

Jan 30, 2020

Bead-free long fragment LSK109 library preparation



In 1 collection

DOI

dx.doi.org/10.17504/protocols.io.7eshjee

John Tyson¹

¹Snutch Lab, UBC, Vancouver, BC, Canada



John Tyson

UBC

OPEN ACCESS



DOI: dx.doi.org/10.17504/protocols.io.7eshjee

External link: https://www.longreadclub.org/mountain-protocol/

Protocol Citation: John Tyson 2020. Bead-free long fragment LSK109 library preparation. protocols.io

https://dx.doi.org/10.17504/protocols.io.7eshjee

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: September 18, 2019

Last Modified: January 30, 2020

Protocol Integer ID: 27826

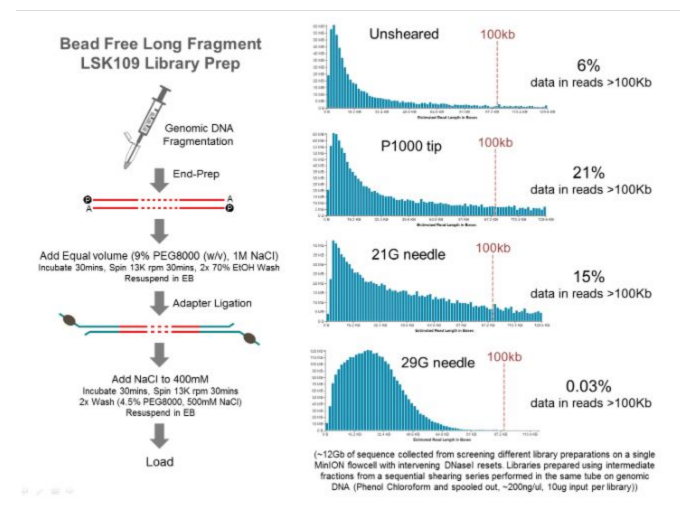
Bead-free Long Fragment LSK109 Library Preparation



Abstract

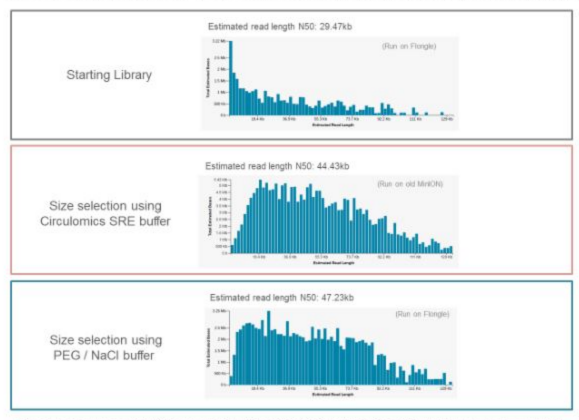
Bead-Free LSK109 ligation prep for ultra-long DNA

The use of AMPureXP beads for reaction cleanup when targeting Ultra-long DNA reads is not optimal as interbead binding results in bead clumping and potential fragmentation as individual beads move around freely in suspension. This is why the original Ultra-read method developed by Josh Quick and Nick Loman used dilution of the Rapid MuA kit and high concentrations of HMW genomic DNA. Saturating the MuA insertion reaction with HMW DNA results in less individual strand fragmentation and produces previously unattainable read lengths. It does not require cleanup or binding and elution on and off magnetic beads and minimized "handling" of the DNA. This therefore mitigates the fragmentation and DNA recovery issue with very long DNA molecules and has allowed the production of individual reads in excess of 1 million bases to be produced. The yield from individual flowcells using this approach is around an order of magnitude less however as there appears to be flowcell surface damage and rapid pore "blockage". While 1Mb+ reads are amazing to see, they are a very small fraction of the total and it is the read N50 that is important. Targeting 100kb+ reads in volume goes a long way to producing substantial improvements in large genome assemblies particularly when trying to span large repeats or low complexity regions refractory to other sequencing technologies. We have been experimenting with DNA size fractionation / precipitation using PEG/NaCl as mentioned above. This provides a means to remove the requirement for magnetic beads with their associated issues, and allows a higher yield of ultra-long reads to be produced. We have found that the DNA produced from the Phenol/Chloroform prep above provides DNA of a size that requires shearing down into a range to provide a mix of length and yield. Shown below is data from a single flowcell that I ran multiple libraries on looking at the effects of sequential shearing to gradually shorter lengths using a single DNA sample and where aliquots were removed before the next smaller shear was performed. Using a P1000 pipette tip or simple needle shear and PEG/NaCl precipitations, you are able to produce up to 21 % of total sequenced data in 100kb+ reads. Combined with DNasel flowcell clears, high yields are looking obtainable. It is early days for this protocol so expect changes to come guickly as we try and dial in better size selection and an even higher proportion of 100kb+ reads. At present we are not really seeing much short read depletion and I think this is to do with suboptimal conditions used for the DNA precipitation. Unknown buffer components in the End-Prep and Ligation buffers appear to be throwing off the PEG/NaCl parameters we have identified for size selection so there is still work to be done to tune this and something we are working on. For the 21G and 29G examples below we did do a final PEG/NaCl precipitation of the finished library but didn't see consistent depletion comparing the two, so still things to be worked out for consistency and to understanding what is happening.



