

Library Preparation Protocol

TCR/BCR Full Length, mRNA Whole Transcriptome Analysis (WTA), BD® AbSeq, and Sample Tag

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### Regulatory information

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#### History

Revision	Date	Change made	
23-24020(01)	2022-01	Initial release.	
23-24020(02)	2022-07	Updated Time considerations workflow.	
		Updated Sequencing section. Added additional library pooling and sequencing recommendation.	
		In Appendix, added Illumina Index 1 (i7) sequences.	

# **Contents**

Introduction	5
Required and recommended materials	5
Required reagents	5
Recommended consumables	5
Equipment	6
Best practices	6
Additional documentation	6
Safety information	6
Time considerations	7
Procedure	8
Preparing BD Rhapsody™ Enhanced Cell Capture Beads for TCR/BCR full length, WTA, BD® AbSo	eq,
and Sample Tag library amplification	8
cDNA synthesis and template switching	8
Denaturation and self-hybridization	9
TCR/BCR extension	10
Treating the sample with Exonuclease I	11
Performing random priming and extension (RPE) on BD Rhapsody™ Enhanced Cell Capture Bead	ls
with cDNA	11
Purifying RPE product	13
Performing RPE PCR	
Purifying RPE PCR amplification product (single-sided cleanup)	14
Performing AbSeq/Sample Tag PCR1	16
Performing TCR/BCR PCR1	17
Purifying TCR/BCR and AbSeq/Sample Tag PCR1 products	19
Quantifying BD® AbSeq/Sample Tag PCR1 products	19
Performing TCR/BCR and Sample Tag PCR2 on the PCR1 products	20
Purifying TCR/BCR and Sample Tag PCR2 products	21
Performing random priming and extension (RPE) on TCR/BCR PCR2 products	22
Purifying TCR/BCR RPE product	23
Performing TCR/BCR and AbSeq/Sample Tag index PCR	24
Purifying TCR/BCR, AbSeq, and Sample Tag index PCR products	26
Performing WTA index PCR	28
Purifying WTA index PCR product (single-sided cleanup)	30

Sequencing	32
Read requirements for libraries	32
Pooling libraries for sequencing	32
Example of pooling with no correction	33
Example of pooling with correction	33
Example of pooling with a mixed population	34
Example of pooling with correction while sequencing AbSeq and Sample Tag libraries separately	35
Additional considerations	35
Sequencing flow cell loading and PhiX concentrations	36
Quantifying libraries	36
AbSeq/Sample Tag libraries	36
WTA and TCR/BCR libraries (with or without AbSeq/Sample Tag)	36
Sequencing analysis pipeline	37
Appendix	38
Illumina Index 1 (i7) sequences	38
Human T cell PCR1 primers	38
Human T cell PCR2 primers	38
Human B cell PCR1 primers	
Human B cell PCR2 primers	39

# Introduction

This protocol enables high-throughput single-cell transcriptome and protein analysis alongside TCR and BCR profiling of individual cells captured on the BD Rhapsody™ system, providing instructions for amplifying Illumina-compatible single-cell barcoded mRNA, TCR, and BCR libraries. The addition of a sample-tag library enables demultiplexing of samples that were pooled before capture on the BD Rhapsody™ system.

After staining of cell with AbSeg and Sample Tag antibodies and partitioning and lysis of cells, cDNA is encoded on BD Rhapsody™ Enhanced Cell Capture beads using both the 3' and 5' ends of transcripts as templates. mRNA, TCR, and BCR libraries are then amplified from these on-bead cDNA libraries using a two-step nested amplification, with TCR and BCR libraries undergoing additional random priming to capture complementarity determining regions (CDR) 1, 2, and 3, as well as framework regions (FR) 1-4. AbSeq and Sample Tag libraries are amplified from the small products recovered after cleanup of larger mRNA, TCR, and BCR PCR1 products. Note that Sample Tag and AbSeq libraries are amplified from the supernatant that was denatured from the beads.

# Required and recommended materials

# Required reagents

Store the reagents at the storage temperature specified on the label.

Material	Supplier	Catalog no.
BD Rhapsody™ WTA Amplification Kit	BD Biosciences	633801
BD Rhapsody™ TCR/BCR Amplification Kit	BD Biosciences	665345
Agencourt <sup>®</sup> AMPure <sup>®</sup> XP magnetic beads	Beckman Coulter	A63880
100% ethyl alcohol	Major supplier	_
Nuclease-free water	Major supplier	_

# Recommended consumables

Material	Supplier	Catalog no.
Pipettes (P10, P20, P200, P1000)	Major supplier	_
Low-retention, filtered pipette tips	Major supplier	_
0.2-mL PCR 8-strip tubes	Major supplier	-
15-mL conical tube	Major supplier	-
DNA LoBind <sup>®</sup> Tubes, 1.5 mL	Eppendorf	0030108051
DNA LoBind <sup>®</sup> Tubes, 5.0 mL	Eppendorf	0030108310
Qubit™ Assay Tubes	Thermo Fisher Scientific	Q32856

# Equipment

Material	Supplier	Catalog no.
Microcentrifuge for 1.5–2.0-mL tubes	Major supplier	_
Microcentrifuge for 0.2-mL tubes	Major supplier	_
Vortexer	Major supplier	_
Digital timer	Major supplier	_
Eppendorf ThermoMixer <sup>®</sup> C	Eppendorf	5382000023
6-tube magnetic separation rack for 1.5-mL tubes	New England Biolabs	S1506S
Low-profile magnetic separation stand for 0.2 mL, 8-strip tubes	V&P Scientific, Inc.	VP772F4-1
Qubit™ 3.0 Fluorometer	Thermo Fisher Scientific	Q33216
Agilent® 2100 Bioanalyzer	Agilent Technologies	G2940CAG
Or,		
Agilent <sup>®</sup> 4200 TapeStation System	Agilent Technologies	G2991AA

# **Best practices**

- Use low-retention filtered pipette tips.
- When working with BD Rhapsody<sup>™</sup> Enhanced Cell Capture Beads, use low-retention filtered tips and LoBind<sup>®</sup>
  Tubes.

# Never vortex the beads. Pipet-mix only.

- Bring Agencourt AMPure XP magnetic beads to room temperature (15–25 °C) before use. See the AMPure XP User's Guide for information.
- Remove supernatants without disturbing AMPure XP magnetic beads.

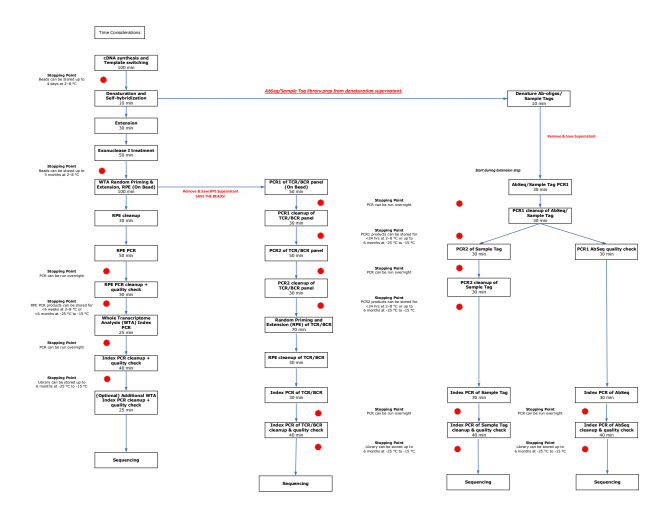
# Additional documentation

- BD Rhapsody™ Single-Cell Analysis System Instrument User Guide (23-21336)
- BD Rhapsody™ Express Single-Cell Analysis System Instrument User Guide (23-21332)

# Safety information

For safety information, see the BD Rhapsody<sup>TM</sup> Single-Cell Analysis Instrument User Guide (23-21336) or the BD Rhapsody<sup>TM</sup> Express Single-Cell Analysis System Instrument User Guide (23-21332).

# Time considerations



# **Procedure**

Perform the experiment on the BD Rhapsody™ Single-Cell Analysis system following either the:

BD Rhapsody™ Single-Cell Analysis System Instrument User Guide (23-21336)

STOP after the section "Washing the Cell Capture Beads" and follow this protocol from Preparing BD Rhapsody™ Enhanced Cell Capture Beads for TCR/BCR full length, WTA, BD<sup>®</sup> AbSeq, and Sample Tag library amplification and subsequent steps.

or

• BD Rhapsody™ Express Single-Cell Analysis System Instrument User Guide (23-21332)

STOP after the section "Washing the Cell Capture Beads" and follow this protocol from Preparing BD Rhapsody™ Enhanced Cell Capture Beads for TCR/BCR full length, WTA, BD<sup>®</sup> AbSeq, and Sample Tag library amplification and subsequent steps.

Ensure that the intended total cell load is between 7,500–20,000 single cells for this protocol. Cell load below or above this recommended range may not be suitable for current protocol configuration. Then proceed as described in the following procedure.

# Preparing BD Rhapsody™ Enhanced Cell Capture Beads for TCR/BCR full length, WTA, BD® AbSeq, and Sample Tag library amplification

# cDNA synthesis and template switching

Thaw reagents (except for the enzymes) in the BD Rhapsody<sup>TM</sup> cDNA Kit (Cat. No. 633773) at room temperature. Keep enzymes at -25 °C to -15 °C.

Note: This section should be performed in the pre-amplification workspace.

- 1 Set a thermomixer to 42 °C.
- 2 In a new 1.5-mL LoBind<sup>®</sup> tube, pipet the following reagents.

