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(NEBNext® Immune Sequencing Kit (Mouse) (NEB #E6330S/L)

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ABSTRACT

The NEBNext Immune Sequencing Kit (Mouse) contains enzymes and buffers that are ideal to convert a small amount of RNA input into indexed libraries for next-generation sequencing on the Illumina MiSeq platform (Illumina, Inc). The workflow of NEBNext Immune Sequencing Kit (Mouse) is very user-friendly and fast with minimal hands-on time. Each of these components must pass rigorous quality control standards and are lot controlled, both individually and as a set

For larger volume requirements, customized and bulk packaging is available by purchasing through the OEM?Bulks department at NEB. Please contact OEM@neb.com for further information.

GUIDELINES

Overview:

Immune repertoire sequencing is a powerful tool that can be used to analyze the immune response to diseases and pathogens from current and past exposures. Areas of particular interest include characterization of autoimmune diseases, oncology, discovery of neutralizing antibodies against infectious disease, or use as a diagnostic tool for residual disease detection.

In order to create the incredible genetic diversity required for antigen recognition by B and T Cells, lymphocytes construct unique antigen receptors via a process known as V(D)J recombination. In this process, each cell selects a single V, D, and J gene segment through genetic recombination, introducing additional non-germline-encoded nucleotides at the junctions. This process generates immune receptor diversity, the majority of which is encoded in the heavy chain complementarity determining region 3 (CDR3) (Figure 1).

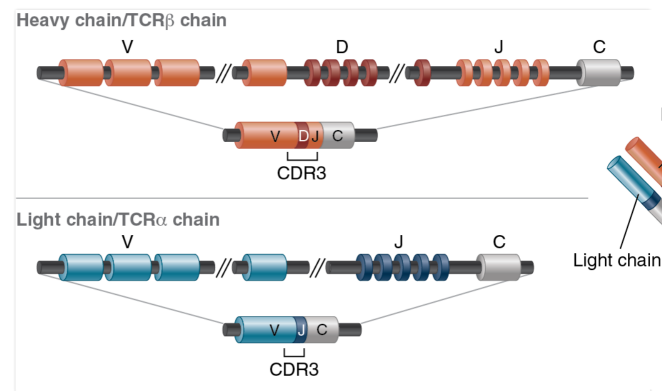
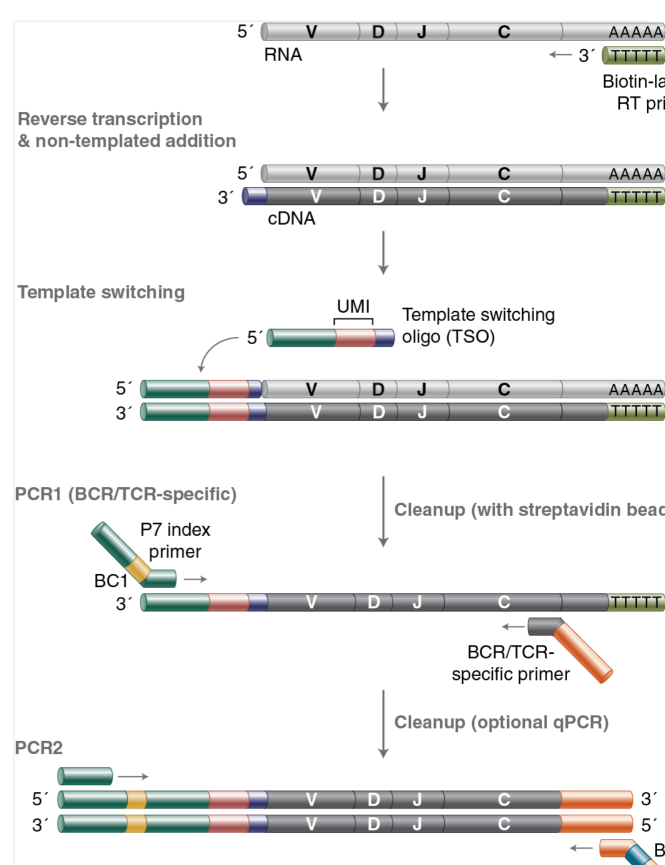


Figure 1. Simplified representation of the structure of BCR or TCR showing the outcome of V(D)J recombination in mature lymphocytes.

Recent improvements in the read length and throughput of next-generation sequencing (NGS) platforms have resulted in a rise in the popularity of immune repertoire sequencing. The complex structure and sequence diversity of antibody-encoding genes have provided significant challenges to the development of a simple and reliable method to sequence immune receptor sequences.

