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Integrated Indexing One-Step RT-PCR with NEB LunaScript: Influenza A Virus Sequencing with Native Oxford Nanopore Software Compatibility V.1

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ABSTRACT

This integrated indexing RT-PCR protocol has been optimized for whole genome sequencing of influenza A virus (IAV) using Oxford Nanopore's Ligation Sequencing Kit V14. The updated protocol is an optimization of an early approach used across several laboratories for research purposes, and uses New England Biolabs reagent sets including LunaScript ® Multiplex One-Step RT-PCR (#E1555), NEBNext dA-Tailing Module (E6053), and the NEBNext Quick Ligation Module (E6056). Previously demonstrated IAV whole genome sequencing approaches for Oxford Nanopore used Superscript III/Platinum Taq RT-PCR reagents input into the Native Indexing reagent set, which results in a complex, multi-step process that requires more than a full working day to carry out.

The improved integrated indexing protocol can be completed within a working day, and incorporation of the molecular index during first-strand cDNA synthesis in the RT-PCR step reduces the likelihood of sample cross-contamination during handling. Recent improvements to the approach additionally include Oxford Nanopore defined constant regions that are required for full compatibility with default informatics packages. This update now enables native demultiplexing within either Guppy or Dorado, effectively serving as a "molecular patch" where previous implementations required software modifications that were often broken by silent updates to MinKNOW.

As designed, this protocol has potential for extension to other viruses of interest by modifying primer sets. A simplified excel workbook is provided for design of new primer sets, for which the first worksheet requires input of gene-specific primer sets, the second worksheet provides fully constructed primers, and the third can be exported directly for submission to a primer synthesis vendor such as IDT DNA.



Protocol status: Working We use this protocol and it's

working

ATTACHMENTS

Process Overview.png

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Last Modified: Jan 19, 2024

PROTOCOL integer ID: 93455

MATERIALS

New England Biolabs Components:

А	В	С	D	E	F
NEB #	Component Name	Component #	Stored at (°C)	Amount	Concentration
E60 56	NEBNext Quick Ligation Reaction Buffer	E6058AVIAL	-20	1 x 0.2 ml	5 X
E60 56	Quick T4 DNA Ligase	E6057AVIAL	-20	1 x 0.1 ml	Not Applicable
E60 53	NEBNext dA-Tailing Reaction Buffer	E6055AVIAL	-20	1 x 0.1 ml	10 X
E60 53	Klenow Fragment (3´→ 5´ exo−)	E6054AVIAL	-20	1 x 0.06 ml	Not Applicable
E15 55	LunaScript‱ Multiplex One-Step RT-PCR Enzyme Mix	M1556SVIA L	-20	1 x 0.05 ml	25 X
E15 55	LunaScript‱ Multiplex One-Step RT-PCR Reaction Mix	M1557SVIA L	-20	1 x 0.25 ml	5X
E15 55	Nuclease-free Water	B1502AVIAL	-20	1 x 1.5 ml	Not Applicable

Oxford Nanopore Components:

A	В	С	D	E	F
ONT #	Component Name	Component #	Stored at (°C)	Amount	Concentratio n
SQK- LSK114	Ligation Sequencing Kit V14		-20	1 x 0.2 ml	6 Runs
	R10.4.1 Sequencing		4		



A	В	С	D	E	F
	Flowcell				

Other Equipment:

А
Component Name
Qubit
Tapestation
Ethanol (molecular biology grade)
Custom Indices

BEFORE START INSTRUCTIONS

Note: High quality isolated Influenza A RNA must be used for this protocol as it is essential for successful complete genome amplification. There are many options for RNA purification, including solid phase reversible immobilization methods (e.g. spin columns, paramagnetic beads), and Trizol derivatives. Freshly isolated RNA with minimal or no freeze thaw cycles typically ensures the highest success rates for complete genome amplification.