### cDNA/template switching mix

Component	For 1 library (μL)	For 1 library with 20% overage (µL)
RT Buffer	40	48
dNTP	20	24
RT 0.1 M DTT	10	12
Bead RT/PCR Enhancer	12	14.4
RNase Inhibitor	10	12
Reverse Transcriptase	10	12
Nuclease-free water	98	117.6
Total	200	240

- 3 Gently vortex mix, briefly centrifuge, and place back on ice.
- 4 Place the tube of washed Enhanced Cell Capture Beads on a 1.5-mL tube magnet for ≥2 minutes. Remove the supernatant.
- 5 Remove the tube from the magnet and pipet 200 µL of cDNA mix into the beads. Pipet-mix.

Note: Keep the prepared cDNA mix with beads on ice until the suspension is transferred in the next step.

- **6** Transfer the bead suspension to a new 1.5-mL LoBind<sup>®</sup> tube.
- 7 Incubate the bead suspension on the thermomixer at 1,200 rpm and 42 °C for 30 minutes.

# Shaking is critical for this incubation!

8 While the bead suspension is still incubating at 1,200 rpm and 42 °C, in a new 1.5-mL LoBind<sup>®</sup> tube, pipet the following reagents.

Note: Prepare the TSO mix approximately within 2 minutes before the 30 minutes incubation at 42 °C is finished.

**USE IMMEDIATELY!** 

#### TSO mix

Component	For 1 library (μL)	For 1 library with 20% overage (µL)
TSO	6	7.2
1M MgCl2	2	2.4
Total	8	9.6

- 9 Gently vortex mix, briefly centrifuge, and keep on ice.
- 10 Add 8 µL of TSO mix to the reaction, gently pipet-mix, and incubate on the thermomixer for another 30 minutes at 1,200 rpm and 42 °C.

STOPPING POINT: BD Rhapsody™ Enhanced Cell Capture Beads can be stored up to 4 days at 2–8 °C after template switching.

If stopping after template switching:

- Place the bead suspension on the 1.5-mL tube magnet until the solution is clear (≤1 minute).
- Carefully remove and appropriately discard the supernatant without disturbing the beads and while leaving the tube on the magnet.
- Remove the tube from the magnet, and with a low-retention tip, pipet 75 µL Elution Buffer to gently resuspend the beads. Do not vortex.
- Store the beads at 2–8 °C for up to 4 days.
- 11 If using the BD Rhapsody™ Single-Cell Analysis System Instrument User Guide, view the BD Rhapsody™ scanner image analysis to see if the analysis metrics passed.

# Denaturation and self-hybridization

Thaw reagents for TCR/BCR Extension at room temperature. Keep TCR/BCR Extension enzyme at -25 °C to -15 °C.

1 Set one thermomixer to 37 °C, a second thermomixer to 25 °C, and a third thermomixer to 95 °C.

**Note**: If the BD Rhapsody<sup>™</sup> Enhanced Cell Capture Beads were stored after template switching, briefly centrifuge and proceed to **step 4**.

- 2 Place the tube of Enhanced Cell Capture Beads with cDNA mix on a 1.5-mL tube magnet for ≤1 minute.
  Remove the supernatant.
- 3 Remove the tube from the magnet and pipet 75  $\mu$ L of Elution Buffer into the tube. Pipet-mix.
- 4 To denature, incubate the tube in the following order:
  - a Ensure that the beads are resuspended. Pipet-mix to resuspend, if needed.
  - **b** Incubate the sample at 95 °C in a thermomixer (no shaking) for 5 minutes. Immediately after the completion of the 95 °C incubation, slightly open the lid of the tube to release air pressure within the tube.
- 5 Briefly centrifuge the tube, then immediately place the tube on a 1.5-mL magnet for ≥30 seconds until clear. Keep the supernatant. Remove the supernatant and transfer to a new 1.5-mL LoBind<sup>®</sup> tube. This contains the AbSeq/ST supernatant products. To minimize AbSeq/ST contamination in the TCR/BCR and WTA libraries, ensure that all liquid is removed from the tube. Keep the supernatant tube at 4 °C until ready to proceed to Performing AbSeq/Sample Tag PCR1 on page 16.
- 6 Resuspend the beads in 1.5 mL of Hybridization Buffer.
- 7 Incubate the bead suspension on the thermomixer at 1,200 rpm and 25 °C for 2 minutes.
- **8** Briefly centrifuge after 25 °C incubation. Be careful when opening the tube lid. If there are droplets on the lid, use a P10 to transfer the volume into the supernatant.

### TCR/BCR extension

- 1 Set a thermomixer to 37 °C.
- 2 Ensure all reagents other than the TCR/BCR Extension enzyme are at room temperature.
- 3 In a new 1.5-mL LoBind<sup>®</sup> tube, pipet the following reagents.

## TCR/BCR extension mix

Component	For 1 librαry (μL)	For 1 library with 20% overage (μL)
TCR/BCR Extension Buffer	20	24
dNTP	20	24
TCR/BCR Extension Enzyme	10	12
Nuclease-free water	150	180
Total	200	240

- 4 Gently vortex mix, briefly centrifuge, and keep at room temperature.
- **5** Briefly spin the tube with the bead suspension.
- 6 Place the tube of Enhanced Cell Capture Beads on a 1.5-mL tube magnet for ≤2 minutes. Remove the supernatant.
- 7 Remove the tubes from magnet and resuspend using 200 µL of TCR/BCR extension mix. Pipet-mix.
- 8 Incubate the bead suspension on a thermomixer at 1,200 rpm and 37 °C for 30 minutes.

**Note:** During TCR/BCR Extension incubation, begin **AbSeq/Sample Tag PCR1**. See Performing AbSeq/Sample Tag PCR1 on page 16. You can leave the AbSeq/Sample Tag PCR1 reaction in the thermocycler when complete. TCR/BCR PCR1 will be performed after Purifying RPE PCR amplification product

(single-sided cleanup) on page 14. All PCR1 product purification (TCR/BCR and AbSeq/Sample Tag) can be done at the same time.

9 Briefly spin the tube with the beads suspension and place the tube on ice.

# Treating the sample with Exonuclease I

Thaw reagents for Exonuclease I treatment at room temperature. Keep Exonuclease I enzyme at -25 °C to -15 °C.

- 1 Set one thermomixer to 37 °C and a second thermomixer to 80 °C.
- 2 In a new 1.5-mL LoBind<sup>®</sup> tube, pipet the following reagents.

### Exonuclease I mix

Kit component	For 1 library (μL)	For 1 library with 20% overage (μL)
10X Exonuclease I Buffer	20	24
Exonuclease I	10	12
Nuclease-free water	170	204
Total	200	240

- 3 Gently vortex mix, briefly centrifuge, and keep at room temperature.
- 4 Place the tube of Enhanced Cell Capture Beads with TCR/BCR Extension mix on a 1.5-mL tube magnet for ≤1 minute. Remove the supernatant.
- 5 Remove the tube from the magnet and pipet 200 µL Exonuclease I mix into the tube. Pipet-mix.
- 6 Incubate the bead suspension on thermomixer at 1,200 rpm and 37 °C for 30 minutes.

**Note:** If only one thermomixer is available, allow it to equilibrate to 80 °C before starting the inactivation incubation. Place the samples on ice until that temperature is reached.

- 7 Incubate the bead suspension on thermomixer (no shaking) at 80 °C for 20 minutes.
- 8 Place the tube on ice for ~1 minute.
- **9** Briefly spin the tube with the bead suspension.
- 10 Place the tube on the magnet for ≤1 minute until clear. Remove the supernatant.
- 11 Remove the tube from the magnet and pipet 200 µL of cold Bead Resuspension Buffer into the tube. Pipetmix.

**STOPPING POINT:** Exonuclease I-treated beads can be stored at 2–8 °C for up to 3 months.

**12** Proceed to library preparation.

# Performing random priming and extension (RPE) on BD Rhapsody™ Enhanced Cell Capture Beads with cDNA

This section describes how to generate random priming products. First, random primers are hybridized to the cDNA on the BD Rhapsody™ Enhanced Cell Capture Beads, followed by extension with an enzyme.

**Note:** Perform this procedure in the pre-amplification workspace.

- 1 Set a heat block to 95 °C, one thermomixer to 37 °C, and one thermomixer to 25 °C.
- 2 In a new 1.5-mL LoBind<sup>®</sup> tube, pipet the following reagents.

## Random primer mix

Kit component	For 1 library (μL)	For 1 library with 20% overage (μL)
WTA Extension Buffer	20	24
WTA Extension Primers	20	24
Nuclease-free water	134	160.8
Total	174	208.8

3 Pipet-mix the Random Primer Mix and keep at room temperature.

**Note:** Use the entire sample of beads. Sub-sampling beads is not recommended for TCR/BCR FL + WTA + AbSeq + Sample Tag combination assays.

- 4 Resuspend the Exonuclease I-treated Enhanced Cell Capture Beads with a pipette..
- **5** Place the tube of Exonuclease I-treated beads in Bead Resuspension Buffer on the 1.5-mL magnet for 2 minutes. Remove the supernatant.
- **6** Briefly centrifuge the tube, then place the tube on a 1.5-mL magnet for 2 minutes. Remove and dispose of the supernatant.
- 7 Remove the tube with the Enhanced Cell Capture Beads from the magnet, and use a low-retention tip to pipet  $174 \mu L$  of Random Primer Mix into the tube. Pipet-mix 10 times to resuspend the beads.
- 8 Incubate the tube in the following order:
  - a 95 °C in a heat block (no shaking) for 5 minutes.
  - **b** Thermomixer at 1,200 rpm and at 37 °C for 5 minutes.
  - c Thermomixer at 1,200 rpm and at 25 °C for 15 minutes.
- 9 Briefly centrifuge the tube and keep it at room temperature.
- **10** In a new 1.5-mL LoBind<sup>®</sup> tube, pipet the following reagents.