The NEBNext Immune Sequencing Library Preparation Kit (Figure 2) provides all the components required for enrichment and sequencing of the B Cell

Receptor (BCR) and T Cell Receptor (TCR) RNA transcripts present in a sample. NEBNext Immune Sequencing Library Preparation Kit has been developed and optimized to provide accurate sequencing of full-length immune transcript repertoires of B Cells and T Cells. This allows for exhaustive somatic mutation profiling across complete V, D and J segments, full isotype information analysis (IgM, IgD, IgG, IgA and IgE), and also allows for synthesis and expression of complete antibody chains for downstream immunological assays. In addition, this approach allows characterization of TRA, TRB, TRD and TRG chains. The method uses a unique molecular identifier scheme specifically designed to barcode each mRNA molecule, allowing PCR copies derived from an individual mRNA to be collapsed into a single consensus sequence. This improves sequence accuracy by resolving PCR and sequencing errors. This also eliminates PCR bias, thus allowing for quantitative digital molecule counting.



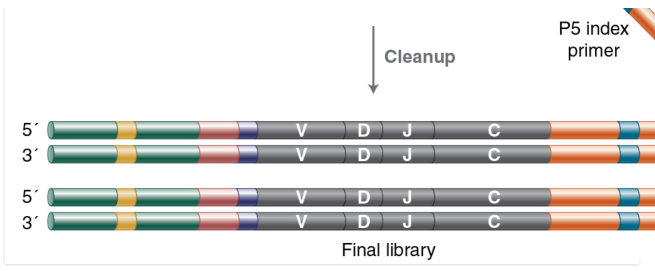


Figure 2. NEBNext Immune Sequencing Library Preparation Kit Workflow.

Defining features of the Immune Sequencing method include:

- 1. Full-length variable sequences are generated (including isotype information), allowing downstream antibody synthesis and functional characterization not possible with approaches sequencing only the CDR3 region.
- 2. Variable region primers are not required, which reduces primer pool complexity and allows for the unbiased and simultaneous recovery of B Cell and T Cell receptor transcripts.
- 3. Unique molecular barcoding approach minimizes PCR biases and improves sequencing accuracy by allowing a consensus to be generated from duplicate sequencing reads originating from the same transcript. Moreover, unique barcoding enables accurate quantitation of each clone present in the sample.
- 4. High target capture efficiency allows for immune repertoire sequencing and analysis from sub-microgram quantities of total RNA.

Step I-Prepare target RNA

Immune Sequencing does not require mRNA isolation prior to library construction. Total RNA can be prepared from any sample or tissue known to contain B Cells or T Cells, such as peripheral blood mononuclear cells, bone marrow, lymph nodes, or any other tissue in which infiltrating lymphocytes are present. Cells or tissues can be frozen, but intact, high quality RNA is required for optimal performance.

Step II-Perform molecular-tagging reverse transcription

During this 50-minute reaction full-length mRNA is reverse transcribed and the 3' end of each cDNA is tagged with a DNA adaptor containing a unique molecular identifier (UMI) barcode.

Step III-Purify with Streptavidin Magnetic Beads

This 30-minute cleanup step removes all salts, reagents and unused cDNA adaptor oligonucleotides (TSO and RT Primers) allowing clean downstream amplification steps.

Step IV-Primary amplification of target fragments

This primary PCR step uses one primer which binds the universal cDNA adaptor and a primer mix targeting constant regions of the desired immune receptors. Products are then purified with Sample Purification Beads.

Step V-Secondary PCR cycle optimization with real-time PCR

This optional real-time PCR allows accurate assessment of target quantity leading to an optimized choice of PCR cycles for the final library PCR.

Step VI-Secondary PCR including Illumina® adaptor addition

This secondary PCR adds additional Illumina adaptor sequences to the target fragments (including Illumina multiplexing index sequences if desired) After Sample Purification Bead cleanup the libraries are ready for Bioanalyzer or TapeStation analysis, quantification and sequencing on the MiSeq platform.

Applications

The NEBNext Immune Sequencing Kit (Mouse) contains enzymes and buffers that are ideal to convert a small amount of RNA input into indexed libraries for next-generation sequencing on the Illumina MiSeq platform (Illumina, Inc). The workflow of NEBNext Immune Sequencing Kit (Mouse) is very user-friendly and fast with minimal hands-on time. Each of these components must pass rigorous quality control standards and are lot controlled, both individually and as a set of reagents.

For larger volume requirements, customized and bulk packaging is available by purchasing through the OEM/Bulks department at NEB. Please contact OEM@neb.com for further information.

