



purePlex™ DNA Library Preparation Kit  
for Illumina® Sequencing Platforms  
(Product Nos. 301067, 301068, 301069, and 301070)

User Guide

V20230327

## Introduction

The purePlex™ DNA Library Preparation Kit enables the streamlined and parallel construction of up to 96 uniquely dual-indexed, sequencing-ready libraries per plate. The use of unique dual indexes (UDIs), where unique i5 and i7 indexes are used to specifically identify each library within a set, allows for the detection and removal of “hopped” reads on patterned flow cells.<sup>1</sup> Additionally, the workflow tolerates a wide DNA input concentration range without the need for protocol modifications and utilizes a novel DNA normalization reagent wherein individual libraries can be pooled and amplified together to reduce final QC burden. Overall, this translates to a reduction in labor and consumable costs. Each purePlex kit allows for flexible batch and pool sizes (see Appendix D) and contains sufficient reagents to process and pool 96 samples.

This multiplexed library prep procedure is optimized for inputs of 5 - 50 ng of purified dsDNA per sample, and typically generates library fragment lengths ranging from 300 – 1,500 bp. Multiple DNA types are suitable input for the kit, ranging in size and complexity from amplicons to genomic DNA. purePlex libraries are compatible with all Illumina sequencers including iSeq, MiniSeq, MiSeq, NextSeq, HiSeq and NovaSeq systems.<sup>2</sup>

Library preparation kits from seqWell utilize proprietary transposase-based reagents to insert barcoded adapters directly into input DNA in two separate steps. In the purePlex workflow, a first barcoding step adds unique i7-barcoded adapters into up to 96 DNA samples in segregated reactions. During the second step, unique i5-barcoded transposases and normalization reagent are added to each reaction. At the end of the second step each well contains a normalized number of unique dual indexed library fragments. After a brief stop reaction, samples are pooled in batches of 24,<sup>3</sup> purified, and amplified in a single PCR reaction using universal library primers (i.e., P5 and P7 primers), making for a highly efficient multiplexed library prep workflow (see Workflow Diagram).

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<sup>1</sup> <https://www.illumina.com/techniques/sequencing/ngs-library-prep/multiplexing/index-hopping.html>

<sup>2</sup> Refer to Illumina technical documentation for specific instructions on how to set up a sequencing run for a Nextera style 10 nt dual-indexed library prior to loading a plexWell library on your model of sequencing system.

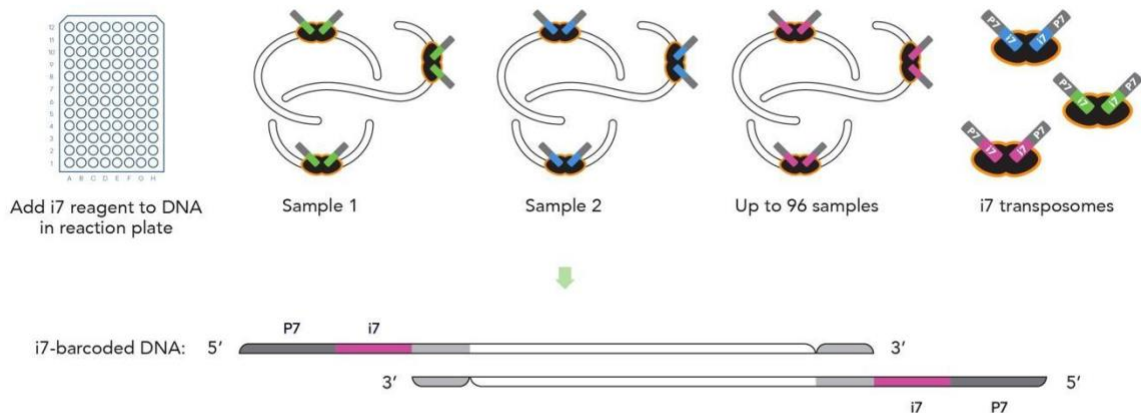
<sup>3</sup> Refer to Appendix D for pooling in batches not equal to 24