# Extension enzyme mix

Kit component	For 1 librαry (μL)	For 1 library with 20% overage (µL)
dNTP	8	12
Bead RT/PCR Enhancer	12	18
WTA Extension Enzyme	6	9
Total	26	39

- 11 Pipet-mix the Extension Enzyme Mix.
- 12 Pipet 26  $\mu$ L of the Extension Enzyme Mix into the sample tube containing the beads (for a total volume of 200  $\mu$ L) and keep at room temperature until ready.
- **13** Program the thermomixer.
  - a 1,200 rpm and at 25 °C for 10 minutes
  - **b** 1,200 rpm and at 37 °C for 15 minutes
  - c 1,200 rpm and at 45 °C for 10 minutes
  - **d** 1,200 rpm and at 55 °C for 10 minutes

**IMPORTANT** Set the ramp rates at maximal and set "Time Mode" to "Temp Control" before the program begins.

- 14 Place the tube from **step 12** in the thermomixer. The program takes approximately 55 minutes. Remove the tube after the program is finished.
- 15 Place the tube in a 1.5-mL tube magnet and remove the supernatant.
- 16 Remove the tube from the magnet and resuspend the beads in 205  $\mu$ L of Elution Buffer using a P200 pipette.
- 17 To denature the random priming products off the beads, pipet to resuspend the beads. Incubate the sample at 95 °C in a heat block for 5 minutes (no shaking). Immediately after the completion of the 95 °C incubation, slightly open the lid of the tube to release air pressure within the tube.

Note: Do not incubate for more than 5 minutes.

- 18 Place the tube in a 1.5-mL tube magnet. Immediately transfer 200 µL of the supernatant containing the Random Primer Extension Product (RPE Product) to a new 1.5-mL LoBind<sup>®</sup> tube and keep at room temperature. Proceed to Purifying RPE product in the following section.
- 19 Pipet 200 µL of cold Bead Resuspension Buffer to the tube with leftover beads. Gently resuspend the beads by pipet-mixing only. Do not vortex. Store the beads on ice or at 4 °C in the pre-amplification workspace until needed.

Note: These beads will be used for TCR/BCR target specific amplification. DO NOT THROW AWAY!

# **Purifying RPE product**

This section describes how to perform a single-sided AMPure cleanup, which removes primer dimers and other small molecular weight by-products. The final product is purified single-stranded DNA.

Note: Perform the purification in the pre-amplification workspace.

1 In a new 15-mL conical tube, prepare 10 mL of fresh 80% (v/v) ethyl alcohol by pipetting 8.0 mL of absolute ethyl alcohol to 2.0 mL of nuclease-free water (from major supplier). Vortex the tube for 10 seconds.

Note: Make fresh 80% ethyl alcohol and use within 24 hours.

- 2 Bring Agencourt AMPure XP magnetic beads to room temperature. Vortex the AMPure XP magnetic beads at high speed for 1 minute until the beads are fully resuspended.
- 3 Pipet 320  $\mu$ L of AMPure XP magnetic beads into the tube containing the 200  $\mu$ L of RPE product supernatant. Pipet-mix at least 10 times, then briefly centrifuge.
- 4 Incubate at room temperature for 10 minutes.
- 5 Place the tube on the 1.5-mL tube magnet for 5 minutes. Remove the supernatant.
- **6** Keeping the tube on the magnet, gently add 1 mL of fresh 80% ethyl alcohol into the tube and incubate for 30 seconds. Remove the supernatant.
- 7 Repeat step 6 for a total of two washes.
- **8** Keeping the tube on the magnet, use a small-volume pipette to remove and discard any residual supernatant from the tube.
- 9 Air-dry the beads at room temperature for 5 minutes or until the beads no longer look glossy.
- 10 Remove the tube from the magnet and resuspend the bead pellet in 40  $\mu$ L of Elution Buffer. Pipet-mix the suspension at least 10 times until the beads are fully suspended.
- 11 Incubate the sample at room temperature for 2 minutes. Briefly centrifuge the tube to collect the contents at the bottom.
- 12 Place the tube on the magnet until the solution is clear, usually ~30 second.
- 13 Pipet the eluate ( $\sim$ 40  $\mu$ L) to a new PCR tube. This is the purified RPE product.

# Performing RPE PCR

This section describes how to generate more RPE product through PCR amplification, so that there are multiple copies of each random-primed molecule.

1 In the pre-amplification workspace, in a new 1.5-mL LoBind<sup>®</sup> tube, pipet the following components.

### **RPE PCR mix**

Kit component	For 1 library (μL)	For 1 library with 20% overage (µL)
PCR MasterMix	60	72
Universal Oligo	10	12
WTA Amplification Primer	10	12
Total	80	96

- 2 Add 80  $\mu$ L of the RPE PCR Mix to the tube with the 40  $\mu$ L of purified RPE product. Pipet-mix 10 times.
- 3 Split the RPE PCR reaction mix into two PCR tubes with 60  $\mu$ L of reaction mix per tube.
- 4 Bring the reaction to the post-amplification workspace and run the following PCR program.

# PCR program

Step	Cycles	Temperature	Time
Hot start	1	95 ℃	3 min
Denaturation	Refer to the	95 ℃	30 s
Annealing	following table, Recommended	60 °C	1 min
Extension	number of PCR cycles.*	72 °C	1 min
Final extension	1	72 °C	2 min
Hold	1	4 °C	∞
*Suggested PCR cycles might need to be optimized for different cell types and cell number.			

### Recommended number of PCR cycles

Number of cells in RPE PCR	Recommended PCR cycles for resting PBMCs
7,500	13
20,000	11

5 When the RPE PCR reaction is complete, briefly centrifuge to collect the contents at the bottom of the tubes.

# Purifying RPE PCR amplification product (single-sided cleanup)

This section describes how to perform a single-sided AMPure cleanup to remove unwanted small molecular weight products from the RPE products. The final product is purified double-stranded DNA (~200–2,000 bp).

**Note:** Perform the purification in the post-amplification workspace.

- 1 Combine the two 60- $\mu$ L RPE PCR reactions into a new 1.5-mL tube.
- **2** Briefly centrifuge the tube with the RPE PCR product.

**IMPORTANT** It is critical for the final volume to be exactly 120  $\mu$ L to achieve the appropriate size selection of the purified RPE PCR product.

3 In a new 15-mL conical tube, prepare 5 mL of fresh 80% (v/v) ethyl alcohol by pipetting 4.0 mL of absolute ethyl alcohol to 1.0 mL of nuclease-free water (from major supplier). Vortex the tube for 10 seconds.

Note: Make fresh 80% ethyl alcohol and use within 24 hours.

- 4 Bring AMPure XP magnetic beads to room temperature. Vortex the AMPure XP magnetic beads at high speed for 1 minute until the beads are fully resuspended.
- 5 Pipet 96  $\mu$ L of AMPure XP magnetic beads into the tube containing 120  $\mu$ L of RPE PCR product. Pipet-mix at least 10 times, then briefly centrifuge the samples.
- 6 Incubate at room temperature for 5 minutes.
- 7 Place the 1.5-mL LoBind<sup>®</sup> tube on the magnet for 5 minutes.
- 8 Keeping the tube on the magnet, gently add 200 µL of fresh 80% ethyl alcohol into the tube and incubate for 30 seconds. Remove the supernatant without disturbing the beads.
- **9** Repeat **step 8** once for a total of two washes.
- **10** Keeping the tube on the magnet, use a small-volume pipette to remove any residual supernatant from the tube.
- 11 Air-dry the beads at room temperature for 3 minutes.
- 12 Remove the tube from the magnet and pipet 40  $\mu$ L of Elution Buffer into the tube. Pipet-mix the suspension at least 10 times until the beads are fully suspended.
- 13 Incubate the sample at room temperature for 2 minutes. Briefly centrifuge the tube to collect the contents at the bottom..
- 14 Place the tube on the magnet until the solution is clear, usually ~30 seconds.
- **15** Pipet the eluate (~40  $\mu$ L) into a new 1.5-mL LoBind<sup>®</sup> tube. The RPE PCR product is ready for WTA Index PCR.

**STOPPING POINT:** The RPE PCR libraries can be stored at -20 °C for up to 6 months or 4 °C for up to 6 weeks.

- 16 Quantify and perform quality control of the RPE PCR products with a Qubit Fluorometer using the Qubit dsDNA HS Assay and the Agilent 2100 Bioanalyzer using the Agilent High Sensitivity DNA Kit.
  - **a** The expected concentration from the Qubit Fluorometer is  $\sim 0.5$  to 10 ng/ $\mu$ L.
  - **b** The Bioanalyzer trace should show a broad peak from ~150 to 2,000 bp. Use the concentration from 200 to 600 bp to calculate how much template to add into Index PCR. Refer to the blue-boxed regions in the sample trace images in Figure 1.

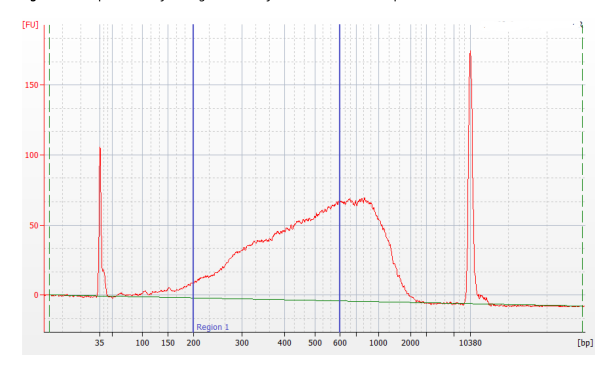


Figure 1 Sample Bioanalyzer High Sensitivity DNA trace - RPE PCR product trace

# Performing AbSeq/Sample Tag PCR1

This section describes how to amplify AbSeq/Sample Tag products through PCR.