The Kit Includes

NEB #E6330S contains E2629S (Pouch with BCR and TCR primers), #E6325-1 (Box 1 of 2), and #E6325-2 (Box 2 of 2) and is sufficient for preparation of up to 24 reactions. NEB #E6330L contains E2629L (Pouch with BCR and TCR primers), #E6325-3 (Box 1 of 2), and #E6325-4 (Box 2 of 2) and is sufficient for preparation of up to 96 reactions. All reagents should be stored at -20°C, 4°C or room temperature as indicated below. Colored bullets represent the color of the cap of the tube containing the reagent.

NEB #E2629S/L, NEBNext Immune Sequencing Primers (Mouse)

Pouch: Store at -20°C.

(blue) NEBNext IS BCR Primers (Mouse)

(blue) NEBNext IS TCR Primers (Mouse)

NFB #F6325-1.

NEBNext Immune Sequencing Kit RT and PCR Reagents (24 reactions) New England Biolabs Catalog #E6320S_____

Box 1 of 2: Store at −20°C.

(lilac) NEBNext IS RT Buffer (4X)

(lilac) NEBNext IS RT Primer

(lilac) NEBNext IS TS Oligos

(lilac) NEBNext First Strand Synthesis Enzyme Mix

(lilac) dNTP Solution Mix

(lilac) NEBNext Cell Lysis Buffer

(blue) Q5 ® Reaction Buffer (5X)

(blue) Q5 Hot Start High-Fidelity DNA Polymerase

(blue) NEBNext i701-i706 Primers

(blue) NEBNext i501-i504 Primers

(blue) NEBNext IS PCR2 Universal Primer

(white) Nuclease-free Water

(white) TE (0.1X)

NEB #E6325-3, NEBNext Immune Sequencing Kit RT and PCR Reagents (NEB #E6330L, 96 reactions)

Box 1 of 2: Store at -20°C.

(lilac) NEBNext IS RT Buffer (4X)

(lilac) NEBNext IS RT Primer

(lilac) NEBNext IS TS Oligos

(lilac) (lilac) NEBNext First Strand Synthesis Enzyme Mix

(lilac) dNTP Solution Mix

(lilac) NEBNext Cell Lysis Buffer

(blue) (blue) Q5 Reaction Buffer (5X)

(blue) (blue) Q5 Hot Start High-Fidelity DNA Polymerase

(blue) NEBNext i701-i712 Primers

(blue) NEBNext i501-i508 Primers

(blue) NEBNext IS PCR2 Universal Primer

(white) Nuclease-free Water

(white) TE (0.1X)

NEB #E6325-2/#E6325-4, NEBNext Immune Sequencing Kit cDNA Purification Reagents

Box 2 of 2: Store at 4°C. Do not freeze.

(white) Streptavidin Magnetic Beads

(white) NEBNext IS Bind and Wash Buffer (2X)

(white) 0.1% Tween

(white) NEBNext Sample Purification Beads*

*This component should be stored at room temperature.

Required Materials Not Included

- SYBR® Green (Life Technologies #S7563)
- 80% Ethanol (freshly prepared)
- DNA LoBind ® Tubes (Eppendorf #022431021)
- OI
- Magnetic rack/stand (NEB #S1515, Alpaqua®, cat. #A001322 or equivalent)
- Thermal cycler
- qPCR machine and associated 96 well plates
- Bioanalyzer®/TapeStation® and associated consumables
- MiSeq® and v3 600 cycle sequencing reagent
- Nuclease-free Water
- Tube rotating mixer
- Magnet stand or equivalent (for washing beads only) (NEB #S1506)
- DNase RNase-free PCR strip tubes (USA Scientific 1402-1708)

NEBNext Immune Sequencing Reverse Transcription and cDNA Synthesis

X

Note

RNA Sample Recommendations: The RNA sample should be free of salts (e.g., Mg²⁺, or guanidine salts) or organics (e.g., phenol and ethanol). High quality RNA with a RIN > 7 is recommended for optimal performance.

Starting Material:

enriched B cell or T Cell or B cell and T Cell total RNA, or up total RNA from peripheral blood mononuclear cells (PBMCs). The maximum volume of RNA input is up 9 µL.