## Table of Contents

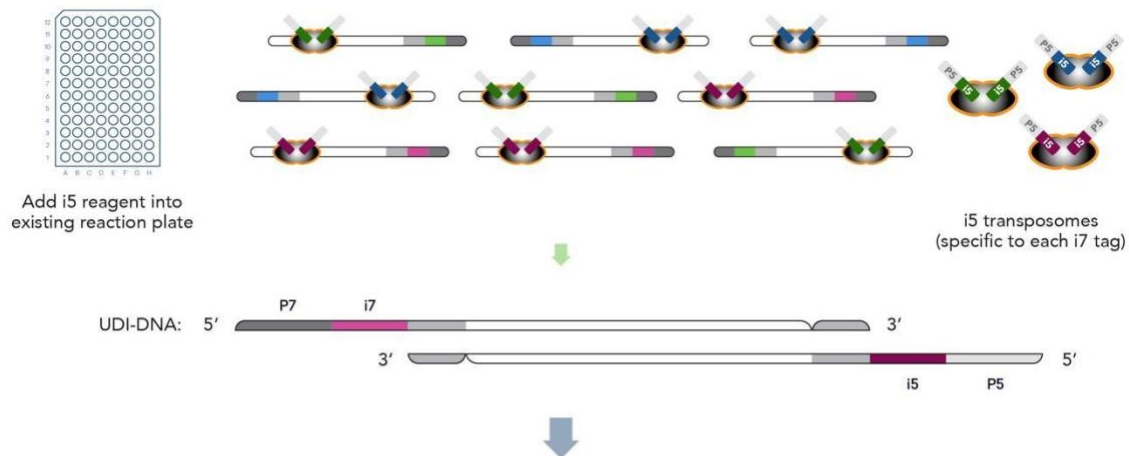
<a href="#">Molecular Diagram</a>	4
<a href="#">Workflow Diagram</a>	5
<a href="#">plexWell Kit Components</a>	6
Considerations before you begin	
<a href="#">Required Equipment, consumables, and reagents</a>	7
<a href="#">Reagent handling</a>	8
<a href="#">purePlex DNA Library Prep detailed protocol</a>	9
<a href="#">i7 Tagging (i7-TR) Reaction</a>	9
<a href="#">i5 Tagging (i5-TR) Reaction</a>	10
<a href="#">Tagging Reaction Stop</a>	10
<a href="#">Tagged Library Pooling</a>	11
<a href="#">Tagged Library Pool Purification</a>	11
<a href="#">Library Amplification</a>	12
<a href="#">Library Purification</a>	13
<a href="#">Library QC</a>	15
<a href="#">Sequencing Loading and Read Configuration</a>	16
Appendices	
A: <a href="#">purePlex™ Index Information</a>	17
B: <a href="#">purePlex™ Library Intermediates</a>	20
C: <a href="#">Sample Sheet and Sequencer Loading Guidelines</a>	21
D: <a href="#">Sample Pooling for Plexities ≠24</a>	22

# purePlex™ Library Prep - Molecular Diagram

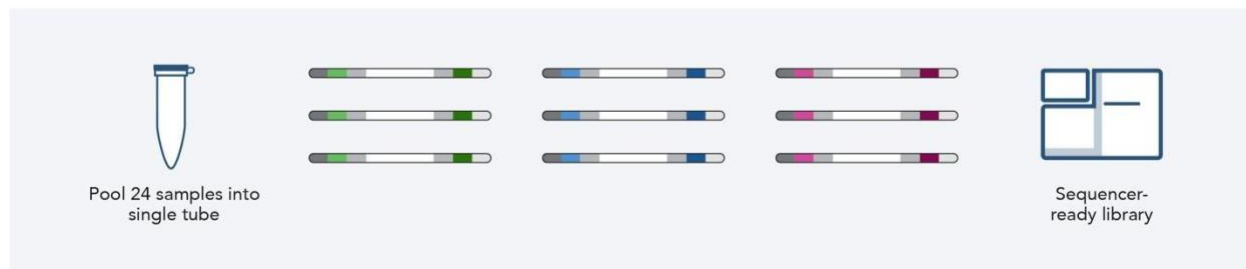
## i7 BARCODING:



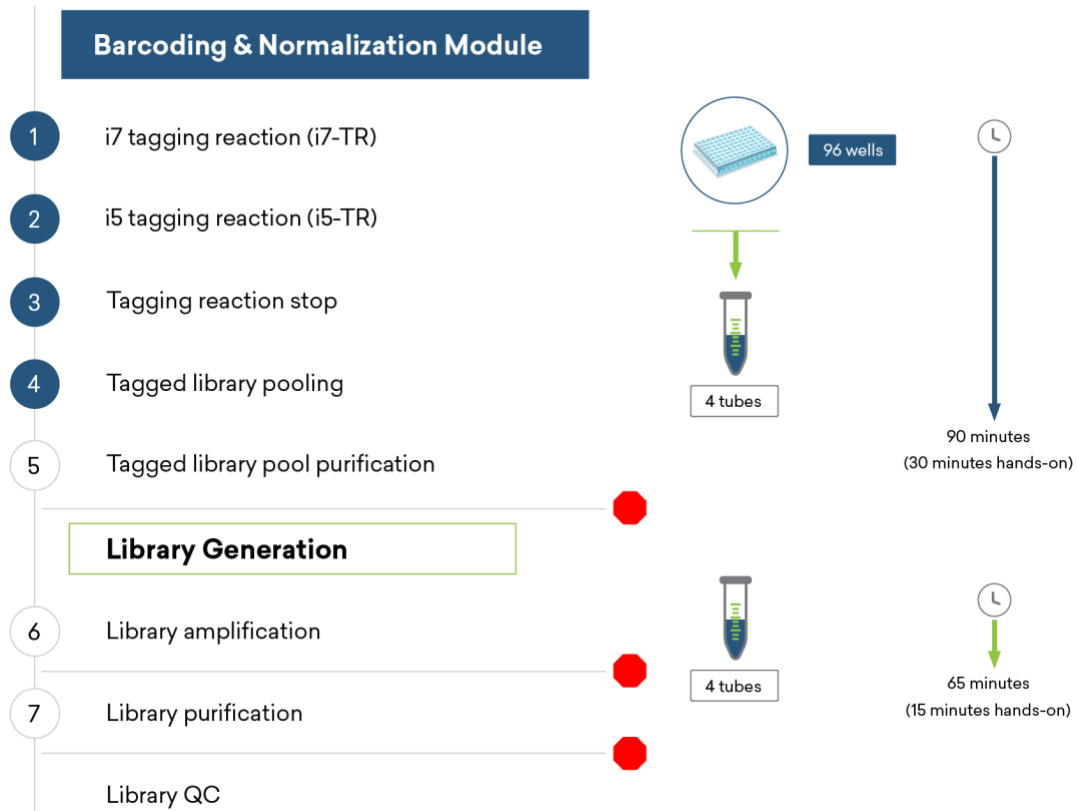
## i5 BARCODING:



## FILL-IN AND LIBRARY AMPLIFICATION:



# purePlex™ Library Prep - Workflow Diagram



**Table 1.** Components of purePlex™ Library Preparation Kit

We currently offer four sets of 96 UDIs: Set 1 (Part No. 301067), Set 2 (Part No. 301068), Set 3 (Part No. 301069), and Set 4 (Part No. 301070). Part numbers for the Tagging Reagent plates are different between the four sets, but all other components are identical (see below).

Important note: Tagging Reagent sets are NOT interchangeable. Please only use the i7-TR96 plate and i5-TR96 plate corresponding to the same set. Do not mix plates from different sets.

