Nanobind® plant nuclei kit

Guide & overview

For extraction of HMW (50–300+ kb) genomic DNA from plant nuclei.
# Table of contents

- Prior to starting ........................................................................................................... 3
- Storage ............................................................................................................................ 3
- Product use ..................................................................................................................... 3
- For all protocols ............................................................................................................. 3
- Equipment and reagent list ........................................................................................ 4
- Introduction .................................................................................................................... 5
  - Kit overview ................................................................................................................. 5
- Workflow – nuclei isolation ....................................................................................... 6
- Workflow – Nanobind DNA extraction .................................................................... 7
- Sample information ....................................................................................................... 8
  - DNA size .................................................................................................................... 10
  - PacBio® sequencing .................................................................................................. 11
- Processing tips ............................................................................................................... 12
  - Magnetic rack handling procedure ......................................................................... 12
  - Pipetting .................................................................................................................... 13
- Nuclei isolation protocols .......................................................................................... 15
  - Nuclei isolation – TissueRuptor plant tissue (102-574-900) .................................... 15
  - Nuclei isolation – LN2 plant tissue (102-574-800) ................................................. 15
- HMW (50 kb – 300+ kb) DNA extraction protocols .................................................. 16
  - HMW DNA extraction – plant nuclei (102-574-200) .............................................. 16
- QC procedures ............................................................................................................. 16
- Storage of DNA ............................................................................................................. 16
- Troubleshooting FAQ ................................................................................................. 16
- PacBio Sequel system sequencing recommendations .............................................. 17
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.

Storage

RNase A should be stored at 4 °C upon arrival.
Nanobind disks and all other buffers should be stored at room temperature (15–30 °C).

Product use

Nanobind plant nuclei kits are intended for research use only.

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.
## Equipment and reagent list

<table>
<thead>
<tr>
<th>Equipment/reagent</th>
<th>Manufacturer (part number)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nanobind plant nuclei kit</td>
<td>PacBio® (102-302-000)</td>
</tr>
<tr>
<td>Microcentrifuge</td>
<td>Eppendorf (5415R)</td>
</tr>
<tr>
<td>HulaMixer</td>
<td>Thermo Fisher (15920D)</td>
</tr>
<tr>
<td>Magnetic tube rack</td>
<td>Thermo Fisher DynaMag-2 (12321D)</td>
</tr>
<tr>
<td>Mini-centrifuge</td>
<td>Ohaus Mini-Centrifuge (FC5306)</td>
</tr>
<tr>
<td>ThermoMixer</td>
<td>Eppendorf (5382000023)</td>
</tr>
<tr>
<td>1.5 mL Protein LoBind microcentrifuge tubes</td>
<td>Eppendorf (022431081)</td>
</tr>
<tr>
<td>Isopropanol (100%)</td>
<td></td>
</tr>
<tr>
<td>Ethanol (96–100%)</td>
<td></td>
</tr>
<tr>
<td>UV/Vis</td>
<td>Thermo Fisher Scientific NanoDrop 2000</td>
</tr>
<tr>
<td>Fluorescent DNA quantification</td>
<td>Thermo Qubit 3.0, dsDNA BR and RNA BR assay kits</td>
</tr>
<tr>
<td>26g blunt end needle</td>
<td>SAI Infusion (B26150)</td>
</tr>
<tr>
<td>1 mL syringe</td>
<td>Fisher Scientific (14-823-30)</td>
</tr>
</tbody>
</table>
Introduction

Nanobind is a novel magnetic disk covered with a high density of micro- and nanostructured silica that can be used for rapid extraction and purification of high-quality DNA and RNA. The high surface area and unique binding mechanism give it an extraordinary binding capacity, allowing isolation of high purity, high molecular weight (HMW) and ultra high molecular weight (UHMW) DNA in a microcentrifuge tube format. It uses a standard lyse, bind, wash, and elute procedure that is common for silica DNA extraction technologies. A single disk is used in each tube. However, unlike magnetic beads and silica spin columns which shear large DNA, Nanobind disks bind and release DNA without fragmentation, to yield DNA up to megabases in length.

Kit overview

The Nanobind plant nuclei kit is used for extraction of HMW (50 kb – 300+ kb) DNA from plant nuclei. First, nuclei are isolated from 1–5 g of plant tissue using one of the recommended nuclei isolation protocols. Then, HMW DNA is extracted from the nuclei using Nanobind disks. Each of the two purification steps (i.e. nuclei isolation + Nanobind extraction) removes different contaminants from the sample, resulting in clean, HMW DNA from even the most challenging plant species.

The extracted DNA is suitable for HiFi sequencing. Process time is approximately 2–3 hours for the nuclei isolation and 60 minutes for the Nanobind DNA extraction.