Mix the following components in a sterile nuclease-free tube:

A	В
Component	Volume
(lilac) NEBNext IS RT Buffer (4X)	5 μΙ
(lilac) NEBNext IS RT Primer	1 μΙ
(lilac) NEBNext IS TS Oligos	1 μΙ
(lilac) dNTP Solution Mix	2 μΙ
(lilac) NEBNext First Strand Synthesis Enzyme Mix	2 μΙ
(lilac) NEBNext Cell Lysis Buffer	0.5 μΙ
Total RNA	1-9 μΙ
white) Nuclease-free Water	Variable
Total Volume	20.5 μΙ

Set a 100 µl or 20 µl pipette to 15 µl and then pipette the entire volume up and down at least 10 times to mix thoroughly. Perform a quick spin to collect all liquid from the sides of the tube.



Note

Note: It is important to mix well. The presence of a small amount of bubbles will not interfere with the performance.

Place in a thermal cycler, with the heated lid set to \geq 80 °C, and run the following program:

A	В	С
CYCLE STEP	TEMP	TIME
Incubation	42°C	40 minutes
Inactivation	70°C	10 minutes
Hold	4°C	∞

Purify the cDNA with Streptavidin Magnetic Beads

17m

- 4 Prepare Bind and Wash Buffer (1X) to be used in Step 14.
 - a. Dilute NEBNext IS Bind and Wash Buffer (2X) to 1X with an equal volume of nuclease-free water (not provided).
 - b. For each sample, you will need 🔼 100 µL Bind and Wash Bufer (1X).
 - Vortex Streptavidin Magnetic Beads to resuspend. Aliquot the total amount of Streptavidin Magnetic Beads needed (4 15 µL of Streptavidin Magnetic Beads per sample) into a clean RNase-free 1.5 ml tube.

5

- Place the tube on a magnetic rack at Room temperature. Once the solution is clear (~ 00:02:00) carefully remove and discard the supernatant without disturbing the bead pellet.
- 2m

- 7
 - Remove the tube from the magnet and wash the beads by adding A 200 µL of undiluted NEBNext IS Bind and Wash Buffer (2X). Vortex briefly to resuspend the beads and quickly spin the tube in a microcentrifuge to collect any sample on the sides of the tube.
 - 8 Place the tube on a magnetic rack. Once the solution is clear, carefully remove and discard the supernatant without disturbing the bead pellet.
 - 9 Repeat Steps 7 and 8 once for a total of two washes.
- Remove the tube from the magnetic rack. Resuspend the beads in 2 20 µL of undiluted NEBNext IS Bind and Wash Buffer (2X) for each sample (e.g., for 4 samples, add 4 x 20 = 2 80 µL NEBNext IS Bind and Wash Buffer (2X). Vortex briefly to resuspend the beads.
- 11 Add 🛕 20 µL beads to the cDNA products (20.5 µl) from Step 3. Vortex briefly to mix.
- 8° ;;
- Place the tube on a rotator for 00:15:00 at 8 Room temperature
- Quickly spin down the tube to collect any sample on the sides of the tube and place the tube on a magnetic rack. Once the solution is clear, carefully remove and discard the supernatant without disturbing the bead pellet.
- Remove the tube from the magnetic rack. Add Δ 100 μL diluted Bind and Wash Buffer (1X) (from Step 4). Vortex briefly to resuspend the beads.

Note: Vortexing is better than flicking the tube, as it minimizes foaming.

- Quickly spin down the tube to collect any sample on the sides and place the tube on a magnetic rack. Once the solution is clear, carefully remove and discard the supernatant without disturbing the bead pellet.
- Remove the tube from the magnetic rack and add 🔼 100 µL of 0.1% Tween. Vortex briefly to resuspend the beads.

15m



Note : This step is critical to remove all the salt from the beads before elution.

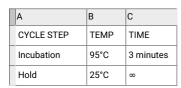
17 Quickly spin down the tube to collect any sample on the sides of the tube and place the tubes on a magnetic rack. Once the solution is clear, carefully remove and discard the supernatant without disturbing the bead pellet. With a 20-µl pipette tip, carefully remove any residual liquid left at the bottom of the tube without disturbing the beads.

Remove the tube from the magnet and add 23 µL of 0.1% Tween. Vortex briefly to resuspend the beads and quickly spin in a microcentrifuge to collect any sample on the sides of the tube.



19

Place the tube in a thermal cycler, with the heated lid set at [105 °C], and run the following program:



20 Remove the tube from the thermal cycler, vortex briefly, spin down the contents and place on the magnetic rack.

X

Once the solution is clear, transfer 🔼 21 µL of the supernatant into a new tube and discard the beads.



21

Note

Safe Stopping Point: It is safe to store the cDNA at T -20 °C Overnight

PCR1 for VDJ Region Amplification and Index Barcode Incorporation

17m

22

Note

Caution:

- Use Option step case 1 for enriching B Cell Receptor (BCR) chains.
- Use Option step case 2 for enriching T Cell Receptor (TCR) chains.
- Use Option step case 3 for enriching both BCR chains and TCR chains.