Box	Component	P/N	Description	Storage	Qty
1	<b>i7 Tagging Reagent Plate</b>	i7-TR96-X	Fully skirted, low profile, 96-well, <b>blue</b> PCR plate.*	-20°C	1
1	<b>i5 Tagging Reagent Plate</b>	i5-TR96-X	Fully skirted, low profile, 96-well, <b>yellow</b> PCR plate.*	-20°C	1
2	<b>Coding Buffer (3X)</b>	CB0384	2 ml tube, violet cap, 1.5 ml	ambient	1
	<b>X Solution</b>	QB0096	2 ml tube, black cap, 1.5 ml	ambient	1
	<b>MAGwise™ Paramagnetic Beads</b>	MG5000	8 ml bottle, 5 ml	4°C	1
3	<b>Normalization Reagent</b>	NRM60	2 ml tube, yellow cap, 600 µl	-20°C	1
	<b>Library Primer Mix</b>	PRM384	0.5 ml tube, natural cap, 24 µl	-20°C	1

\*i7 and i5 tagging reagent plates may be supplied as colored plates as denoted in the table above OR by blue and yellow (i7 and i5, respectively) stickers located on the plate skirt. For best results, read the plate label to ensure the correct reagents are used at each step.

# User-Supplied Reagents, Equipment & Consumables, and Thermal Cycler Programs

## Reagents

- 80% Ethanol (freshly prepared)
- Tris-HCl, pH 8.0
- PicoGreen® DNA assay (recommended) or other validated dsDNA quantification assay
- KAPA HiFi HotStart ReadyMix (2X) (P/N: KK2602 or KK2601) for library amplification

## Equipment & Consumables

- Single-channel pipettors (1-20 µl, 20-200 µl, 100-1,000 µl)
- Multi-channel pipettors (1-10 µl, 10-200 µl)
- Pipette tips (low-retention barrier tips)
- Eppendorf Tubes® 1.5 ml, DNA LoBind Tubes
- PCR plate seals (must be evaporation-resistant)
- Thermal cycler
- Magnetic stand for 1.5 ml and 2 ml tubes
- 0.2 ml PCR 8-tube strips and caps/seals
- Benchtop centrifuge to pulse-spin tubes and PCR 8-tube strips
- Plate centrifuge
- Vortex mixer

## Thermal Cycler Programs (all with lid-heating on)

- **TAG:** 55°C for 15 minutes; 25°C hold.
- **STOP:** 68°C for 10 min; 25°C hold.
- **FILL AMP8:**
  - 72°C for 10 minutes (fill-in)
  - 95°C for 3 minutes (initial denaturation)
  - 
  - 98°C for 30 seconds
  - 64°C for 15 seconds
  - 72°C for 30 seconds
  - 
  - 72°C for 3 minutes (final extension)
  - 4°C hold

} 8 cycles

## Before starting the procedure:

**Adjust input DNA concentration.** Assay the DNA concentration of each sample using a PicoGreen assay. This protocol tolerates a range of 5 - 50 ng of purified genomic DNA input per sample. Before starting library preparation, adjust each input DNA concentration to be 1.0 – 10.0 ng/μl in 10 mM Tris-HCl, pH 8.0 (do not dilute input DNA with TE, or other EDTA-containing buffers).

**Program thermal cycler(s).** For convenience, set-up the thermal cycler programs listed on the previous page before starting.

**Pulse-spin kit components.** Liquids can condense and shift location inside containers during shipment or storage. Before opening the **i7 and i5 TR plates** and before dispensing from reagent tubes, pulse-spin in a suitable centrifuge to gather the reagents at the bottom of the well or tube. If kit components freeze, thaw, mix and pulse-spin before use.

**Equilibrate MAGwise paramagnetic beads to room temperature.** MAGwise beads can be stored for up to 2 weeks at room temperature, or, for longer periods at 2° - 8°C. Equilibrate to room temperature for 30 minutes before use. Vortex to thoroughly resuspend the magnetic beads prior to use. Pipette slowly and do not pre-wet pipette tips to transfer volumes accurately.