1 In the pre-amplification workspace, pipet the following reagents into a new 1.5-mL LoBind® tube on ice.

AbSeq/Sam	ple Tag PC	R1 reaction mix
-----------	------------	-----------------

Component	For 1 library (μL)	For 1 library with 20% overage (µL)
PCR MasterMix	100	120
Universal Oligo	10	12
Sample Tag PCR1 Primer	1	1.2
BD® AbSeq Primer	10	12
Nuclease-free water	12	14.4
Total	133	159.6

- 2 Gently vortex mix, briefly centrifuge, and place back on ice.
- 3 In a new 1.5-mL tube, pipet 133  $\mu$ L of the AbSeq/Sample Tag PCR1 reaction mix. Add 67  $\mu$ L of the AbSeq/Sample Tag product from **step 5** in Denaturation and self-hybridization on page 9. Pipet-mix 10 times. Do not vortex.
- 4 Pipet 50  $\mu$ L AbSeq/Sample Tag PCR1 reaction mix into each of four 0.2-mL PCR tubes. Transfer any residual mix to one of the tubes.
- **5** Bring the reaction mix to the post-amplification workspace.
- 6 Program the thermal cycler.

# Thermal cycler program

Step	Cycles	Temperature	Time
Hot start	1	95 ℃	3 min
Denaturation		95 ℃	30 s
Annealing	10-11*	60 °C	30 s
Extension		72 °C	1 min
Final extension	1	72 °C	5 min
Hold	1	4°C	∞
*Suggested PCR cycles might need to be optimized for different cell types and cell number.			

# Recommended number of PCR cycles

Number of cells in PCR1	Recommended PCR cycles for resting PBMCs
7,500 - 10,000	11
20,000	10

**STOPPING POINT:** The PCR can run overnight.

- **7** After PCR, briefly centrifuge the tubes.
- **8** Pipet-mix and combine the four reactions into a new 1.5-mL LoBind<sup>®</sup> tube, labeled AbSeq/Sample Tag PCR1. Keep the tube on ice.

# Performing TCR/BCR PCR1

1 Obtain beads from Step 19 on page 13 of "Performing random priming and extension (RPE) on BD Rhapsody™ Enhanced Cell Capture Beads with cDNA."

Note: Use the entire sample of beads. Sub-sampling beads is not recommended for TCR/BCR FL + WTA + AbSeq + Sample Tag combination assays.

2 In the pre-amplification workspace, pipet the following reagents into a new 1.5-mL LoBind® tube.

# PCR1 reaction mix

Component	For 1 library (μL)	For 1 library with 20% overage (µL)
PCR MasterMix	100	120
TCR/BCR Universal Oligo N1	10	12
Bead RT/PCR Enhancer	12	14.4
*TCR N1 primer	2.4	2.88
*BCR N1 primer	2.4	2.88
Nuclease-free water	73.2	87.84
Total	200	240

\*If only doing TCR or BCR amplification, replace N1 primer volume with nuclease-free water. For example, if only doing TCR amplification, replace BCR N1 primer with nuclease-free water.

- 3 Gently vortex mix, briefly centrifuge, and place back on ice.
- 4 Briefly spin the tube with the bead suspension. Place the tube of beads in Bead Resuspension Buffer on a 1.5-mL magnet for ≤1 minute. Remove the supernatant.
- 5 Remove the tube from the magnet and resuspend the beads in 200  $\mu$ L of PCR1 reaction mix. Do not vortex.
- **6** Ensuring that the beads are fully resuspended, pipet 50 μL of PCR1 reaction mix with beads into each of four 0.2-mL PCR tubes. Transfer any residual mix to one of the tubes.
- 7 Bring the reaction mix to the post-amplification workspace.
- 8 Program the thermal cycler as follows.

### Thermal cycler program

Step	Cycles	Temperature	Time
Hot start	1	95 °C*	3 min
Denaturation		95 ℃	30 s
Annealing	10-11**	60 °C	1 min
Extension		72 °C	1 min
Final extension	1	72 °C	5 min
Hold	1	4°C	∞

<sup>\*</sup>To avoid beads settling due to prolonged incubation time on the thermal cycler before the denaturation step, it is critical to pause the instrument at 95 °C before loading the samples. Different thermal cyclers might have different pause time settings. In certain brands of thermal cyclers, however, we have observed a step-skipping error with the pause/unpause functions. To ensure that the full 3-minute denaturation is not skipped, verify that the pause/unpause functions are working correctly on your thermal cycler. To avoid the step-skipping problem, a 1-minute 95 °C pause step can be added immediately before the 3-minute 95 °C denaturation step.

\*\*Suggested PCR cycles might need to be optimized for different cell types and cell number.

# Suggested number of PCR cycles

Number of cells in PCR1	Suggested PCR cycles for resting PBMCs
7,500 - 10,000	11
20,000	10

9 Ramp the heated lid and heat block of the post-amplification thermal cycler to ≥95 °C by starting the thermal cycler program and then pausing it.

**Note:** Do not proceed to thermal cycling until each tube is gently mixed by pipette to ensure uniform bead suspension.

**10** For each 0.2-mL PCR tube, gently pipet-mix, immediately place the tube in thermal cycler, and unpause the thermal cycler program.

**STOPPING POINT:** The PCR can run overnight, but proceed with purification within 24 hours after PCR.

- **11** After PCR, briefly centrifuge the tubes.
- 12 Pipet-mix and combine the four reactions into a new 1.5-mL LoBind $^{\textcircled{8}}$  tube.
- 13 Place the 1.5-mL tube on the magnet for ≤1 minute. Retain the supernatant that contains the TCR/BCR PCR1 products. Carefully pipet the supernatant (TCR/BCR PCR1 products) into the new 1.5-mL LoBind<sup>®</sup> tube without disturbing the beads.

**Note:** (Optional) Remove the tube with the BD Rhapsody<sup>TM</sup> Enhanced Cell Capture Beads from the magnet and pipet 200  $\mu$ L of cold Bead Resuspension Buffer into the tube. Pipet-mix. Do not vortex. Store the beads at 2–8 °C in the post-amplification workspace.

# Purifying TCR/BCR and AbSeq/Sample Tag PCR1 products

This section describes how to perform a single-sided AMPure cleanup to remove primer dimers from the TCR/BCR and AbSeq/Sample Tag PCR1 products. The final product is purified double-stranded DNA.

**Note:** Perform the purification in the post-amplification workspace.

1 In a new 5.0-mL LoBind<sup>®</sup> tube, prepare 5 mL of fresh 80% (v/v) ethyl alcohol by combining 4.0 mL absolute ethyl alcohol, molecular biology grade, with 1.0 mL nuclease-free water. Vortex the tube for 10 seconds to mix.

Note: Make fresh 80% ethyl alcohol and use it within 24 hours.

- 2 Bring the AMPure XP magnetic beads to room temperature. Vortex on high speed for 1 minute until the beads are fully resuspended.
- 3 To 200  $\mu$ L of PCR1 products, pipet:
  - AbSeq/Sample Tag libraries: 280 μL AMPure beads. From step 8 on page 17 in "Performing AbSeq/Sample Tag PCR1."
  - TCR/BCR libraries: 140 µL AMPure beads. From step 13 on page 18 in "Performing TCR/BCR PCR1."
- 4 Pipet-mix 10 times. Incubate at room temperature for 5 minutes.
- 5 Place the 1.5-mL LoBind<sup>®</sup> tube on the magnet for 5 minutes.
- **6** Keeping the tube on the magnet, gently add 500  $\mu$ L of fresh 80% ethyl alcohol into the tube and incubate for 30 seconds. Remove the supernatant.
- 7 Repeat step 6 once for a total of two washes.
- **8** Keeping the tube on the magnet, use a small-volume pipette to remove and discard any residual supernatant from the tube.
- **9** Air-dry the beads at room temperature for 5 minutes.
- 10 Remove the tube from the magnet and resuspend the bead pellet in 50  $\mu$ L of Elution Buffer. Vigorously pipet-mix until the beads are uniformly dispersed. Small clumps do not affect performance.
- 11 Incubate at room temperature for 2 minutes and briefly centrifuge.
- 12 Place the tube on the magnet until the solution is clear, usually ~30 seconds.
- 13 Pipet the eluate ( $\sim$ 50  $\mu$ L) into a new 1.5-mL LoBind<sup>®</sup> tube separately (purified TCR/BCR PCR1 and AbSeq/Sample Tag PCR1 products).

**STOPPING POINT:** Store at 2–8 °C before proceeding within 24 hours or at -25 °C to -15 °C for up to 6 months.

# Quantifying BD® AbSeq/Sample Tag PCR1 products

- 1 Measure the yield of the largest peak of the BD<sup>®</sup> AbSeq/ Sample Tag PCR1 product (~150 bp) by using the Agilent 2100 Bioanalyzer with the High Sensitivity Kit assay. Follow the manufacturer's instructions.
- 2 Based on the yield of the largest peak, dilute an aliquot of BD<sup>®</sup> AbSeq/Sample Tag PCR1 product to 0.1–1.1 ng/μL with Nuclease-Free water before index PCR of BD<sup>®</sup> AbSeq PCR1 products. See Performing TCR/BCR and AbSeq/Sample Tag index PCR on page 24.