Step 22 includes a Step case.

B Cell Receptor Chains PCR 1

T Cell Receptor Chains PCR 1

B and T Cell Receptor Chains PCR 1

step case

B Cell Receptor Chains PCR 1

23 Add the following components to a sterile strip tube:



ſ	Δ	В
ľ	COMPONENT	VOLUME
	COMI CIVEIVI	VOLOIVIL

A	В
Purified cDNA (Step 21)	21 μΙ
(blue) Q5 Reaction Buffer (5X)	10 μΙ
(lilac) dNTP Solution Mix	1 μΙ
(blue) Q5 Hot Start High-Fidelity DNA Polymerase	1 μΙ
(blue) NEBNext IS BCR Primers (Mouse)	2 μΙ
(blue) NEBNext i7 Index Primer*	1 μΙ
(white) Nuclease-free Water	14 μΙ
Total Volume	50 μΙ

^{*}Use only one i7 Primer per sample.



24

Set a 100 μ l or 200 μ l pipette to 40 μ l and then pipette the entire volume up and down at least 10 times to mix thoroughly. Perform a quick spin to collect all liquid from the sides of the tube.

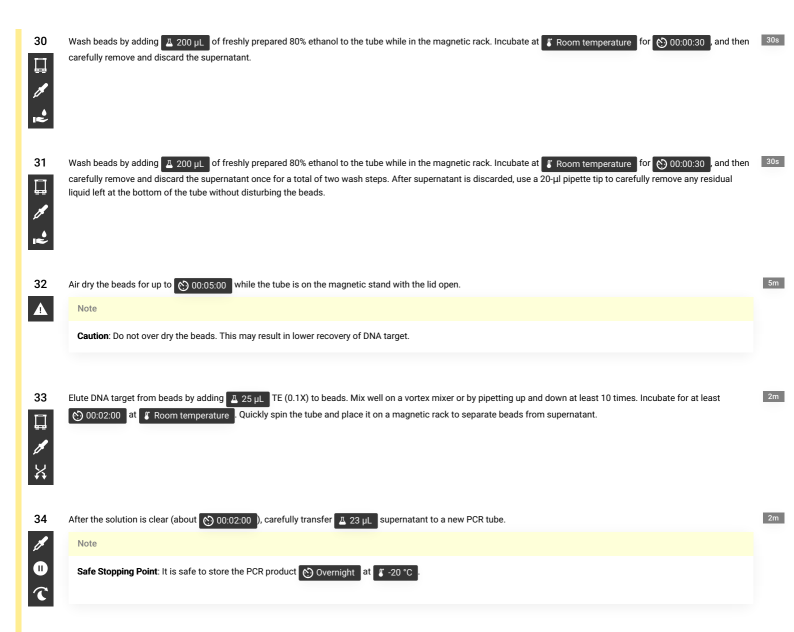
25

Place the tube on a thermal cycler and perform PCR amplification using the following PCR cycling program:



A	В	С	D
CYCLE STEP	TEMP	TIME	CYCLES
Initial Denaturation	98°C	60 seconds	1
Denaturation	98°C	10 seconds	
Annealing	64°C	30 seconds	12
Extension	72°C	30 seconds	
Final Extension	72°C	60 seconds	1
Hold	4°C	∞	

Cleanup of PCR1 Amplification 26 Vortex NEBNext Sample Purification Beads to resuspend. SPRIselect and AMPure® XP Beads can be used as well. If using AMPure XP Beads, please allow the beads to warm to resuspended NEBNext Sample Purification Beads to each 50 µL PCR reactions. Mix well by pipetting or brief vortexing. 27 Add 50 µL of resuspended NEBNext Sample Purification Beads to each 50 µL PCR reactions. Mix well by pipetting or brief vortexing. 28 Incubate for 00:005:00 at Room temperature . 29 Quickly spin the tube in a microcentrifuge and place it on a magnetic rack. Once the solution is clear (00:005:00) carefully remove and discard the supernatant without disturbing the bead pellet. Note Caution: Do not discard beads.



qPCR for PCR2 Cycle Optimization

35



Note

The qPCR step is optional; it is only used to identify the optimal final PCR cycling conditions, as each sample can contain varying amounts of amplifiable molecules. One can achieve a similar result by manually taking an aliquot of the PCR reaction every 5 cycles and visualizing each aliquot on Bioanalyzer or TapeStation to identify the optimal cycling number that yields the highest product with the lowest background. The number of cycles should be high enough to provide sufficient library fragments for a successful sequencing run, but low enough to avoid PCR artifacts and over-cycling (high molecular weight fragments on Bioanalyzer). Typically, most samples will amplify best in the range of 6–18 cycles.