**Check the X Solution for precipitate before use.** If a precipitate is visible, incubate at 37°C for 5 minutes (or longer). Mix gently by inversion until the precipitate dissolves (do not vortex). Note: X Solution contains SDS and will precipitate if stored below room temperature. Overly vigorous mixing will cause foaming.

**Please note that Coding Buffer is viscous.** Store **Coding Buffer (3X)** at room temperature. To transfer volumes accurately, pipette slowly and do not pre-wet pipette. While adding **Coding Buffer** to reactions, mix in completely by pipetting up and down several times with the same pipette tip(s) used for addition. Always change pipette tips before adding **Coding Buffer** to different reactions.

**Prepare 80% ethanol fresh daily.**

**Prepare 10 mM Tris-HCl, pH 8.0.** Prepare 10 mM Tris-HCl, pH 8.0 from a concentrated stock solution diluted with ultrapure water (both molecular-biology/PCR grade). Do not use EDTA-containing solutions (e.g., TE) to dilute input DNA or elute DNA from beads as EDTA can inhibit enzymatic activity.

**Safe-stopping points are indicated in the procedure.** For optimal results, proceed directly to the next step unless an optional safe-stopping point is indicated.



## Procedure

Before beginning the procedure, review the adjusting sample concentration guidelines on the previous page to ensure that each sample has been adjusted to 1.0 - 10.0 ng/μl using 10 mM Tris-HCl, pH 8.0.

The protocol herein has been validated for pools of 24 samples, such that columns 1-3, 4-6, 7-9, and 10-12 each have their own pool. If processing <96 samples at one time, take careful note of storage conditions especially for i7 and i5-TR plates. Appendix D includes tips for alternate pooling in plex sizes other than 24.

### 1. i7 Tagging (i7-TR) Reaction

- a. Transfer 5 μl of DNA sample (1.0-10 ng/μl) to a 96-well PCR plate (the REACTION PLATE).
- b. Pulse-spin the **i7-TR plate** in a centrifuge. Take care when removing the seal to avoid splashing/contamination of reagents. Only remove the seal from the wells that will be used.

**Useful Tip:** *If processing <96 samples, use a razor blade to cut the plate seal on the i7-TR plate around the wells that will be used. Then remove only that portion of the plate seal.*

- c. Next, add 5 μl from the i7 tagging reagent to the DNA samples. Carefully pipette 5 μl from the i7-TR plate and transfer to the REACTION PLATE already containing the DNA. Mix thoroughly and slowly by pipetting up and down (5 times at 5 μl), being careful not to introduce excessive bubbles. Take note of which i7-TR reagent is used for each sample.

**Note:** If <96 samples are being processed, reseal and store the **i7-TR plate** at -20°C, marking on the seal what columns have been used.

- d. Next, carefully pipette 5 μl of **Coding Buffer (3X)** to each well of the REACTION PLATE, using new tips for each transfer. Mix thoroughly and slowly by pipetting up and down (10 times at 5 μl), being careful not to introduce excessive bubbles.

**Useful Tip:** *Aliquot 70 μl of **Coding Buffer (3X)** into each well of a PCR 8-tube strip, and then use a multichannel pipettor to dispense 5 μl into the REACTION PLATE. The coding buffer can be stored in the PCR strip at room temperature for up to 4 weeks for future use.*

- e. Seal the REACTION PLATE, pulse-spin, then transfer to a thermal cycler, and run the TAG program, below, with lid-heating on:

55°C for 15 minutes

25°C hold

## 2. i5 Tagging (i5-TR) Reaction

- a. Pulse-spin the REACTION PLATE in a centrifuge and carefully remove the seal.
- b. Add 5  $\mu$ l of **Normalization Reagent** to each well being used in the REACTION PLATE using new tips for each transfer. Mix thoroughly and slowly by pipetting up and down (10 times at 5  $\mu$ l), being careful not to introduce excessive bubbles.