Protocol for Integrated Indexing of Influenza A Virus



Prepare the One-Step Reverse Transcriptase Multi-Segment PCR for Influenza A Virus as described below:

A	В
Reagent	Volume (uL)
Purified Viral RNA	5
5x Buffer (Reaction mix)	10
Nuclease Free Water (NFW)	31
Indexed Primer (MBTuni12-NB##)	1

	A	В
	Unindexed Primer (MBTuni13)	1
Г	25x Enzyme	2
Г	Total Volume	50

^{*}Resulting amplicons should range from 700 bp to 2.1kb

2 Flick the tube or pipet up and down to mix at minimum 10 times followed by a short spin



3 Incubate reaction in thermo-cycler following these steps:



A	В	С	D
Step	Temp (C)	Duration (min)	Cycles
RT Priming	55	2	1
Reverse Transcription	42	20	1
RT kill/ PCR Hot Start	98	2	1
PCR Round One	95	.5	5
	44	.5	
	68	3.5	
PCR Round Two	95	.5	26
	57	.5	
	68	3.5	
Final Extension	68	10	1
Hold	10	∞	∞

3. Library Pooling and Purification

4 Quantify unpurified PCR product using Qubit High Sensitivity DNA reagents

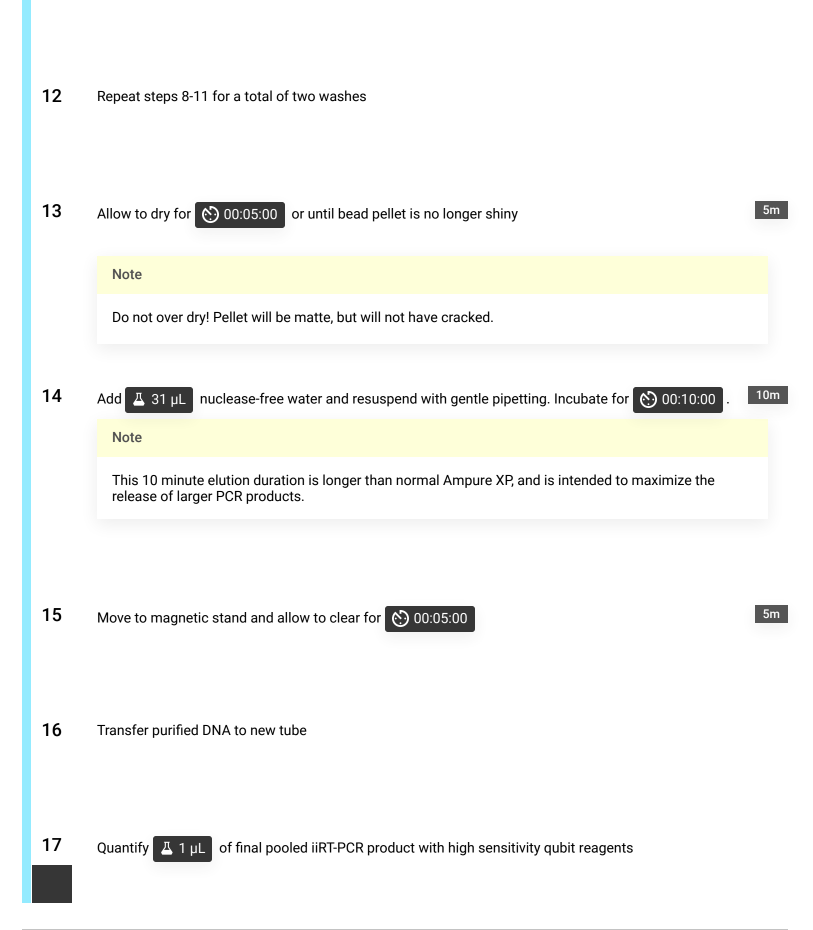
^{**} For no template control replace purified viral RNA with NFW

- 5 Pool indexed PCR products together by equal mass into a single tube for purification
- 6 Add .6 volumes of AMPure XP to the pooled iiRT-PCR product and mix by finger flicking

Note

Example: 60uL AMPure per 100uL pooled library

- 7 at room temperature or if available incubate on hula mixer at low speeds or IK. 5m Incubate for 00:05:00 plate shaker at 500 RPM
- 8 Move to magnetic stand to separate beads, allowing them to sit until solution clears
- 9 Remove and discard supernatant
- 10 Add 4 500 µL of freshly made 80% ethanol
- 11 Remove tube from magnet and finger flick





18 (Optional) Following purification of iiRT-PCR products, samples can be analyzed on an agarose gel, Tapestation, or similar platform.

