

SQK-NBD114 Post Ligation Pooling (PLP) High Yield Protocol

Overview

This protocol is built on top of the high-yield version (see SQK-NBD114 high yield protocol) and specifically adjusts the pooling strategy to reduce barcode crosstalk. In the original workflow, samples were pooled immediately after barcode addition and EDTA quench, followed by shared cleanup and adapter ligation. In this modified version, each sample is cleaned individually after barcode + EDTA, then kept separate for per-sample adapter ligation. Only after all ligation steps are complete are the libraries pooled, followed by a final SFB cleanup step. This preserves the yield benefits of the high-input–agnostic protocol while physically separating ligation reactions, thereby reducing opportunities for barcode misassignment and cross-ligation between samples.

Here's a reference-card style protocol tailored for MinION, 20 ng input per sample with SQK-NBD114.96, adapted from high yield version of ONT NBE_9171_v114_revT_03Oct2025. I've highlighted our modifications vs original ONT protocol so you can see exactly what changed.

Materials (as per ONT document unless noted)

- SQK-NBD114.96 kit (barcodes, ligation reagents, adapters)
- NEBNext® FFPE DNA Repair Mix (NEB, M6630), NEBNext® Ultra™ II End Repair/dA-Tailing Module (NEB, E7546)
- Nuclease-free water (e.g. ThermoFisher, AM9937)
- SFB Expansion (EXP-SFB001) or molecular biology grade ethanol (see note below).
- 1.5 ml Eppendorf DNA LoBind tubes, Eppendorf twin.tec® PCR plate 96 LoBind, semi skirted (Merck, EP0030129504) with heat seals
- P1000, P200, P20 and P10 pipette and tips
- Vortex mixer
- Hula mixer or rotator mixer
- Thermal cycler
- Ice bucket with ice
- Magnetic rack
- Qubit (optional for spot checks)

Note on SFB vs ethanol (EtOH)

Before ONT's revT protocol, the default post barcoding cleanup used 80% ethanol. In revT version, ONT changed to SFB washes after barcoding pooling and indicates SFB Expansion has to be purchased for this step. We evaluated both versions (pre-revT EtOH workflow, i.e. PLP protocol) and revT with SFB, i.e. PLP + SFB protocol). Yield were comparable when EtOH washes were done carefully.

1. DNA repaire and end-prep (~20 minutes)

Prepare the NEBNext FFPE DNA Repair Mix and NEBNext Ultra II End Repair / dA tailing Module reagents in accordance with manufacturer's or Nanopore's original instructions, and place on ice.

In a clean 96-well plate, aliquot each sample and negative control. Make up each sample to **6 μ l** using nuclease-free water. Mix gently by pipetting and spin down.

Combine the following components per well:

Reagent	Volume
DNA sample	6 μ l
NEBNext FFPE DNA Repair Buffer	0.4375 μ l
Ultra II End-prep Reaction Buffer	0.4375 μ l
Ultra II End-prep Enzyme Mix	0.375 μ l
NEBNext FFPE DNA Repair Mix	0.25 μ l
Total	7.5 μ l

Note on volume modification

To mitigate dead-volume losses, we use half ($0.5 \times$) of ONT's recommended volumes for all reagents while maintaining the same stoichiometries specified by ONT.

Note: We recommend mixing NEBNext FFPE DNA Repair Buffer, Ultra II End-prep Reaction Buffer, Ultra II End-prep Enzyme Mix and NEBNext FFPE DNA Repair Mix as a master mix, thoroughly mixed by pipetting, and add 1.5 μ l of the master mix for each well to simplify the loading of this step.

Between each addition, pipette mix 10-20 times.

Ensure the components are thoroughly mixed by pipetting and spin down briefly.

Using a thermal cycler, incubate at 20°C for 5 minutes and 65°C for 5 minutes.

2. Native barcoding ligation (~60 minutes)

Prepare the NEB Blunt/TA Ligase Master Mix according to the manufacturers instructions, and place on ice.

Thaw the AMPure XP Beads (AXP) at room temperature and mix by vortexing. Keep the beads at room temperature.

Thaw the EDTA at room temperature and mix by vortexing. Then spin down and place on ice.

If you plan to use SFB in the washing step, thaw SFB at room temperature and mix by vortexing. Place on ice.

Thaw the Native Barcodes (NB01-96) required for your number of samples at room temperature. Individually mix the barcodes by pipetting, spin down, and place them on ice.

Select a unique barcode for every sample to be run together on the same flow cell. Up to 96 samples can be barcoded and combined in one experiment. Only use one barcode per sample.

In a new 96-well plate, add the reagents in the following order per well mixing well by pipetting between each addition:

Reagent	Volume
End-prepped DNA	7.5 μ l
Native Barcode (NB01-96)	2.5 μ l
Blunt/TA Ligase Master Mix	10 μ l
Total	20 μ l

Note on volume modification

In this step, we doubled each reagent in the reaction system to make sure all 7.5 μl of end-prepped DNA go to the barcoding and following step.