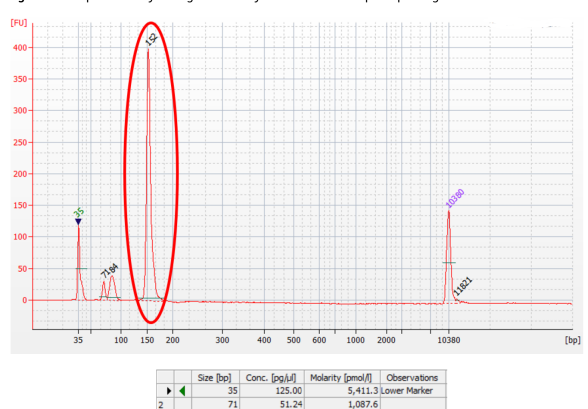


Figure 2 Sample Bioanalyzer High Sensitivity DNA trace - AbSeq/Sample Tag PCR1

# Performing TCR/BCR and Sample Tag PCR2 on the PCR1

84

# **products**This section describes how to amplify TCR/BCR and Sample Tag products through PCR. The PCR primers include

123.99

2,225.8

partial Illumina sequencing adapters that enable the additions of full-length Illumina sequencing indices in the next PCR.

1 In the pre-amplification workspace, pipet reagents into a new 1.5-mL LoBind $^{\circledR}$  tube on ice.

TCR/RCR	DCD2	roaction	miv
II K/DI K	PIRI	reaction	mıx

Component	For 1 library (μL)	For 1 library with 20% overage (µL)
PCR MasterMix	25	30
TCR/BCR Universal Oligo N2	2	2.4
*TCR or BCR N2 primer	6	7.2
Nuclease-free water	12	14.4
Total	45	54
*PCR2 reaction mixes for TCR and BCR are made separately.		

# Sample Tag PCR2 reaction mix

Component	For 1 library (μL)	For 1 library with 20% overage (µL)
PCR MasterMix	25	30
Universal Oligo	2	2.4
Sample Tag PCR2 Primer	3	3.6
Nuclease-free water	15	18
Total	45	54

- **2** Gently vortex mix, briefly centrifuge, and place back on ice.
- 3 Bring the PCR2 reaction mix to the post-amplification workspace.
- 4 Pipet 5.0  $\mu$ L of PCR1 products into 45  $\mu$ L of TCR/BCR or Sample Tag PCR2 reaction mix.
- **5** Gently vortex and briefly centrifuge.
- 6 Program the thermal cycler.

# Sample Tag PCR2

Step	Cycles	Temperature	Time
Hot start	1	95 ℃	3 min
Denaturation		95 ℃	30 s
Annealing	10*	66 °C	30 s
Extension		72 °C	1 min
Final extension	1	72 °C	5 min
Hold	1	4 °C	∞
* Suggested PCR cycles might need to be optimized for different cell types and cell number.			

### TCR and BCR PCR2

Step	Cycles	Temperature	Time
Hot start	1	95 ℃	3 min
Denaturation		95 ℃	30 s
Annealing	20	60 °C	1 min
Extension		72 °C	1 min
Final extension	1	72 °C	5 min
Hold	1	4 °C	∞

**STOPPING POINT:** The PCR can run overnight.

# Purifying TCR/BCR and Sample Tag PCR2 products

This section describes how to perform a single-sided AMPure cleanup to remove primer dimers from the TCR/BCR PCR2 products. The final product is purified double-stranded DNA.

Note: Perform PCR2 purification in the post-amplification workspace.

1 In a new 5.0-mL LoBind<sup>®</sup> tube, prepare 5 mL fresh 80% (v/v) ethyl alcohol by combining 4 mL absolute ethyl alcohol, molecular biology grade, with 1 mL of nuclease-free water. Vortex the tube for 10 seconds to mix.

Note: Make fresh 80% ethyl alcohol and use it within 24 hours.

- 2 Bring AMPure XP beads to room temperature and vortex at high speed for 1 minute until beads are fully resuspended.
- **3** To 50.0 μL PCR2 products, pipet:
  - AbSeq/Sample Tag libraries: 60 μL AMPure beads.
  - TCR/BCR libraries: 35 µL AMPure beads.
- 4 Pipet-mix 10 times and incubate at room temperature for 5 minutes.
- 5 Place the tube on the strip tube magnet for 3 minutes. Remove the supernatant.
- **6** Keeping the tube on the magnet, gently add 200  $\mu$ L of fresh 80% ethyl alcohol into the tube and incubate for 30 seconds. Remove the supernatant.
- 7 Repeat step 6 once for a total of two washes.
- **8** Keeping the tube on the magnet, use a small-volume pipette to remove and discard any residual supernatant from the tube.
- 9 Air-dry the beads at room temperature for 3 minutes.
- 10 Remove the tube from the magnet and resuspend the bead pellet in 50  $\mu$ L of Elution Buffer. Pipet-mix until the beads are fully resuspended.
- 11 Incubate at room temperature for 2 minutes and briefly centrifuge.
- 12 Place the tube on the magnet until the solution is clear, usually ~30 seconds.
- 13 Pipet the entire eluate ( $\sim$ 50  $\mu$ L) into a new 1.5-mL LoBind<sup>®</sup> tube separately (purified TCR/BCR and Sample Tag PCR2 products).
  - **STOPPING POINT:** Store at 2–8  $^{\circ}$ C before proceeding on the same day, or at –25  $^{\circ}$ C to –15  $^{\circ}$ C for up to 6 months.
- 14. Estimate the concentration with a Qubit Fluorometer using the Qubit dsDNA HS Assay Kit. Follow the manufacturer's instructions.

# Performing random priming and extension (RPE) on TCR/BCR PCR2 products

Note: Perform TCR/BCR Random Priming the purification in the post-amplification workspace.

- 1 Dilute an aliquot of the TCR/BCR PCR2 products with nuclease-free water to 1.0  $ng/\mu L$ .
- 2 In pre-amplification workspace, pipet reagents into a new 1.5 mL LoBind<sup>®</sup> tube:

### Random primer mix

Component	For 1 librαry (μL)	For 1 library with 20% overage (µL)
TCR/BCR Extension Buffer	5	6
TCR/BCR Extension Primers	2.5	3
Nuclease-free water	34	40.8
Total	41.5	49.8

- 3 Pipet-mix the Random Primer Mix and keep at room temperature.
- 4 Add 41.5  $\mu$ L of Random Primer Mix + 5  $\mu$ L of 1 ng/ $\mu$ L diluted TCR/BCR PCR2 products (5 ng total concentration). Total volume of the reaction will be 46.5  $\mu$ L for Random Priming.
- 5 Perform denaturation and random priming on thermocycler using the following program:

# Program

Temperature	Time	Cycles
95 ℃	5 min	
37 °C	5 min	1
25 °C	15 min	

- **6** Briefly centrifuge the tube and keep at room temperature.
- 7 In pre-amplification workspace, pipet reagents into a new 1.5 mL LoBind<sup>®</sup> tube:

# **Primer Extension Enzyme mix**

Component	For 1 library (μL)	For 1 library with 20% overage (µL)
dNTP	2	2.4
TCR/BCR Extension Enzyme	1.5	1.8
Total	3.5	4.2

- 8 Gently vortex mix, centrifuge, and keep at room temperature.
- 9 Add 3.5  $\mu$ L Primer Extension Enzyme Mix to Random Priming Rxn tube to bring total volume up to 50  $\mu$ L. Run the following protocol on a thermocycler for Extension.

# **Protocol**

Temperature	Time	Cycles
25 °C	10 min	
37 °C	15 min	1
45 °C	10 min	ı
55 °C	10 min	

10 Remove tubes from thermocycler and prepare to purify RPE product.

# **Purifying TCR/BCR RPE product**

**Note:** Perform purification in the post-amplification workspace.

- 1 In a new 5.0-mL LoBind® tube, prepare 5 mL fresh 80% (v/v) ethyl alcohol by combining 4 mL absolute ethyl alcohol, molecular biology grade, with 1 mL of nuclease-free water. Vortex the tube for 10 seconds to mix.
  - Note: Make fresh 80% ethyl alcohol and use it within 24 hours.
- 2 Bring AMPure XP beads to room temperature and vortex at high speed for 1 minute until beads are fully resuspended.
- 3 To the TCR and BCR RPE products, add 90  $\mu$ L AMPure beads.
- 4 Pipet-mix 10 times and incubate at room temperature for 5 minutes.
- 5 Place the tube on the strip tube magnet for 3 minutes. Remove the supernatant.
- **6** Keeping the tube on the magnet, gently add 200  $\mu$ L of fresh 80% ethyl alcohol into the tube and incubate for 30 seconds. Remove the supernatant.
- 7 Repeat step 6 once for a total of two washes.
- **8** Keeping the tube on the magnet, use a small-volume pipette to remove and discard any residual supernatant from the tube.
- 9 Air-dry the beads at room temperature for 1 minute.
- 10 Remove tubes from the magnet and add 50  $\mu$ L of Elution Buffer.
- 11 Incubate at room temperature for 2 minutes and briefly centrifuge.
- 12 Place the tube on the magnet until the solution is clear, usually ≤30 seconds.
- 13 Pipet the entire eluate ( $\sim$ 50  $\mu$ L) into a new 1.5-mL LoBind<sup>®</sup> tube separately (purified TCR/BCR RPE products).

# Performing TCR/BCR and AbSeq/Sample Tag index PCR

This section describes how to generate TCR/BCR and AbSeq/Sample Tag libraries compatible with the Illumina sequencing platform, by adding full-length Illumina sequencing adapters and indices through PCR.