**Useful Tip:** Aliquot 70  $\mu$ l of **Normalization Reagent** into each well of a PCR 8-tube strip, and then use a multichannel pipettor to dispense 5  $\mu$ l into the REACTION PLATE. If processing < 96 samples, store the remaining **Normalization Reagent** in the strip tube at -20°C for up to 4 weeks.

- c. Pulse-spin the i5-TR plate in a centrifuge. Take care when removing the seal to avoid splashing/contamination of reagents. Only remove the seal from the wells that will be used.

**Useful Tip:** If processing <96 samples, use a razor blade to cut the plate seal on the i5-TR plate around the wells that will be used. Then remove only that portion of the plate seal.

- d. Transfer 5  $\mu$ l from the i5-TR plate to the REACTION PLATE well that used the i7-TR from the same well. Use new tips for each transfer. Mix thoroughly and slowly by pipetting up and down (10 times at 5  $\mu$ l), being careful not to introduce excessive bubbles.

**Note:** The unique dual index i7+i5 combinations have been specifically paired, thus the i7-TR in A01 should ONLY be used with the i5-TR in A01.

**Note:** If less than 96 samples are being processed, reseal and store the i5-TR plate at -20°C, marking on the seal which columns have been used.

- e. Seal the REACTION PLATE, pulse-spin, then transfer to a thermal cycler, and run the TAG program, below, with lid-heating on:

55°C for 15 minutes

25°C hold

## 3. Tagging Reaction Stop

- a. Add 12.5  $\mu$ l of **X Solution** to each well of the REACTION PLATE. Pipette up and down slowly 10 times to mix, being careful not to introduce excessive bubbles. Use clean tips for each addition of **X Solution**.

**Useful Tip:** Aliquot 160  $\mu$ l of **X Solution** into a PCR 8-tube strip, and then use a multichannel pipettor to transfer 12.5  $\mu$ l from the strip to the REACTION PLATE and mix. The **X Solution** can be stored in the PCR strip at room temperature for up to 4 weeks for future use.

- b. Securely reseal and pulse-spin the REACTION PLATE, then transfer to a thermal cycler and run the STOP program, below, with lid heating on:  
68°C for 10 minutes  
25°C hold

#### 4. Tagged Library Pooling

- a. Pulse-spin the REACTION PLATE and then remove the seal.
- b. For every 24 samples being processed, set out and label a 1.5 ml LoBind tube. Label tubes as A-D if processing 96 samples. for each pool of 24 samples as A-D.

**Note:** See Appendix D for recommendations for plex sizes other than 24-plexes.

- c. Transfer 32 µl of stopped tagging reaction from every well of Cols 1-3 into tube A. Repeat the process for the samples in columns 4-6 to tube B, columns 7-9 to tube C, and columns 10-12 to tube D, such that you have 24 samples per LoBind tube.

**Useful Tip:** Pool down each column into a row of a deep well or midi plate using a multi-channel pipette then transfer to a LoBind tube.

- d. The total volume of the pool in each tube will be approximately 768 µl.

#### 5. Tagged Library Pool Purification

- a. Vortex (or vigorously pipette) room temperature **MAGwise Paramagnetic Beads** to ensure that the beads are fully resuspended.
- b. Add 576 µl (0.75 volume equivalent) of **MAGwise** to each Library Pool and mix thoroughly by pipetting.
- c. Incubate in a tube rack for ≥5 minutes to allow the DNA to bind.
- d. Place tube(s) on magnetic stand and let beads settle, 5 minutes. A pellet should form on one side of the tube and the supernatant should be visibly clear after 5 minutes.
- e. Remove and discard supernatant with pipette. Be careful not to disturb the pellet.  
**Useful tip:** Use a large pipette tip to remove most of the supernatant and then if necessary, use a smaller one to remove the remaining supernatant.
- f. Wash the bead pellet with 80% ethanol.

