraction from plant nuclei for ultra long nanopore sequencing.

# Equipment and reagent list

Equipment/reagent	Manufacturer (part number)
Nanobind plant nuclei kit	PacBio® (102-302-000)
Nanobind UL library prep kit	PacBio® (NB-900-601-01)
HulaMixer	Thermo Fisher (15920D)
Magnetic tube rack	Thermo Fisher DynaMag-2 (12321D)
Mini-centrifuge	Ohaus (FC5306)
Micro-centrifuge	Eppendorf (5404000413)
1.5 mL Protein LoBind microcentrifuge tubes	Eppendorf (022431081)
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

## For all protocols

Eppendorf Protein LoBind tubes (Eppendorf #022431081) 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.

#### **Prior to starting**

Buffer PW1 is supplied as a concentrate. This kit uses PW1 with a 70% final ethanol concentration. Before using, add the appropriate amount of ethanol (96-100%) as indicated on the bottle.

## Kit storage

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

Nanobind disks and all other buffers should be stored at room temperature (18-25 °C).



#### **Product use**

Nanobind plant nuclei kits are intended for research use only.

## HMW (50 kb - 300+ kb) DNA extraction protocol

This protocol describes the extraction of HMW DNA from plant nuclei. It is recommended for HiFi sequencing.

#### Input requirements

Amount: Nuclei pellet from 1 g of plant tissue. Input will vary by tissue type.

- Prepare a plant nuclei pellet with sufficient material for 5–20 μg of isolated DNA. This typically requires 1–10 g of plant tissue.
- Samples can be pooled or diluted to yield the appropriate amount of DNA for HiFi library preparation.
- The size of the nuclei pellet will vary based on the plant species, nuclei isolation method, and amount of starting material. Typical pellets range from 10–100 µL in volume. Smaller pellet volumes are acceptable but will likely result in lower DNA yields; larger pellet volumes may require additional Proteinase K and Buffer PL1.
- This protocol has been validated on pepper leaf.
- Young leaf material typically results in the highest quality DNA and the highest ratio of DNA to input tissue
  mass.
- The resultant eluate can be pooled or diluted to obtain the correct DNA input amount for library prep.

#### HMW DNA extraction – plant nuclei

- Add 30 μL of Proteinase K to a 1.5 mL Protein LoBind tube containing plant nuclei from 1 g of tissue and gently pipette mix with a wide bore P200 pipette until the nuclei pellet is fully resuspended. Spin tube on a mini-centrifuge for 2 s to remove liquid from the cap.
  - Do not vortex the tube.
  - If pellet is not thoroughly resuspended by wide bore pipette mixing, mixing with a standard P200 pipette can be performed to homogenize the pellet without damaging high molecular weight DNA.
  - If removal of RNA is not necessary, proceed directly to step 3.
  - The input amount can be scaled from 0.5 5 g as necessary depending on plant species, age, and nuclei isolation efficiency.
- 2. Optional for removal of RNA: add 10  $\mu$ L of RNase A and gently pipette mix 5X with a wide bore P200 pipette. Incubate at RT (18–25 °C) for 3 min.
- 3. Add 80  $\mu$ L of Buffer PL1 followed by 30  $\mu$ L of Buffer CS. Gently pipette mix 10X with a wide bore P200 pipette set to 200  $\mu$ L.
  - Thorough mixing at this step is necessary to ensure complete lysis of the nuclei. Sample should appear cloudy and viscous but homogeneous.
- 4. Incubate on a ThermoMixer at 55 °C and 900 rpm for 15 min.
  - If a ThermoMixer is not available, a heat block or water bath can be used instead. Inversion mix 5X every 5 min.
- 5. Gently pipette mix 10X with a wide bore P200 pipette set to 200 μL.

## Quick tip

The nuclei suspension must be mixed with Proteinase K until it appears completely homogeneous. Insufficient mixing will result in low purity and poor yield. This typically takes 15–20 pipette mixes.



- 6. Incubate the tube at RT (18–25 °C) for 30 min. Gently pipette mix 10X with a wide bore P200 pipette set to 200  $\mu$ L every 10 min.
- 7. Centrifuge the lysate at 16,000 x g at RT (18-25 °C) for 5 min.
- 8. Transfer the supernatant to a new 1.5 mL Protein LoBind tube using a wide bore pipette.
- 9. Add a Nanobind disk to the tube followed by  $\sim$ 120  $\mu$ L (1X volume) of isopropanol. Gently invert the tube 5X, or until the supernatant-isopropanol mixture is homogeneous, and place onto a HulaMixer with the cap-side down.
  - If the volume of supernatant is greater or less than 120 μL, the volume of isopropanol should be adjusted accordingly.
  - The Nanobind disk must be added before the isopropanol.
- 10. Carefully mix with the tube-oriented cap-side down, with all fluid and the Nanobind disk contained in the cap, at RT (18–25 °C) for 20 min. Ensure that the fluid is continuously mixing and that the Nanobind disk remains submerged in the solution throughout the binding process.
  - We recommend the following HulaMixer settings:

Step	Setting	Time (s)
Rotation	9 rpm	OFF
Tilting	70°	12
Vibration	2°	1

- If a HulaMixer is not available, manually mix the tube by inversion (e.g., 5X inversions every 2-3 min) to facilitate binding.
- DNA binding is optimal when binding occurs with tube cap-side down to maximize mixing volume. A tube rotator is not recommended as there is not enough fluid for adequate mixing.
- 11. Place tubes on the magnetic tube rack.
  - Use the method described in the Nanobind big DNA kit Guide & overview "Magnetic rack handling procedure" section.
- 12. Discard the supernatant with a pipette, taking care to avoid pipetting the DNA.
  - Refer to the Nanobind DNA kit Guide & overview "Pipetting" section for details.
- 13. Add 500  $\mu$ L of Buffer PW1, remove tube rack from magnetic base, inversion mix 4X, replace tube rack on the magnetic base, and discard the supernatant.
  - Remove excess liquid from the tube cap to minimize carryover contamination.
- 14. Repeat step 13.
- 15. 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.

#### Quick tip

The Protein LoBind tubes will improve UV 260/230 ratios by up to 0.1–0.4 by preventing carryover of contaminants stuck to the tube surfaces.



- If the Nanobind disk is blocking the bottom of the tube, gently push it aside with the tip of the pipette. At this stage, DNA is tightly bound to the disk and gently manipulating the disk with a pipette tip should not cause any damage.
- 16. Repeat step 15.
- 17. Add 150 μL of Buffer EB+. Incubate at RT (18-25 °C) for 10 min.
- 18. Collect DNA by transferring eluate to a new 1.5 mL microcentrifuge tube using a wide bore P200 pipette.
  - A standard P200 pipette can be used to aid in the removal of residual liquid after most of the eluate has been removed with a wide bore pipette.
- 19. 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.
  - A small amount of liquid or gel-like material may remain on the Nanobind disk after transferring the eluate in step 18. This clear gel is DNA! The 5 s spin in step 19 will allow DNA to slide off the Nanobind disk into the bottom of the tube, after which it can be pipetted out and combined with the previously transferred eluate.
- 20. Let eluate rest at RT overnight to allow DNA to solubilize.
  - Visible "jellies" should disperse after resting.
- 21. Analyze the recovery and purity of the DNA by Nanodrop and Qubit as described in QC procedures.
- 22. Confirm triplicate Nanodrop measurements are within range to give optimal sequencing results.
  - Samples can be pooled or diluted to yield the appropriate amount of DNA for HiFi library preparation.
- 23. Proceed to library preparation. If not proceeding immediately, store the DNA at 4 °C and prepare libraries within one week.

#### Quick tip

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



# QC procedures

It is recommended that QC is performed after the DNA has been allowed to rest at RT overnight and appears homogenous under visual examination and when pipetting.

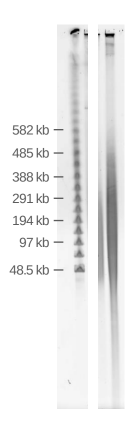
- 1. Perform triplicate NanoDrop UV/Vis measurements from top, middle, and bottom of the tube to determine total nucleic acid concentration as well as purity (A260/A280, A260/230).
  - UHMW DNA will be viscous and inhomogeneous. Take 3-5 measurements, sampling from the top, middle, and bottom of the tube, to get an accurate concentration reading. We typically see concentration %CV values of <50%. However, if the DNA is very large, the %CV can exceed 100%. In this case, take additional measurements to ensure that the concentration is accurate. Homogenization of the sample to reduce measurement CV is not required.</li>
  - If 260/280 and 260/230 ratios deviate significantly from 1.8, Nanodrop nucleic acid concentration measurements may need to be adjusted accordingly to account for contamination. UV ratios outside of this range do not necessarily indicate that sequencing will be poor.
- 2. Use Qubit dsDNA BR Assay to determine DNA concentration.
  - We recommend taking a single measurement to get an approximate DNA concentration reading.
  - We recommend the Qubit 3.0 (Thermo Fisher Scientific) with the dsDNA BR Assay kit. We do not recommend the dsDNA HS Assay kit as we have found the concentration measurements to be unreliable.
  - If Qubit DNA measurements disagree with Nanodrop measurements, we recommend using the Nanodrop data as we have found the Qubit DNA measurements to provide inconsistent readings with UHMW DNA.
- 3. Use Qubit RNA BR Assay to determine RNA concentration.
  - We recommend taking a single measurement to get an approximate RNA concentration reading.
  - We recommend the Qubit 3.0 (Thermo Fisher Scientific) with the RNA BR Assay kit.
  - If Qubit RNA measurements indicate high RNA (>50%), Nanodrop nucleic acid concentration measurements may need to be reduced to account for RNA.
  - Use Agilent Femto pulse for HMW DNA size QC.