1 In the pre-amplification workspace, pipet reagents into a new 1.5-mL LoBind $^{\textcircled{8}}$  tube on ice.

### AbSeq and Sample Tag index PCR mix

Component	For 1 library (μL)	For 1 library with 20% overage (μL)		
PCR MasterMix	25	30		
Library Forward Primer	2	2.4		
*Library Reverse Primer 1 - 4	2	2.4		
Nuclease-free water	18 21.6			
Total 47 56.4				
*For more than one library, use different Library Reverse Primers for each AbSeq and Sample Tag library.				

# TCR/BCR index PCR mix

Component	For 1 librαry (μL)	For 1 library with 20% overage (μL)		
PCR MasterMix	25	30		
Library Forward Primer	2	2.4		
*Library Reverse Primer 1 - 4	2	2.4		
Total 29 34.8				
*For more than one library, use different Library Reverse Primers for each TCR or BCR library.				

- 2 Gently vortex mix, briefly centrifuge, and place back on ice.
- 3 Bring the TCR/BCR and AbSeq/Sample Tag Index PCR mix to post-amplification workspace.
- 4 In new 0.2 mL PCR tubes,
  - For TCR and BCR libraries, pipet 21  $\mu$ L of TCR/BCR RPE purified products into 29  $\mu$ L of TCR/BCR Index PCR mix.
  - For AbSeq library, pipet 3.0 μL of 0.1–1.1 ng/μL AbSeq PCR1 product into 47 μL AbSeq Index PCR mix. (From Quantifying BD® AbSeq/Sample Tag PCR1 products on page 19). See Conc. Index PCR input for AbSeq/Sample Tag libraries (ng/μL) on page 26.
  - For Sample Tag libraries, pipet 3.0 μL of 0.1–1.1 ng/μL Sample Tag PCR2 products into 47.0 μL index PCR mix. See Conc. Index PCR input for AbSeq/Sample Tag libraries (ng/μL) on page 26.
- **5** Gently vortex, and briefly centrifuge.
- 6 Program the thermal cycler.

# TCR/BCR index PCR

Step	Cycles	Temperature	Time
Hot start	1	95 ℃	3 min
Denaturation		95 ℃	30 s
Annealing	10	60 °C	30 s
Extension		72 °C	30 s
Final extension	1	72 ℃	1 min
Hold	1	4 °C	∞

# AbSeq/Sample Tag index PCR

Step	Cycles	Temperature	Time
Hot start	1	95 ℃	3 min
Denaturation	Refer to the	95 ℃	30 s
Annealing	following table Recommended number of PCR cycles*	60 °C	30 s
Extension		72 °C	30 s
Final extension	1	72 °C	1 min
Hold	1	4°C	∞
*Cycle number varies based on the concentration of the Sample Tag PCR2 product.			

### Recommended number of PCR cycles

Conc. Index PCR input for AbSeq/Sample Tag libraries (ng/µL)	Recommended number of PCR cycles
0.5–1.1	6
0.25–0.5	7
0.1–0.25	8

**STOPPING POINT:** The PCR can run overnight.

# Purifying TCR/BCR, AbSeq, and Sample Tag index PCR products

This section describes how to perform a single-sided AMPure cleanup to remove primer dimers from the TCR/BCR and AbSeq/Sample Tag Index PCR products. The final product is purified double-stranded DNA with full-length Illumina adapter sequences.

Note: Perform Index PCR purification in the post-amplification workspace.

1 In a new 5.0-mL LoBind<sup>®</sup> tube, prepare 5 mL fresh 80% (v/v) ethyl alcohol by combining 4 mL absolute ethyl alcohol, molecular biology grade, with 1 mL of nuclease-free water. Vortex the tube for 10 seconds to mix.

Note: Make fresh 80% ethyl alcohol, and use it within 24 hours.

- 2 Bring AMPure XP beads to room temperature and vortex at high speed for 1 minute until the beads are fully resuspended.
- 3 Briefly centrifuge all the index PCR products.
- 4 To  $50.0 \mu L$  of the AbSeq/Sample Tag index PCR products, pipet:
  - AbSeq/Sample Tag library: 40 µL AMPure beads.

Transfer 40 μL of the TCR and/or BCR index PCR product(s) to a new strip tube(s), pipet:

- TCR and BCR libraries: 26 μL AMPure beads.
- 5 Pipet-mix 10 times and incubate at room temperature for 5 minutes.
- 6 Place the tube on the strip tube magnet for 3 minutes. Remove the supernatant.
- 7 Keeping the tube on the magnet, gently add 200  $\mu$ L of fresh 80% ethyl alcohol into the tube and incubate for 30 seconds. Remove the supernatant.
- 8 Repeat step 7 for a total of two washes.
- **9** Keeping the tube on the magnet, use a small-volume pipette to remove and discard the residual supernatant from the tube.
- **10** Air-dry the beads at room temperature for 3 minutes.
- 11 Remove the tube from the magnet and resuspend the bead pellet in 50  $\mu$ L of Elution Buffer. Pipet-mix until the beads are fully resuspended.
- 12 Incubate at room temperature for 2 minutes, and briefly centrifuge.
- 13 Place the tube on the magnet until the solution is clear, usually ~30 seconds.
- 14 Pipet the entire eluate ( $\sim$ 50  $\mu$ L) into a new 1.5-mL LoBind<sup>®</sup> tube (final sequencing libraries).

**STOPPING POINT:** Store at -25 °C to -15 °C for up to 6 months until sequencing.

15 Estimate the concentration by quantifying 2  $\mu$ L of the final sequencing library with a Qubit Fluorometer using the Qubit dsDNA HS Kit to obtain an approximate concentration of PCR products to dilute for

quantification on an Agilent 2100 Bioanalyzer system using the Agilent High Sensitivity D1000. Follow the manufacturer's instructions.

Figure 3 Sample Bioanalyzer High Sensitivity DNA trace - Sample Tag index PCR product

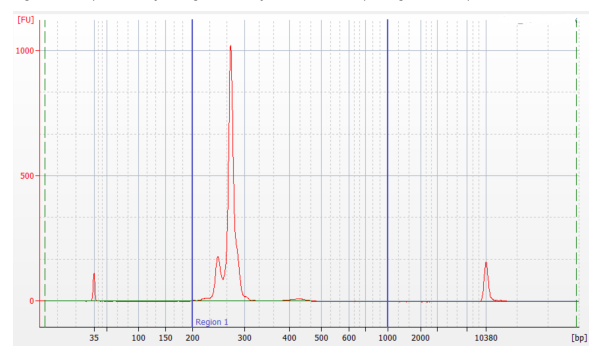


Figure 4 Sample Bioanalyzer High Sensitivity DNA trace - AbSeq index PCR product

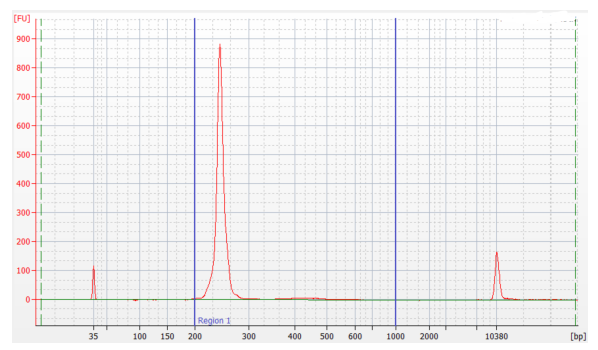
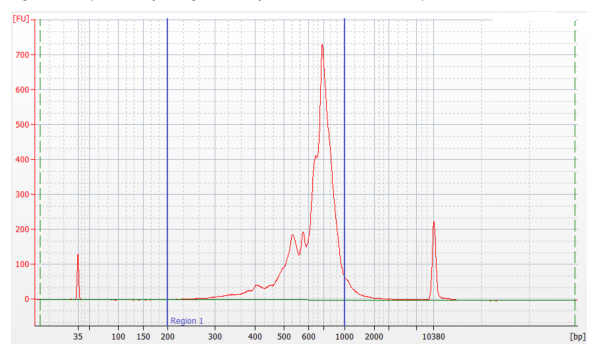


Figure 5 Sample Bioanalyzer High Sensitivity DNA trace - TCR index PCR product





# Performing WTA index PCR

This section describes how to generate mRNA libraries compatible with the Illumina sequencing platform, by adding full-length Illumina sequencing adapters and indices through PCR.

Note: Perform this procedure in the post-amplification workspace.

1 Dilute the RPE PCR products from Purifying RPE PCR amplification product (single-sided cleanup) on page 14 with Nuclease-Free water such that the concentration of the 200–600 bp peak is 2 nM. If the product concentration is <2 nM, do not dilute and continue.

For example, if the Bioanalyzer measurement of the 200–600 bp peak is 6 nM, then dilute the sample threefold with Nuclease-Free water to 2 nM.

**2** In a new 1.5-mL tube, pipet the following components.

# WTA index PCR mix

Kit component	For 1 library (μL)	For 1 library with 20% overage (µL)			
PCR MasterMix	25	30			
Library Forward Primer	5	6			
*Library Reverse Primer 1 - 4	5	6			
Nuclease-free water	5	6			
Total 40 48					
*For more than one library, use different Library Reverse Primers for each library.					

- 3 Gently vortex mix, briefly centrifuge, and place back on ice.
- 4 In a new 0.2-mL PCR tube, combine WTA Index PCR Mix with diluted RPE PCR products as follows:
  - $\alpha$  For one sample, combine 40  $\mu$ L of WTA Index PCR Mix with 10  $\mu$ L of 2 nM of RPE PCR product.
  - **b** For multiple samples, combine 35  $\mu$ L of WTA Index PCR Mix with 5  $\mu$ L of Library Reverse Primer and 10  $\mu$ L of 2 nM of RPE PCR products.
- 5 Pipet-mix 10 times.
- 6 Run the following PCR program.