As a fall back, assuming little size selection can be implemented during the prep for now, we looked again at the size selection step at the end of the library preparation when eluted into EB buffer (no other salts or other components to throw size selective precipitation off). This is shown below comparing the Circulomics SRE buffer and 9 % PEG8000 (w/v) 1 M NaCl on precipitation and recovery using an equal volume addition to a P1000/26G needle sheared Bead-free LSK109 prep. Take this with a grain of salt at the moment as one used an old MinION flowcell (for Circulomics) and the others (library before selection (control), and PEG/NaCl size selection) were run on flongles so data volume is a little low with long frag blockage. We are seeing the SRE buffer performing better at short fragment removal, but want to investigate what is going on with the upper end and read N50.

Size selection of a finished P1000/26G needle sheared bead-free LSK109 ligation library



(final library split and diluted 1:1 with Circulomics SRE buffer or 9%PEG8000/1M NaCl/10mMTris-CI (pH8.0), incubated at room temp for 30 mins. followed by centrifugation at 13.5Krpm for 30 mins, supernatant removed and DNA resuspended in 20ul EB of EB).

Materials

- HMW genomic DNA
- Buffer EB
- Qubit DNA HS kit
- Ethanol
- Ultra II End-Prep Buffer (NEB)
- Ultra II End-Prep Enzyme mix (NEB)
- LNB Buffer
- AMX adapter
- Quick T4 DNA Ligase (NEB)
- NaCl
- PEG/NaCl precipitation buffer [9 % PEG8000 (w/v), 1 M NaCl, 10 M Tris-Cl (pH8.0)]



Safety warnings

• Please see SDS (Safety Data Sheet) for hazards and safety warnings.



- 1 $\underline{\underline{L}}$ 10 μg of **HMW genomic DNA** in $\underline{\underline{L}}$ 51 μL of **EB** was either used unsheared or sheared (see previous protocols).
- Quantify $\underline{\underline{A}} 1 \mu \underline{\underline{L}}$ of the DNA sample using the **Qubit DNA HS kit** to confirm DNA recovery and then set up the following End-Prep reaction:
 - Δ 50 μL DNA sample
 - 4 7 μL Ultra II End-Prep Buffer (NEB)
 - 🚨 3 µL Ultra II End-Prep Enzyme mix (NEB)
- Incubate at $3 \times 20 \, ^{\circ}\text{C}$ for $3 \times 20 \, ^{\circ}\text{C$
- Add Δ 60 μL of a **PEG/NaCl precipitation buffer** [9 % PEG8000 (w/v), [M] 1 Molarity (M) NaCl, [M] 10 Molarity (M) Tris-Cl (pH8.0)] and mixed by flicking tube.
- 5 Incubate at & Room temperature for 🕙 00:30:00 .
- 6 Centrifuge at (3) 13500 rpm (4) 13500 rpm for (5) 00:30:00 at (3)
- Remove supernatant and wash 2x with Δ 200 μL of **70** % **ethanol** by simple addition to the inside of the tube wall and centrifuging for 5 00:05:00 followed by removal.

Note

Be careful to avoid pelleted DNA when doing this (you may not see DNA so keep away from appropriate area of tube).

- 8 Allow DNA to air dry for a couple of mins.
- 9 Add \perp 41 μ L **EB buffer** (10 mM Tris-Cl pH 8.0).



10 Leave DNA in fridge to resuspend overnight.

Note

Depending on DNA fragment size time can be shortened.

- 11 Mix DNA by flicking and quantify $\Delta 1 \mu$ using the **Qubit DNA HS kit** to confirm DNA recovery and then set up the following Ligation reaction:
 - 🚨 33.5 µL DNA sample
 - 4 12.5 μL LNB buffer
 - Δ 2 μL AMX adapter
 - ∠ 2 μL Quick T4 DNA Ligase (NEB)
- 12 Incubate at \$\mathbb{L}\$ 20 °C / \$\mathbb{L}\$ Room temperature for \bigodeta 01:00:00 \ .
- 13 Add Δ 4.4 μL [M] 5 Molarity (M) NaCl.
- 14 Mix by flicking.
- 15 Incubate for 00:30:00 at Room temperature to precipitate the DNA.
- 16 Centrifuged at (3500 rpm) for (500:30:00) at (8 Room temperature).
- 17 Remove supernatant and wash 2x with $\Delta 200 \, \mu$ L of a 2 fold dilution of the previously used **PEG/NaCl precipitation buffer** above.

Do not disturb the pellet and centrifuge for 00:05:00 between each addition before removing wash solution.

- 18 Add $\stackrel{\blacksquare}{\Delta}$ 26 μ L **EB buffer** (10 mM Tris-Cl pH 8.0).
- 19 Leave adapted DNA in fridge to resuspend overnight.



Note

Depending on DNA fragment size time can be shortened.

- 20 Mix DNA by flicking and quantify \perp 1 μ L using the **Qubit DNA HS kit** to confirm DNA recovery.
- 21 [Possible round of size selection with Circulomics / PEG-NaCl buffer (see figure above).]
- 22 Library ready to be combined with SQB for loading onto flowcell (400 ng - \perp 1.3 µg depending on size and recovery).