Thoroughly mix the reaction by gently pipetting and briefly spinning down.

Incubate for 20 minutes at room temperature.

Add 4 μl EDTA (blue cap) or 2 μl EDTA (clear cap) to each well and mix thoroughly by pipetting and spin down briefly.

Add 0.4x AMPure XP Beads (AXP) to each reaction, and mix by pipetting.

Note on separate washing

We add AMPure XP Beads on each sample. For example, if you use blue cap EDTA, your sample volume is 24 μl , and you would add 9.6 μl AXP Beads to each sample. Otherwise, if you use clear cap EDTA, you would add 8.8 μl AXP Beads to each sample.

Incubate on a Hula mixer (rotator mixer) for 10 minutes at room temperature.

Spin down the sample and pellet on a magnet for 5 minutes. Keep the plate on the magnetic rack until the eluate is clear and colourless, and pipette off the supernatant.

Wash the beads with 115 μl of Short Fragment Buffer (SFB). Flick the beads to resuspend, spin down, then return the sample to the magnetic rack and allow the beads to pellet. Remove the buffer using a pipette and discard.

Note on SFB volume

In this step, we use 115 μl SFB/EtOH per sample for the wash step because the original protocol specifies a single 700 μl SFB/EtOH wash on the pooled library. In our example, we process 6 samples, so $115 \mu\text{l} \times 6 \approx 700 \mu\text{l}$, keeping the total SFB/EtOH usage comparable to the original method while performing washes separately. For other batch sizes, the SFB/EtOH volume per sample can be adjusted proportionally so that the sum across all samples approximates the 700 μl used in the original protocol.

Repeat the previous step.

Spin down and place the tube back on the magnetic rack. Pipette off any residual buffer.

Remove the tube from the magnetic rack and resuspend the pellet in 17 μl nuclease-free water by gently flicking.

Note on volume modification

Note: Here we modified 35 μl into 17 μl because the next step requires 15 μl for each barcoded sample. For those who want to check DNA concentration after washing, we leave 1 μl available for optional Qubit quantification after the washing step while maintaining sufficient volume for the next reaction. Therefore, 17 μl is selected to maximize recovery/yield without over-diluting the library.

Incubate for 10 minutes at 37°C. Every 2 minutes, agitate the sample by gently flicking for 10 seconds to encourage DNA elution.

Pellet the beads on a magnetic rack until the eluate is clear and colourless.

Remove and retain 17 μl of eluate into a clean 1.5 ml Eppendorf DNA LoBind tube.

3. Adapter ligation and clean-up (~50 minutes)

Prepare the NEBNext Quick Ligation Reaction Module according to the manufacturer's or ONT's instructions, and place on ice.

Spin down the Native Adapter (NA) and Quick T4 DNA Ligase, pipette mix and place on ice.

Thaw the Elution Buffer (EB) at room temperature and mix by vortexing. Then spin down and place on ice.

Thaw either Long Fragment Buffer (LFB) or Short Fragment Buffer (SFB) at room temperature and mix by vortexing. Place on ice.

Prepare diluted NA based on number of samples in your library as follows:

Reagent	Volume for 2 samples	for 12 samples	for 24 samples
Original NA	5 μ l	5 μ l	5 μ l
Molecular grade water	0 μ l	25 μ l	55 μ l
Total	5 μ l	30 μ l	60 μ l

In 1.5 ml Eppendorf LoBind tube for each sample, mix in the following order: Between each addition, pipette mix 10-20 times.

Reagent	Volume
Barcoded sample	15 μ l
Diluted NA	2.5 μ l
NEBNext Quick Ligation Reaction Buffer (5X)	5 μ l
Quick T4 DNA Ligase	2.5 μ l
Total	25 μ l

Note on adapter dilution and volume modification

We halve the reaction volumes at this stage because pooling is intentionally deferred until after adapter ligation to mitigate barcode crosstalk. Without volume reduction, keeping all samples separate through ligation would lead to an excessively large total volume at the subsequent pooling step. To avoid this while maintaining performance, we scale all components to 0.5 \times of the ONT-recommended volumes, preserving the original reagent ratios so that reaction stoichiometry and product quality are not affected. For the adapter, the original protocol specifies 5 μ l per batch; in this modified workflow, we dilute and redistribute the adapter so that the effective adapter amount per sample matches the reduced reaction volume and remains within ONT's intended range. This prevents overloading the library with excess adapter, which could otherwise occupy nanopores as free adapter complexes and reduce the fraction of productive, library-derived reads during sequencing.

Note

We recommend mixing diluted NA and NEBNext Quick Ligation Reaction Buffer (5X) as a master mix, thoroughly mixed by pipetting, and add 7.5 μ l of the master mix for each tube to simplify the loading of this step.

Incubate the reaction for 20 minutes at room temperature.

Pool all samples together.

Add 0.4x AMP beads, and mixed by pipetting.

All downstream steps are the same as Nanopore's recommended.

4. Priming and loading the MinION and GridION Flow Cell (~10 minutes)

All steps are the same as Nanopore's recommended.