Add the following components to a sterile strip tube:

A	В
COMPONENT	VOLUM E
Purified PCR1 DNA (Step 34)	5 μΙ
(blue) Q5 Reaction Buffer (5X)	5 μΙ
(lilac) dNTP Solution Mix	0.5 μΙ
(blue) Q5 Hot Start High-Fidelity DNA Polymerase	0.5 μΙ
(blue) NEBNext IS PCR2 Universal Primer	1 μΙ
(blue) NEBNext i5 Index Primer*	1 μΙ

A	В
SYBR Green (20X)**	0.25 μΙ
(white) Nuclease-free Water	11.75 µl
Total Volume	25 μΙ

^{*} Use i5 Primer per PCR reaction.

Set a 100 μ l or 20 μ l pipette to 20 μ l and then pipette the entire volume up and down at least 10 times to mix thoroughly. Perform a quick spin to collect all liquid from the sides of the tube.



Place the tube on a qPCR instrument and run the following PCR cycling program:



A	В	С	D
CYCLE STEP	TEMP	TIME	CYCLES
Initial Denaturation	98°C	60 seconds	1
Denaturation	98°C	10 seconds	
Annealing and Extension	72°C	30 seconds (fluorescence on*)	30
Hold	4°C	∞	

^{*} Select the SYBR channel.

The amplification curve for each sample is used to determine the optimal number of PCR cycles to use during PCR2 (Section: **PCR2 for Final Library Amplification**). The PCR cycle number is approximated by determining the cycle number (round down to the nearest whole number) corresponding to 2/3 of the peak fluorescence. (see example in Figure 3; in this example 13 PCR cycles will be used in Section: PCR2 for Final Library).

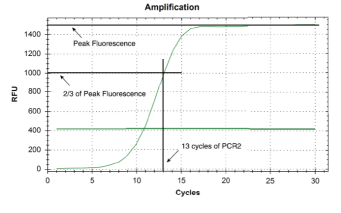


Figure 3. Determining cycle number for PCR2 using amplification curve of sample from a qPCR run.

^{** 20}X working stock of SYBR Green can be made by diluting \square 1 μ L (10,000X) SYBR Green (Life Technologies #S7563) into \square 399 μ L DMSO and stored at \square 4 °C for up to a month. You only need 0.2 x SYBR final in the qPCR reaction.

See example in Figure 1; in this example 13 PCR cycles will be used in Section: PCR2 for Final Library Amplification.

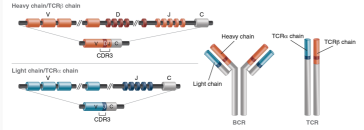


Figure 1. Simplified representation of the structure of BCR or TCR showing the outcome of V(D)J recombination in mature lymphocytes.

PCR2 for Final Library Amplification

Add the following components to a sterile strip tube:



A	В
COMPONENT	VOLUME
Purified PCR1 DNA (Step 34)	10 μΙ
(blue) Q5 Reaction Buffer (5X)	10 μΙ
(lilac) dNTP Solution Mix	1 μΙ
(blue) Q5 Hot Start High-Fidelity DNA Polymerase	1 μΙ
(blue) NEBNext IS PCR2 Universal Primer	2 μΙ
(blue) NEBNext i5 Primer*	2 μΙ
(white) Nuclease-free Water	24 μΙ
Total Volume	50 μΙ

^{*} Use i5 Primer per PCR reaction.

40



Set a 100 μ l pipette to 40 μ l then pipette the entire volume up and down at least 10 times to mix thoroughly. Perform a quick spin to collect all liquid from the sides of the tube.

41 Place the tube on a thermal cycler and run the following PCR cycling program:



A	В	С	D
CYCLE STEP	TEMP	TIME	CYCLES
Initial Denaturation	98°C	60 seconds	1
Denaturation	98°C	10 seconds	X*
Annealing and Extension	72°C	30 seconds	^"
Hold	4°C	∞	

^{*} Use number of cycles determined from Step 38. If qPCR was not done, see Table 1 for recommended number of PCR cycles. In this case the number of PCR cycles should be chosen based on input amount and sample type. The number of cycles should be high enough to provide sufficient library fragments for a successful sequencing run, but low enough to avoid PCR artifacts and over-cycling (high molecular weight fragments on Bioanalyzer).