## Results

- Nuclei isolation was performed on 5 g of pepper leaf. The resultant nuclei pellet was split into five. Five separate Nanobind extractions were performed and pooled for library preparation.
- The measurements below are of the pooled DNA (750 μL total).
- Qubit dsDNA measurements are used as a sanity check only. We find that the numbers do not accurately reflect UHMW DNA concentration even when replicate measurements are performed and go by Nanodrop instead.
- Qubit RNA measurements are used to verify that high RNA levels will not throw off Nanodrop DNA concentration measurements.

Sample	260/	260/	Nanodrop	Nanodrop	Nanodrop	Nanodrop	Qubit dsDNA	Qubit RNA
	280	230	top (ng/μL)	middle (ng/µL)	bottom (ng/µL)	avg (ng/μL)	(ng/μL)	(ng/μL)
Pepper leaf	2.7	2.2	92.7	34.4	19.1	48.7	40.7	N/A



22 hour Pulsed Field Gel Electrophoresis (PFGE) image of pepper leaf gDNA.



## Troubleshooting FAQ

#### 1. How can I increase DNA recovery?

- Adequately disrupt plant tissue prior to beginning the nuclei isolation protocol. If using LN2 grinding, grind
  the tissue for >20 min to maximize DNA recovery. If using TissueRuptor disruption, include 1 or 2 additional
  30 s disruptions to further disrupt the tissue. Inadequate tissue disruption often leads to complete failure of
  the nuclei preparation and subsequent Nanobind DNA extraction.
- Ensure that the starting nuclei pellet does not contain too much debris by selecting the proper size filter during nuclei isolation (<25 µm pore size). Improper filtration during nuclei isolation can lead to nuclei pellets which contain too much cellular debris and are exceedingly large in volume, causing poor lysis efficiency and failure of the downstream DNA extraction.
- Thoroughly resuspend the nuclei pellet during both Proteinase K and PL1 additions (steps 1 and 3). Pipette
  mixing during these steps will not substantially decrease the DNA size and can significantly increase DNA
  recovery.
- Assess the homogeneity of the eluted DNA to ensure accurate quantification. If the CV of Nanodrop measurements is high (> 50%), additional Nanodrop measurements can be taken for a more precise average.
- If above suggestions have been addressed, input tissue mass can be increased or multiple nuclei pellets can be extracted in parallel to increase DNA recovery. Some plant samples have a lower DNA to tissue ratio, particularly mature tissues; more than 1 g of starting material may be necessary to obtain adequate DNA for downstream applications.
  - o If DNA recovery is <5 µg per gram of tissue, increase the input material to up to 5 g per nuclei pellet.
  - o If DNA recovery is >5 μg per gram of tissue, prepare multiple nuclei pellets, perform DNA extraction in parallel, and combine the final eluates.

#### 2. Why are the A260/A280 or A260/A230 signals low?

- Low UV ratios do not necessarily indicate that sequencing will be poor. As long as the UV ratios are close to the stated ranges (260/260 = 1.71-2.0, 260/230 = 1.1-2.3), sequencing performance should be good.
- Poor mixing during nuclei resuspension and lysis can result in low purities. Ensure that the nuclei pellet is thoroughly resuspended during both Proteinase K and PL1 additions (steps 1 and 3). Pipette mixing during these steps will not substantially decrease DNA size and can significantly increase DNA purity and recovery.
- Insufficient washing during either nuclei isolation or DNA extraction (steps **13** and **14**) can result in lower purities. An additional wash in NIB during the nuclei isolation protocol, or an additional was in PW1 during the DNA extraction protocol (step **13**), can be included to increase DNA purity.
- Tissue input may be too high. If overall DNA recovery is adequate, sample input can be decreased to avoid overwhelming the lysis chemistry, resulting in increased purity.

#### 3. How can I recover residual DNA from the Nanobind disk after elution?

• The DNA can be eluted by spinning the tube on a mini-centrifuge for 5–10 s or centrifuging the tube at 10,000 x g for 5 s. This spin can be repeated multiple times to ensure full elution. This should not take more than 1-2 spins. We do not recommend performing a 2nd elution or heated elution as these methods dilute the eluate and can damage DNA.



- Consider lower tissue inputs. High sample input and high DNA concentration can result in DNA that is difficult to completely solubilize. For high nuclei inputs, a larger elution volume can be used to facilitate elution of the DNA.
- Do not allow bound DNA to dry after wash steps. Over-drying the Nanobind disk will lead to DNA that is difficult to elute.
- Verify that the proper amount of 100% ethanol was added to Buffer PW1 (page 2).

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