The Sample information section provides example extraction and sequencing results from a wide variety of plant species.
Workflow – nuclei isolation

1. **Disrupt Plant Tissue**
   - Grind plant tissue in LN2 for >20 min
   - Optional: TissueRuptor

2. **Lyse and Filter**
   - Lyse ground plant tissue
   - Centrifuge at 3000 x g (>1 Mb) or 7000 x g (< 1 Mb)

3. **Wash**
   - Wash 3X
   - Resuspend nuclei pellet using a paintbrush

4. **Pellet Nuclei**
   - Flash freeze or proceed to Nanobind DNA extraction

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Workflow – Nanobind DNA extraction

Step 1: Lysis
- Resuspend plant nuclei pellet in Proteinase K
- Lyse nuclei with Buffer PL1
- Pellet debris

Step 2: Transfer Supernatant
- Transfer supernatant to a new tube
- Add Nanobind + isopropanol
- Gentle mixing in cap
- Add wash buffer (PW1)

Step 3: Bind DNA
- Wash 2X
- Discard supernatant + add Buffer EB

Step 4: Wash
- Bluate with HMW DNA
Sample information

The following tables illustrate extraction data from a variety of plant species. All samples have been validated for PacBio sequencing.

Each sample was extracted using either the Liquid Nitrogen (LN2) and/or TissueRuptor II (TR) nuclei isolation Protocol. These protocols differ primarily in the upfront homogenization approach. For most plant species, either protocol may be used with similar results. If only a single protocol is listed, this does not indicate that the other protocol is incompatible.

Tissue disruption with TissueRuptor is faster and typically results in higher extraction yields than LN2 grinding. For some plant species, LN2 grinding may result in improved DNA size. It is recommended that users start with the TissueRuptor protocol.

Either fresh or frozen plant material can be used. Up to 5 g of plant material can be input into the LN2 protocol and up to 4 g of plant material can be put into the TissueRuptor protocol. A nuclei pellet containing up to ~20 µg of DNA can be input into each Nanobind DNA extraction process. For higher yields, parallel extractions can be performed or the Nanobind DNA extraction processes can be scaled up.

<table>
<thead>
<tr>
<th>Nanobind plant nuclei kit – validated plant tissues</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample</td>
</tr>
<tr>
<td>----------------------------------------------------</td>
</tr>
<tr>
<td>Arabidopsis thaliana²</td>
</tr>
<tr>
<td>Baby’s Breath³</td>
</tr>
<tr>
<td>Banana leaf</td>
</tr>
<tr>
<td>Brazilian Hyacinth⁴</td>
</tr>
<tr>
<td>Coastal Redwood⁵</td>
</tr>
<tr>
<td>English Holly</td>
</tr>
<tr>
<td>Giant Sequoia⁵</td>
</tr>
<tr>
<td>Lavender⁶</td>
</tr>
<tr>
<td>Melon⁷</td>
</tr>
<tr>
<td>Pepper leaf⁷</td>
</tr>
<tr>
<td>Rapeseed⁶</td>
</tr>
<tr>
<td>Rice⁴</td>
</tr>
<tr>
<td>Setaria spp.²</td>
</tr>
<tr>
<td>Spinach</td>
</tr>
<tr>
<td>Sugar beet⁷</td>
</tr>
</tbody>
</table>

LN2            Liquid nitrogen nuclei isolation protocol
TR             TissueRuptor nuclei isolation protocol
*              Denotes the protocol used to generate the data presented in the table.
1             Input levels can be scaled up or down as necessary. Optimization may be required.
2             Work performed in collaboration with Peter Thielen at Johns Hopkins University Applied Physics Laboratory.
3 Work performed in collaboration with Dr. Charlyn Partridge at Grand Valley State University.
4 Work performed in collaboration Arizona Genomics Institute at University of Arizona.
5 Work performed in collaboration with Timp Lab at Johns Hopkins University.
6 Work performed in collaboration with Buell Lab at Michigan State University.
7 Work performed in collaboration with KeyGene N.V.
Y Yes
C Customer extracted and sequenced
DNA size

The size of the genomic DNA will vary depending on plant species, the quality of the starting material, the nuclei isolation protocol used, and processing parameters during Nanobind purification. The HMW DNA extraction protocol typically yields DNA in the 50 kb – 300+ kb size range, with some samples sizing larger and some samples sizing smaller.

Figure 2. Pulsed Field Gel Electrophoresis (PFGE) illustrating DNA extracted from coastal redwood nuclei using the Nanobind Plant Nuclei Extraction Kit. The gel images show HMW DNA processed with standard (left) vs. wide-bore pipette tips (right).
PacBio® sequencing

HMW DNA was extracted from various plant samples. The samples were then sequenced on PacBio RS II, Sequel®, or Sequel II system using CLR or HiFi workflows. HMW DNA is recommended for HiFi sequencing.