- i. With the tube in the magnetic stand and without disturbing the beads, add 1.5 ml of 80% ethanol. The beads should be completely covered by 80% ethanol.
- ii. After 30 seconds, slowly remove and discard supernatant, being careful not to disturb the pellet.

- g. Wash beads with 80% ethanol a 2<sup>nd</sup> time.
  - i. With the tube in the magnetic stand and without disturbing the beads, add 1.5 ml of 80% ethanol. The beads should be completely covered by 80% ethanol.
  - ii. Perform the next steps quickly, working 1-2 tubes at a time
    - 1. After  $\geq 30$  seconds, remove and discard supernatant, being careful not to disturb the pellet.
    - 2. Cap tube, pulse-spin and return to magnet, letting beads settle ( $< 30$  seconds). Use a small pipet tip ( $\leq 20$   $\mu$ l) to remove any residual ethanol at the bottom of the tube.
    - 3. Add 48  $\mu$ l of 10 mM Tris to bead pellet, remove from magnetic stand and pipette the solution along the inner wall of the tube multiple times to thoroughly resuspend the bead pellet. Place tube in rack on bench. **DO NOT air-dry bead pellet** prior to Tris addition or the DNA recovery will be compromised.
- h. Incubate the tube(s) on the bench for at least 5 minutes to elute the purified tagged reaction pool from the beads.
- i. Return tube to magnetic stand and allow a bead pellet to reform on the inner wall of the tube (~ 2 minutes).
- j. When the supernatant has cleared completely, carefully transfer 46  $\mu$ l of DNA eluate from each purification tube to a PCR tube (1 for each 24 plex). The transferred eluate contains the purified library pool.

#### SAFE STOPPING POINT

Proceed immediately to the next step, or store the purified pool at -20°C.

## 6. Library Amplification

- a. Add 4  $\mu$ l of **Library Primer Mix** to each PCR tube containing 46  $\mu$ l of the purified 24-plex pools.
- b. Add 50  $\mu$ l of Kapa HiFi Hot Start ReadyMix (2X) to each and mix well by pipetting. Total PCR reaction volume is 100  $\mu$ l.

- c. Close the PCR tubes, pulse-spin and run the FILL\_AMP8 program, below, with lid heating on:

Fill-in: 72°C for 10 min  
Initial denaturation: 95°C for 3 min  
8 Cycles of: 98°C for 30 seconds  
64°C for 15 seconds  
72°C for 30 seconds  
1 Cycle of: 72°C for 3 minutes  
4°C hold

#### SAFE STOPPING POINT

Proceed immediately to the next step, or freeze the amplified library at -20°C.

## 7. Library Purification

Following library amplification, it is necessary to remove residual primers and short library fragments by MAGwise purification.

- After PCR, pulse-spin the strip tubes containing the library amplification reactions.
- Add 25 µl of 10 mM Tris to each reaction to bring the total volume up to 125 µl.
- Transfer 120 µl of each diluted reaction to its own labeled 1.5 ml LoBind Tube.

*Retain the residual ~5 µL of the unpurified diluted library amplification reactions for TapeStation QC.*

- Vortex (or vigorously pipette) room temperature **MAGwise Paramagnetic Beads** to ensure they are completely resuspended.
- Add 90 µl (0.75x volume equivalent) of MAGwise beads to each 1.5 ml tube containing the amplified library pools. Mix thoroughly by pipetting up and down.

**Useful Tip:** *Aspirate beads slowly then wipe outside of pipet tip with a kimwipe to ensure accurate transfer of bead volume.*

- Incubate in a tube rack on the bench for 5 minutes to allow the DNA to bind.
- Transfer the 1.5 ml tube to a magnetic stand and let the beads settle completely. A bead pellet will form along one side of the tube and the supernatant should appear completely clear after 3 minutes.