DNA A-Tailing with NEBNext dA-Tailing Module

42m 30s

30s

- 19 Thaw all NEB consumables on ice and once thawed allow mixture to adjust to room temperature. Then pipette the buffer up and down to. Break up the precipitate and then a (5) 00:00:30
- 20 Place 100-200 fmol of barcoded, pooled and purified DNA into a 1.5 Eppendorf tube and adjust volume to Δ 30 μL with nuclease free water
- 21 Mix in a 0.2 ml PCR plate and then flick and spin down:

A	В
Reagents	Volume (uL)
DNA	30
NEBNext dA-Tailing. Reaction Buffer (10x)	5
Klenow Fragment (3'> 5' exo-)	3
Nuclease Free Water (NFW)	12
Total Volume	50

22 35m Incubate plate in thermal cycler at \$\mathbb{I}\$ 37 °C for \bigodamberrow 00:30:00 and then § 65 °C for 00:05:00

Oct 19 2024

23 Transfer DNA into 1.5 ml tubes

- 24 Add \coprod 60 μ L of AMPure beads to the tube and flick to mix
- 25 Incubate on hula mixer for © 00:05:00

5m

26 Spin down and place on magnet. Once solution is clear pipette off supernatant



- 28 Repeat step 27
- Take tube off of magnet and spin down, then place back on magnet and remove any residual ethanol
- Remove tubes from magnet and resuspend pellets in 🔼 61 µL of nuclease free water



31 Incubate for 00:02:00 at room temperature

2m

32 Place tubes on magnet and once solution is clear remove supernatant and place in clean 1.5 ml tubes

33 Quantify $\Delta 1 \mu L$ using qubit

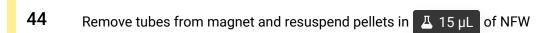
Adapter ligation and clean-up

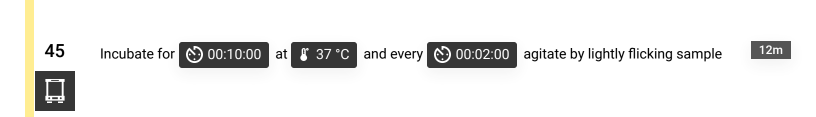
37m

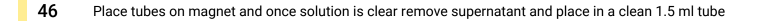
Thaw all NEB reagents at room temperature, spin down for 5 seconds, and mix by pipetting. DON'T VORTEX T4 DNA Ligase. Spin down the Ligation Adapter and Quick T4 DNA Ligase, pipette to mix then place on ice. Thaw the Elution Buffer and Short Fragment Buffer at room temp, mix by vortexing, spin down, and then place on ice.

35 Mix by pipetting and spin down the following reaction:

A	В
Reagents	Volume (uL)
Pooled Barcoded Sample	30
Ligation Adapter (LA)	5
NEBNext Quick Ligation Reaction Buffer (5X)	10
Quick T4 DNA Ligase	5
Total	50







Note

This is the final clean library, and should be stored on ice or at 4C until sequencing

Final Library Quantification

- 47 Quantify Δ 1 μL of the final clean library using a Qubit DNA fluorometer
- 48 Optional: Run final library on a Tapestation to evaluate average peak size
- 49 Calculate the final volume required for 200fmol to be sequenced on a MinION flowcell

Note

Tip: Use the NEBioCalculator https://nebiocalculator.neb.com/#!/dsdnaamt. Assuming 1kb product mean, a final loading concentration of 200fmol requires 125ng of library into the flowcell.



50

Proceed to load the Oxford Nanopore flowcell using the most up to date protocols from the vendor.