### PacBio Sequel system CLR sequencing

<table>
<thead>
<tr>
<th>Sample</th>
<th>Protocol</th>
<th>Library prep</th>
<th>Polymerase length N50 (bp)</th>
<th>Subread length N50 (bp)</th>
<th>Total data (Gb)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coastal redwood1</td>
<td>LN2</td>
<td>5X NS, 30 kb BP express template 1.0</td>
<td>35,558</td>
<td>35,558</td>
<td>5.6</td>
</tr>
<tr>
<td>Rice2</td>
<td>LN2</td>
<td>No shear, 30 kb BP express template 1.0</td>
<td>31,750</td>
<td>29,750</td>
<td>8.5</td>
</tr>
<tr>
<td>Brazilian hyacinth2</td>
<td>LN2</td>
<td>No shear, 30 kb BP express template 1.0</td>
<td>31,750</td>
<td>29,750</td>
<td>11.0</td>
</tr>
<tr>
<td>Baby’s breath3</td>
<td>LN2</td>
<td>No shear, 10-50 kb BP express template 1.0</td>
<td>N/A</td>
<td>24,000</td>
<td>9.0</td>
</tr>
</tbody>
</table>

### PacBio Sequel II system HiFi sequencing

<table>
<thead>
<tr>
<th>Sample</th>
<th>Protocol</th>
<th>Library prep</th>
<th>Total bases (Gb)</th>
<th>≥Q20 mean read length (bp)</th>
<th>≥Q20 read yield (Gb)</th>
<th>≥Q20 median read quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Setaria spp4</td>
<td>TR</td>
<td>SageELF express template 2.0</td>
<td>380</td>
<td>14,894</td>
<td>26.0</td>
<td>30</td>
</tr>
<tr>
<td>Coastal redwood5</td>
<td>LN2</td>
<td>SageELF express template 2.0</td>
<td>-</td>
<td>24,397</td>
<td>23.1</td>
<td>27</td>
</tr>
</tbody>
</table>

LN2        Liquid nitrogen nuclei isolation protocol  
TR         TissueRuptor nuclei isolation protocol  
NS         Needle shear  
BP         BluePippin size selection  
1          Work performed in collaboration with Timp Lab at Johns Hopkins University  
2          Data generated in collaboration Arizona Genomics Institute at University of Arizona and PacBio®.  
3          Work performed in collaboration with Dr. Charlyn Partridge at Grand Valley State University.  
4          Work performed in collaboration with Peter Thielen at Johns Hopkins University Applied Physics Laboratory.  
5          Data generated by PacBio.
**Processing tips**

**Magnetic rack handling procedure**

To capture the Nanobind disk and enable simple processing, the microcentrifuge tubes are placed in a tube rack that is used with a magnetic base. Although DNA is bound quite robustly, proper pipetting and handling will ensure thorough washing and minimize disturbance of the bound DNA. For best results, the Nanobind disk should be captured near the top of the tube so that fluid can be easily removed from the bottom of the tube. The following procedure is recommended.

1. **Remove tube rack from magnetic base and insert tubes**
2. **Invert tube rack, ensuring Nanobind and liquid settle into lid**
3. **Place magnetic base onto inverted tube rack**
4. **Gently rotate the magnetic base**
5. **Rotate magnetic base upright, ensuring Nanobind remains captured near the top**

Recommended procedure for capturing Nanobind disk on a tube rack and magnetic base. This procedure ensures that the Nanobind disk is captured near the top of the liquid interface, minimizing disturbance of the bound DNA and facilitating processing.
Pipetting

When removing liquid from the microcentrifuge tube, the Nanobind disk should not be disturbed. Carefully insert the pipette tip against the wall opposite the Nanobind disk and remove liquid by pipetting from the liquid surface. This will minimize the chances of accidentally pipetting bound DNA. Likewise, when adding liquid, dispense against the wall opposite the Nanobind disk.

Pipetting procedure for removal (left) and addition (right) of liquid during wash steps.
Heterogeneity and viscosity (HMW DNA only)

The extracted HMW DNA can be highly viscous and heterogeneous. This is normal and is one of the challenges of working with HMW DNA. The heterogeneity and viscosity of the DNA eluate will vary depending on sample type, DNA size, sample input, and processing parameters. More gentle processing will yield larger DNA size but will also result in higher heterogeneity and larger amounts of highly viscous, unsolubilized “jellies.” Processing that is too gentle can dramatically reduce DNA purity and yield. To minimize the challenges of heterogeneity and viscosity, we recommend that new users err on the side of being overly aggressive. Listed below are tips for working with HMW DNA.