# PCR program

Step	Cycles	Temperature	Time		
Hot start	1	95 ℃	3 min		
Denaturation	Refer to the	95 ℃	30 s		
Annealing	following table, Recommended	60 °C	30 s		
Extension	number of PCR cycles.*	72 °C	30 s		
Final extension	1	72 °C	1 min		
Hold	1	4 °C	∞		
*Cycle number varies based on the concentration of the RPE PCR product.					

# Recommended number of PCR cycles

Concentration of diluted RPE PCR products	Recommended number of PCR cycles
1 to <2 nM	9
2 nM	8

If the concentrations of diluted RPE PCR products are <1 nM, additional PCR cycles might be needed.

**STOPPING POINT:** The PCR can run overnight.

7 When the WTA Index PCR is complete, briefly centrifuge to collect the contents at the bottom of the tubes.

# Purifying WTA index PCR product (single-sided cleanup)

This section describes how to perform a single-sided AMPure cleanup for Illumina sequencing. The final product is purified double-stranded DNA with full-length Illumina adapter sequences.

Note: Perform the purification in the post-amplification workspace.

- 1 Add 60 µL of nuclease-free water to the WTA Index PCR product for a final volume of 110 µL.
- 2 Transfer 100  $\mu$ L of WTA Index PCR product into a new 0.2-mL PCR tube.
- 3 In a new 5.0-mL LoBind<sup>®</sup> tube, prepare 5 mL fresh 80% (v/v) ethyl alcohol by combining 4.0 mL absolute ethyl alcohol, molecular biology grade, with 1.0 mL of nuclease-free water. Vortex the tube for 10 seconds to mix.

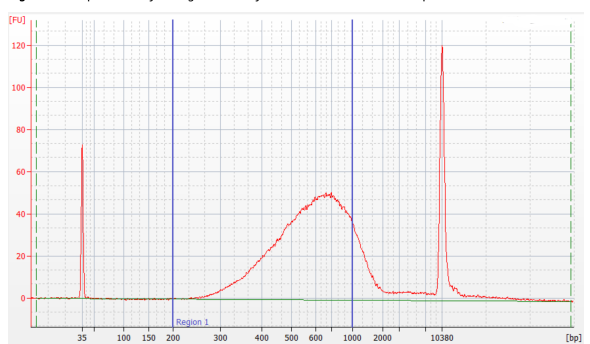
Note: Make fresh 80% ethyl alcohol, and use it within 24 hours.

- 4 Bring AMPure XP beads to room temperature and vortex at high speed for 1 minute until the beads are fully resuspended.
- 5 Add 65  $\mu$ L of AMPure XP magnetic beads to the 0.2-mL PCR tube from step 2.
- 6 Pipet-mix 10 times and incubate at room temperature for 5 minutes.
- 7 Place the tube on the strip tube magnet for 3 minutes. Remove the supernatant.
- **8** Keeping the tube on the magnet, gently add 200  $\mu$ L of fresh 80% ethyl alcohol into the tube and incubate for 30 seconds. Remove the supernatant.
- 9 Repeat step 8 for a total of two washes.
- 10 Keeping the tubes on the magnet, use a small-volume pipette to remove any residual supernatant from the tube.
- 11 Leave the tubes open on the magnet to dry the AMPure XP magnetic beads at room temperature for ~1 minute. Do not over-dry the AMPure XP magnetic beads.
- 12 Pipet 30  $\mu$ L of Elution Buffer into the tubes and pipet-mix to completely resuspend the AMPure XP magnetic beads.
- 13 Incubate the samples at room temperature for 2 minutes.
- **14** Briefly centrifuge the tubes to collect the contents at the bottom.
- 15 Place the tubes on the magnet until the solution is clear, usually ~30 seconds.
- 16 Pipet the eluate ( $\sim$ 30  $\mu$ L) into new 1.5-mL LoBind<sup>®</sup> tubes. The WTA Index PCR eluate is the final sequencing libraries.

STOPPING POINT: The Index PCR libraries can be stored at -20 °C for up to 6 months until sequencing.

- 17 Quantify and perform quality control of the Index PCR libraries with a Qubit Fluorometer using the Qubit dsDNA HS Assay and the Agilent 2100 Bioanalyzer using the Agilent High Sensitivity DNA Kit.
  - **a** The expected concentration from the Qubit Fluorometer is >1  $ng/\mu L$ .
  - **b** The Bioanalyzer trace should show a peak from ~300 to 2,000 bp.

Figure 7 Sample Bioanalyzer High Sensitivity DNA trace - WTA index PCR product



# Sequencing

The sequencing depth for each library is dependent on application. For cell-type clustering, shallow sequencing is sufficient. However, for in-depth analysis, such as comparison across multiple samples, deep sequencing is advised. We recommend meeting the requirement for recursive substitution error correction (RSEC) sequencing depth of  $\geq 6$  to reach the threshold of sequencing saturation where most molecules of the library have been recovered, approximately 80%. The RSEC sequencing depth and sequencing saturation are both reported by the analysis pipeline. The actual sequencing reads/cell required to achieve this depth can vary as it is dependent on the gene expression levels, number of cells, and sequencing run quality. Below are the recommended reads/cell for WTA mRNA, Sample Tag, AbSeq, TCR, and BCR libraries.

# Read requirements for libraries

Library	Read requirement for data analysis		
WTA mRNA	~10,000-100,000 reads/cell		
Sample Tag: samples from the same type of cell (combining different donor PBMCs)	120 reads/cell		
Sample Tag: samples from different types of cells (combining cell lines with PBMCs)	600 reads/cell		
AbSeq	1,000 reads/cell/AbSeq*		
TCR	~5,000 reads/T cell		
BCR	~5,000 reads/B cell		
* The amount of sequencing needed for BD® AbSeq libraries will vary depending on application, BD® AbSeq panel plexy, and cell type. We have observed that using 40,000 sequencing reads per cell for 40-plex BD® AbSeq libraries prepared from resting PBMCs			

achieves an RSEC sequencing depth of ~2.

Sequencing depth can vary depending on whether the sample contains high- or low-content RNA cells. For resting PBMCs, we recommend:

- 10,000 reads per cell for shallow sequencing. Genes per cell and UMI per cell detected is generally lower but can be useful for cell type identification.
- 50,000 reads per cell for moderate sequencing.
- 100,000 reads per cell for highly saturated deep sequencing to identify the majority of UMIs in the library.

# Pooling libraries for sequencing

The efficiency of sequencing on Illumina instruments is influenced by many conditions, library size being one of them. The TCR and BCR libraries are ~200 – 300 bp larger than the WTA mRNA library and ~600 bp larger than the AbSeq and Sample Tag libraries which will cause them to produce less sequencing data if pooled in a 1:1 ratio with the other libraries. To overcome the difference in sequencing efficiency, more DNA of the TCR and BCR libraries needs to be included in the pool than would be expected when calculating ratios based on read depth. The following tables show examples of different pooling strategies and the expected sequencing outcome, with and without correction for the size of the TCR and BCR libraries. Validation data indicates a 5x

volume correction factor is needed for sequencing TCR and BCR libraries with WTA mRNA, AbSeq, and Sample Tag libraries.

**Important note:** AbSeq and Sample Tag libraries can be sequenced together or separately from WTA mRNA and TCR/BCR libraries. For optimal clustering of TCR/BCR libraries on Illumina platforms, however, we recommend sequencing AbSeq libraries separately. Recommendations for sequencing AbSeq and Sample Tag libraries alone or in a pool with other libraries are in the next section.

# Example of pooling with no correction

In this example, a total of 5,000 enriched T cells were processed as a combination of two samples (two sample tags) with a 10-plex AbSeq panel. These calculations assume the TCR library, and BCR library if included, sequences at 1/5 the efficiency of the WTA mRNA, AbSeq, and Sample Tag libraries, supported by internal testing.

A	В	С	D	E	F	G
Library type	Number of cells	Expected reads/cell	Reads needed	-	Sequencing results	Sequencing results (reads/cell)
WTA mRNA	5,000	25,000	125,000,000	62.3%	139,236,902	27,847
TCR	5,000	5,000	25,000,000	12.5%	5,000,000	1,000
AbSeq	5,000	10,000	50,000,000	24.9%	55,694,761	11,139
Sample Tag	5,000	120	600,000	0.3%	668,337	134
		Total	200,600,000	100%	200,600,000	

After sequencing, the reads/cell for the TCR library (Column G) does not match with and are much lower than the expectation (Column C), because it does not sequence as efficiently as the WTA mRNA, AbSeq, nor Sample Tag library. The remaining reads are allotted to the WTA mRNA, AbSeq, and Sample Tag libraries resulting in more reads than required. To obtain the desired number of reads/cell for each library, a correction factor is required for pooling calculations.

# Example of pooling with correction

In this example, the same sample as in the previous one was pooled using a correction factor of 5 for the TCR library to overcome the differences in sequencing efficiency. The amount of data needing to be generated (Column D) is based on the cell number (Column B) and required number of reads per cell (Column C). Based on this example, 200.6 million reads are needed to achieve the appropriate read depths. Changing the pooling ratios by correcting for the lower TCR sequencing efficiency will help ensure the correct amount of data is generated for each library. This modified pooling scheme, however, does not change the total amount of data needing to be generated, 200.6 million reads.