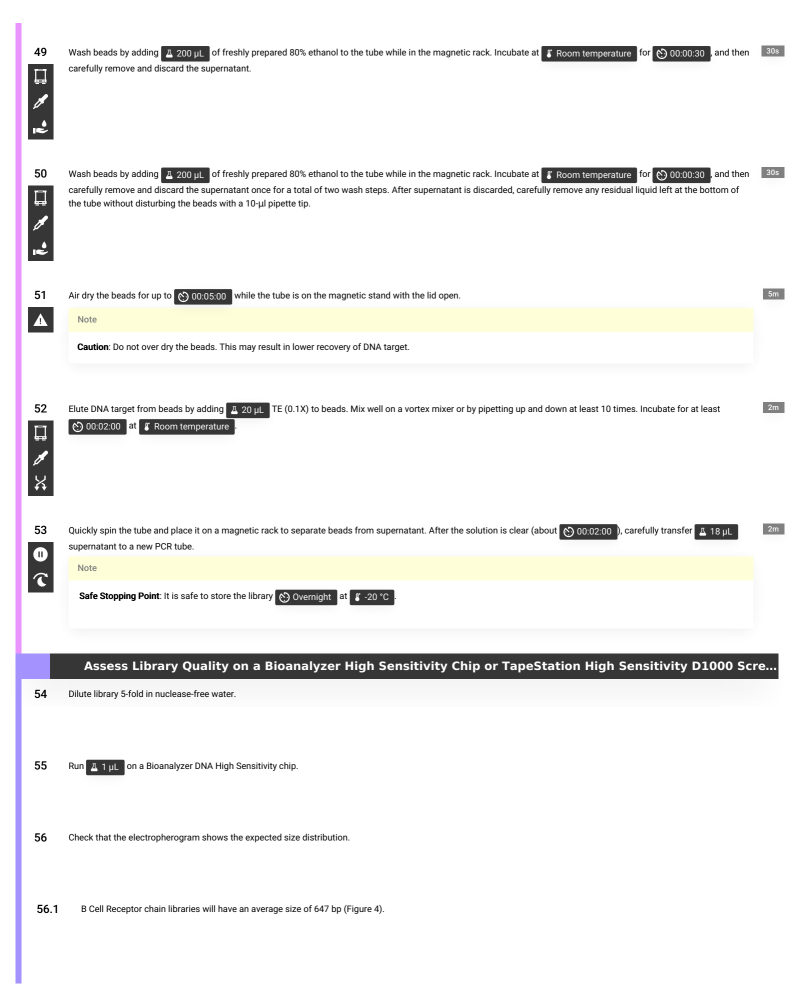
Table 1. For Mouse Spleen RNA input, PCR2 cycle number is recommended as in the table:

A	В	С
TOTAL RNA INPUT	PCR1 ASSAY	PCR2 CYCLE NUMBER
Mouse Spleen RNA 1,000 ng	BCR	6-7
Mouse Spleen RNA 100 ng	BCR	9-10
Mouse Spleen RNA 10 ng	BCR	15-16
Mouse Spleen RNA 1,000 ng	TCR	12-13
Mouse Spleen RNA 100 ng	TCR	14-15
Mouse Spleen RNA 10 ng	TCR	20-21

Note: if B and T cells are combined, choose a PCR2 cycle number in between the recommendations for B cell and T cell RNA.

Cleanup of PCR2 Amplification to Obtain Final Libraries 42 Vortex NEBNext Sample Purification Beads to resuspend. SPRIselect and AMPure XP Beads can be used as well. If using AMPure XP Beads, please allow the beads to warm to 8 Room temperature for at least 👏 00:30:00 before use. 43 Add 🗸 50 µL of TE (0.1X) to each 🗓 50 µL PCR reaction to obtain 👢 100 µL mixture, and then add 🗓 50 µL resuspended NEBNext Sample Purification Beads to the 🔼 100 µL mixture. Mix well by pipetting or brief vortexing. 44 5m Incubate for 👏 00:05:00 at 🛭 Room temperature 45 Quickly spin the tube in a microcentrifuge and place it on a magnetic rack. Once the solution is clear (👏 00:05:00) carefully transfer the 🔼 150 μ L supernatant to a 5m new tube without disturbing the bead pellet. Save the supernatant and discard beads. Add 🔼 15 µL of resuspended NEBNext Sample Purification Beads to supernatant. Mix well by pipetting or brief vortexing. 47 Incubate for 👏 00:05:00 at 🖇 Room temperature 48 5m Quickly spin the tube in a microcentrifuge and place it on a magnetic rack. Once the solution is clear (👏 00:05:00) carefully remove and discard the supernatant without disturbing the bead pellet. Note

Caution: Do not discard beads



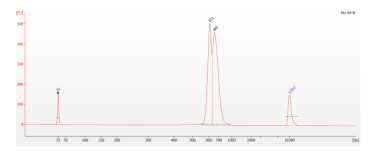


Figure 4. Example of B Cell Receptor library size distribution on a Bioanalyzer.