Following elution of the HMW DNA:

Pipette mix the extracted DNA 5-10X with a standard P200 pipette. Pipette mixing will help to loosen and coax the viscous DNA into solution. Moderate amounts of pipette mixing will not significantly impact DNA length. Pipette mixing is a standard part of our DNA elution process; we routinely use it for all long-read sequencing and optical mapping applications. For greater accuracy, the pipette mixed DNA should be left overnight at RT before quantifying the concentration.

In some cases, the extracted DNA will be very heterogeneous and contain large amounts of unsolubilized “jellies”:

The most common reason for high sample heterogeneity and low purity is insufficient mixing during lysis. More aggressive mixing will result in samples with improved purity due to more efficient lysis and digestion. Improved sample purity will lead to improved DNA homogeneity and reduced “jellies.” Aggressive mixing during lysis will not significantly impact DNA length. Mixing may be carefully scaled back by skilled users to achieve bigger DNA. For UHMW DNA, users should follow the appropriate UHMW DNA extraction protocol.

To accurately quantify the HMW DNA:

Pipette mix the DNA 5X with a standard P200 pipette again. Perform triplicate Nanodrop readings by sampling the top, middle, and bottom of the eluate. If the concentration %CV > 30, perform an additional 5X pipette mixing using a standard P200 pipette. Let the DNA rest for at least 1 hr and repeat the Nanodrop measurements.

To accurately determine the concentration of dsDNA, we recommend making triplicate measurements using the Qubit dsDNA BR assay.

If the extracted DNA needs to be used immediately after extraction:

The extracted DNA can be sheared 5X using a 26g blunt stainless-steel needle and 1 mL syringe. The needle-sheared DNA can be used immediately for library preparation. Moderate amounts of needle shearing will not significantly impact DNA length. Nearly all samples we sequence have been 5X needle sheared.
Nuclei isolation protocols

As of the document release date, the following protocols are available for nuclei isolation from plant samples. They have been optimized for use in downstream HMW and UHMW DNA extraction with the Nanobind plant nuclei kit (102-302-000).

Tissue disruption with TissueRuptor is faster and typically results in higher extraction yields than LN2 grinding. For some plant species, LN2 grinding may result in improved DNA size. It is recommended that users start with the TissueRuptor protocol.

The Sample information section provides additional guidance regarding protocol selection.

**Nuclei isolation – TissueRuptor plant tissue (102-574-900)**

This protocol describes the isolation of nuclei from plant tissues using QIAGEN TissueRuptor II for tissue disruption. This protocol has been validated on plants including banana, lavender, melon, pepper, rapeseed, and sugar beet.

**Nuclei isolation – LN2 plant tissue (102-574-800)**

This protocol describes the isolation of nuclei from plant tissues using liquid nitrogen grinding for tissue disruption. This protocol has been validated on plants including baby’s breath, coastal redwood, English holly, lavender, melon, pepper, rapeseed, and sugar beet.
HMW (50 kb – 300+ kb) DNA extraction protocols

As of the document release date, the following protocols are available for HMW (50–300+ kb) DNA extraction. They are recommended for most long-read sequencing applications. This includes PacBio HiFi sequencing.

HMW DNA extraction – plant nuclei (102-574-200)

This protocol describes the extraction of HMW DNA from plant nuclei. It is recommended for all standard long-read sequencing applications on PacBio instruments. This protocol has been validated on baby's breath, banana, coastal redwood, English holly, fescue, giant sequoia, lavender, melon, pepper, rapeseed, rice, and sugar beet. This protocol requires the Nanobind plant nuclei kit (102-302-000).

QC procedures

Accurate quantification of HMW and UHMW DNA can be challenging due to sample inhomogeneity, often leading to concentration measurements with high concentration CVs. We recommend performing replicate Nanodrop UV/Vis, replicate Qubit BR DNA assay measurements, and a single, optional Qubit BR RNA Assay measurement. See individual HMW and UHMW DNA extraction protocols for detailed guidance.

Storage of DNA

DNA can be stored in Buffer EB at 4 °C for several months. Long term storage at -20 °C or -80 °C can be used if necessary. Avoid freeze/thaw cycles since this can degrade high molecular weight DNA.

Troubleshooting FAQ

See individual DNA extraction protocols for details.
PacBio Sequel system sequencing recommendations

Below are our standard conditions and tips for sequencing on PacBio Sequel systems.

1. Isolate HMW DNA using the Nanobind big DNA kit.
2. Prepare DNA for sequencing using the procedure & checklist - Preparing gDNA libraries using the SMRTbell express template preparation kit SPK 3.0.