A	В	С	D	E	F	G	Н	I	J
Library type	Number of cells	Expected reads/cell	Reads needed	Pooling ratio before correction	Correction	Reads needed for pooling	Pooling ratio with correction	Sequencing results	Sequencing results (reads/cell)
WTA mRNA	5,000	25,000	125,000,000	62.3%	n/a	125,000,000	41.6%	125,000,000	25,000
TCR	5,000	5,000	25,000,000	12.5%	5*	125,000,000	41.6%	25,000,000	5,000
AbSeq	5,000	10,000	50,000,000	24.9%	n/a	50,000,000	16.6%	50,000,000	10,000
Sample Tag	5,000	120	600,000	0.3%	n/a	600,000	0.2%	600,000	120
		Total	200,600,000**	100%		300,600,000+	100%	200,600,000	

<sup>\*</sup>The 5x correction factor is a recommended starting point and some fine tuning may be required to achieve the optimal library balance.

After sequencing, the total amount of data generated (Column I) as well as the reads/cell for each library (Column J) are as expected (Columns D and C). The correction for library pooling did not change the amount of data generated (200.6 million reads) but helped ensure the data was spread out appropriately to each library.

# Example of pooling with a mixed population

The table below shows the pooling logic for two samples of purified B and T cells, each stained with a different sample tag and 10-plex AbSeq panel.

Α	В	С	D	E	F	G	Н	I	J
Library type	Number of cells		Reads needed	Pooling ratio before correction	Correction	Reads needed for pooling	Pooling ratio with correction	Sequencing results	Sequencing results (reads/cell)
WTA mRNA	10,000	25,000	250,000,000	62%	n/a	250,000,000	41%	250,000,000	25,000
TCR	7,000	5,000	35,000,000	9%	5*	175,000,000	29%	35,000,000	5,000
BCR	3,000	5,000	15,000,000	3%	5*	75,000,000	12%	15,000,000	5,000
AbSeq	10,000	10,000	100,000,000	25%	n/a	100,000,000	17%	100,000,000	10,000
Sample Tag	10,000	600	6,000,000	1%	n/a	6,000,000	1%	6,000,000	600
	Total 406,000,000					606,000,000+	100%	406,000,000	

 $<sup>^*</sup>$ The 5x correction factor is a recommended starting point and some fine tuning may be required to achieve the optimal library balance.

 $<sup>^{\</sup>star\star}\text{Total}$  amount of data to be requested from the sequencing facility plus PhiX.

<sup>†</sup>Read total only for pooling purposes.

<sup>\*\*</sup>Total amount of data to be requested from the sequencing facility plus 3% PhiX.

<sup>†</sup>Read total only for pooling purposes.

# Example of pooling with correction while sequencing AbSeq and Sample Tag libraries separately

The table below shows the pooling logic for the previous example of a mixed population where the AbSeq and Sample Tag libraries are sequenced together and the WTA mRNA, TCR, and BCR libraries are sequenced together.

## Pooling for AbSeq and Sample Tag libraries

Α	В	С	D	E	F	G	
Library type	Number of cells	Expected reads/cell	Reads needed	Pooling ratio		Sequencing results (reads/cell)	
AbSeq	10,000	10,000	100,000,000	94%	100,000,000	10,000	
Sample Tag	10,000	600	6,000,000	6%	6,000,000	600	
Total 106,000,000* 100% 106,000,000							
*Total amou	*Total amount of data to be requested from the sequencing facility plus 15% PhiX.						

### Pooling for WTA mRNA, TCR, and BCR libraries

Α	В	С	D	E	F	G	Н	I	J
Library type		Expected reads/cell	Reads needed	Pooling ratio before correction	Correction	needed for	Pooling ratio with correction	Sequencing results	Sequencing results (reads/cell)
WTA mRNA	10,000	25,000	250,000,000	83%	n/a	250,000,000	50%	250,000,000	25,000
TCR	7,000	5,000	35,000,000	12%	5*	175,000,000	35%	35,000,000	5,000
BCR	3,000	5,000	15,000,000	5%	5*	75,000,000	15%	15,000,000	5,000
		Total	300,000,000**	100%		500,000,000+	100%	300,000,000	

<sup>\*</sup>The 5x correction factor is a recommended starting point and some fine tuning may be required to achieve the optimal library balance. \*\*Total amount of data to be requested from the sequencing facility plus 3% PhiX.

#### Additional considerations

- 1. The 5x volume correction factor is a recommended starting place for pooling these libraries. This may need to be adjusted to accommodate different types of flow cells (for example, patterned vs non-patterned).
- 2. It can be easier to achieve the desired sequencing depth when sequencing multiple TCR or BCR libraries alone since all the libraries are the same size. Pooling will not require a correction and will only be dependent on the number of cells and the reads/cell. This scheme, however, would require 10 15% PhiX, rather than the 3% when sequencing with the WTA mRNA library.
- 3. Sequencing AbSeq and Sample Tag libraries separately can help ensure each library receives enough data during sequencing due to size discrepancies but will require 15% PhiX. However, these libraries can be successfully sequenced in combination with WTA mRNA, TCR, and BCR libraries.
- 4. All libraries derived from the same cartridge can be indexed with the same Illumina indices or reverse index primer from the BD Rhapsody™ reagents. The primary analysis pipeline can differentiate the library types

<sup>†</sup>Read total only for pooling purposes.

(for example, WTA mRNA vs TCR) based on their structure and sequences. Demultiplexing statistics are reported from the pipeline, but should these statistics be desired prior to running the pipeline, then unique indices will be required for each library.

For additional support with pooling and sequencing, please reach out to your local Field Application Specialist (FAS) or scomix@bdscomix.bd.com.

# Sequencing flow cell loading and PhiX concentrations

# **Quantifying libraries**

Calculate the molar concentration of WTA, AbSeq, and TCR/BCR libraries using Qubit quantitation concentration (ng/ $\mu$ L) and average Bioanalyzer size (200 bp - 1000 bp). For TCR/BCR libraries, the expected Qubit concentration should be >1.5 ng/ $\mu$ L. Use the calculated molar concentrations to pool libraries.

# AbSeq/Sample Tag libraries

For a NextSeq High or Mid Output and MiniSeq High or Mid Output runs, load the flow cell at a concentration between 1.8-2.2 pM with 15% PhiX for a sequencing run.

# **Required parameters**

Parameter	Requirement
Platform	Illumina: 150 cycle kit
Paired-end reads	Minimum of 51 x 75 paired read length
PhiX	Required (15%)
Analysis	See the BD <sup>®</sup> Single-Cell Multiomics Bioinformatics Handbook (23-21713)

# WTA and TCR/BCR libraries (with or without AbSeq/Sample Tag)

For a NextSeq High or Mid Output and MiniSeq High or Mid Output runs, load the flow cell at a concentration between 1.4-1.8 pM with 3% PhiX\*. For other sequencers follow Illumina recommendations for loading concentration and use 3% PhiX.

**Note:** \*if using less than 10,000 reads/cell for the WTA library increase PhiX percentage to 5-10% to account for lower library diversity.

Set up sequencing run on Illumina<sup>®</sup> BaseSpace. Enter the pooled libraries as one sample if libraries were made with the same Library Forward primer but with different i7 indices.

# Required parameters

Parameter	Requirement		
Platform	Illumina: 300 cycle kit		
Paired-end reads	Minimum of 85 x 215* paired read length		
PhiX	Required (3%)		
Analysis See the BD® Single-Cell Multiomics Bioinformatics Handbook (23-21713)			
*R2 length of 215 is recommended for optimal assembly. If necessary 150 x 150 read lengths can be used.			

# Sequencing analysis pipeline

Contact customer support at <a href="mailto:scomix.bd.com">scomix.bd.com</a> for access to the latest whole transcriptome sequencing analysis pipeline.

# **Appendix**

# Illumina Index 1 (i7) sequences

Library Reverse Primer	Sequence
1	GCTACGCT
2	CGAGGCTG
3	AAGAGGCA
4	GTAGAGGA

# Human T cell PCR1 primers

Primer name	Primer sequence (5' - 3')
TRAC_N1	CTGGAATAATGCTGTTGTTGAAGG
TRBC_N1	AGCCCGTAGAACTGGACTT
TRDC_N1	CTTCAAAGTCAGTGGAGTGCA
TRGC_N1	CACCGTTAACCAGCTAAATTTCATG

# Human T cell PCR2 primers

Primer name	Primer sequence (5' - 3')
TRAC_N2	ATCAAAATCGGTGAATAGGCAGAC
TRBC_N2	GATCTCTGCTTCTGATGGCTCA
TRDC_N2	ATATCCTTGGGGTAGAATTCCTTC
TRGC_N2	GGGAAACATCTGCATCAAGTTG

# Human B cell PCR1 primers

Primer name	Primer sequence (5' - 3')
IGHA_N1	CACAGTCACATCCTGGCT
IGHD_N1	GATCTCCTTCTTACTCTTGCTGG
IGHE_N1	CGCTGAAGGTTTTGTTGTCG
IGHG_N1	TGTTGCTGGGCTTGTGAT
IGHM_N1	сбттстттстттбттбссбт
IGKC_N1	TTTGTGTTTCTCGTAGTCTGCT
IGLC_N1	TGTAGCTTCTGTGGGACTTC

# Human B cell PCR2 primers

Primer name	Primer sequence (5' - 3')
IGHA_N2	CTTTCGCTCCAGGTCACACT
IGHD_N2	TGTCTGCACCCTGATATGATGG
IGHE_N2	GTCAAGGGGAAGACGGATG
IGHG_N2	AAGTAGTCCTTGACCAGGCA
IGHM_N2	ACAGGAGACGAGGGGAAAA
IGKC_N2	TCAGATGGCGGGAAGATGAA
IGLC_N2	ACCAGTGTGGCCTTGTTG

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