T cell receptor chain libraries will have an average size of 630 bp (Figure 5).

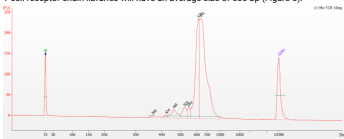


Figure 5. Example of T Cell Receptor library size distribution on a Bioanalyzer.

56.3 BCR + TCR chain libraries will have an average size of 632 bp (Figure 6).

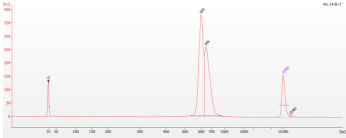


Figure 6. Example of B Cell Receptor chains and T Cell Receptor chains library size distribution on a Bioanalyzer.

Sequencing and Data Analysis

57 Sequencing should be performed on an Illumina MiSeq, using Illumina V3 sequencing chemistry. We recommend running 300 PE using a 600-cycle V3 MiSeq reagent kit with the index read between the first and second read.

Note

For information on data processing please refer to Data Usage Guideline Page within the "Usage Guidelines & Tips" panel in the "Other Tools & Resources" tab at: www.neb.com/E6330

We have developed and optimized a pipeline for processing of data directly from FASTQ files using open-source bioinformatics tools. Details on this pipeline can be found at: https://usegalaxy.org/u/bradlanghorst/w/presto-nebnext-immune-seq-workflow-v320.

58 Index Pooling Guidelines

For more detailed indexing information please refer to the manual for NEBNext Multiplex Oligos for Illumina (Dual Index Primers Set 1), NEB #E7600.

For multiplexing samples, use Table 1 for some valid index combinations.

Table 1. Examples of valid index combinations.

A	В	С
PLEX	i7 PRIMERS	i5 PRIMERS
2	i701 and i702 i703 and i704 i705 and i706 i707 and i708 i709 and i710 i711 and i712	Any i5 primer
3	i701, i702 and i703 i703, i704 and i705 i705, i706 and i707 i707, i708 and i709 i709, i710 and i711	Any i5 primer
4	i701, i702, i703 and i704 i703, i704, i705 and i706 i705, i706, i707 and i708 i707, i708, i709, and i710 i709, i710, i711, and i712	Any i5 primer
5-12	Any valid i7 4-plex with any other i7 primer.	Any i5 primer
7-12	Any 3 plex combination with any other i7 primer (as needed)	i501, i502 i503, i504 i505, i506 i507, i508
Greater than 12	Any 4 plex combination with any other i7 primer (as needed)	i501, i502 and any other i5 primer (as needed) i503, i504 and any other i5 primer (as needed) i505, i506 and any other i5 primer (as needed) i507, i508 and any other i5 primer (as needed)

Table 2. Index sequences of NEBNext i501 Primer – NEBNext i508 Primer:

A	В
PRODUCT	EXPECTED INDEX READ
	MiSeq
NEBNext i501 Primer	TATAGCCT
NEBNext i502 Primer	ATAGAGGC
NEBNext i503 Primer	CCTATCCT
NEBNext i504 Primer	GGCTCTGA
NEBNext i505 Primer	AGGCGAAG
NEBNext i506 Primer	TAATCTTA
NEBNext i507 Primer	CAGGACGT
NEBNext i508 Primer	GTACTGAC

Table 3. Index sequences of NEBNext i701 Primer - NEBNext i712 Primer

A	В
PRODUCT	EXPECTED INDEX READ
NEBNext i701 Primer	ATTACTCG
NEBNext i702 Primer	TCCGGAGA
NEBNext i703 Primer	CGCTCATT
NEBNext i704 Primer	GAGATTCC
NEBNext i705 Primer	ATTCAGAA
NEBNext i706 Primer	GAATTCGT
NEBNext i707 Primer	CTGAAGCT
NEBNext i708 Primer	TAATGCGC
NEBNext i709 Primer	CGGCTATG
NEBNext i710 Primer	TCCGCGAA
NEBNext i711 Primer	TCTCGCGC
NEBNext i712 Primer	AGCGATAG

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