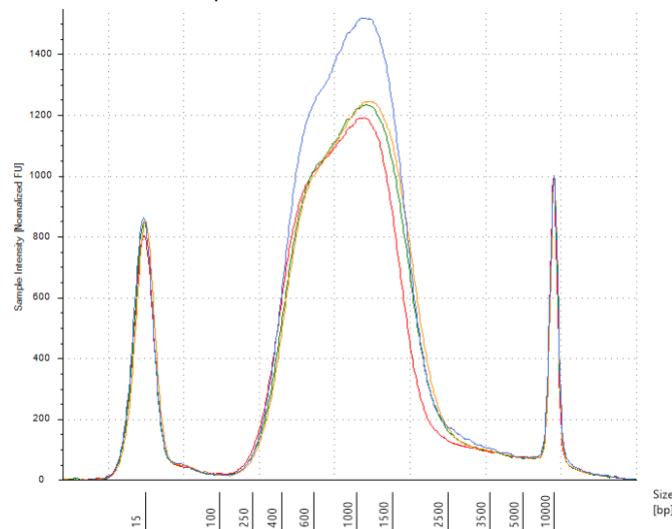
- h. Slowly remove the supernatant with a pipettor and discard. Be careful not to disturb the bead pellet. Use a large pipettor to remove most of the supernatant and then use a smaller pipettor (e.g., P20) to remove the residual supernatant.
- i. Wash beads with 80% ethanol.
  - i. With tube in the magnetic stand, add 300 µl of 80% ethanol without disturbing beads.
  - ii. After ≥30 seconds, remove and discard supernatant, being careful not to disturb the pellet.
- j. Wash beads with 80% ethanol a 2<sup>nd</sup> time.
  - i. With tube in the magnetic stand, add 300 µl of 80% ethanol without disturbing beads.
  - ii. Perform the next steps quickly, working 1-2 tubes at a time
    1. After ≥30 seconds, remove and discard supernatant, being careful not to disturb the pellet.
    2. Cap tube, pulse-spin and return to magnet, letting beads settle (<30 seconds). Use a small pipet tip (≤20 µl) to remove any residual ethanol at the bottom of the tube.
    3. Add 32 µl of 10 mM Tris to bead pellet, remove from magnetic stand and pipette the solution along the inner wall of the tube multiple times to thoroughly resuspend the bead pellet. Place tube in rack on bench. **DO NOT air-dry bead pellet** prior to Tris addition or the DNA recovery will be compromised.
- k. Incubate for 5 minutes on the bench to elute the purified library from the magnetic beads.
- l. Return tubes to magnetic stand and allow bead pellet to form on the inner wall of the tube (~ 2 minutes).
- m. When the supernatant has cleared completely, carefully transfer 30 µl of DNA eluate to a new 1.5 ml LoBind tube. The transferred eluate contains the purified, size-selected multiplexed library.

#### SAFE STOPPING POINT

Store the purified, multiplexed library at -20°C, or proceed directly to library QC.

## Library QC

**Electrophoretic analysis:** Run an aliquot of purified library through electrophoretic analysis such as the Agilent Bioanalyzer (High Sensitivity DNA or DNA7500 kits), Tapestation (High Sensitivity D5000 or D5000 kits), or Fragment Analyzer (High Sensitivity NGS Fragment Analysis Kit, DNF-474). The majority of library fragments should be between 300 and 1,500 bp in length. Typical results on the Tapestation High Sensitivity D5000 assay shown in Figure 1, below. Follow the manufacturer's instructions for these instruments and dilute the library, if appropriate, prior to running. Typically purePlex™ libraries are 80-100 nM. To determine the average fragment size, use a region analysis for 200-1500 bp and use this value for size adjustment with picogreen based assays or SYBR based qPCR assays for determining library concentration.



**Figure 1.** Representative purePlex™ library traces generated using a Tapestation 2200 with HSD5000 DNA reagents and tapescreens. The library was prepared from human genomic DNA (input range of 5-50ng) and used a final purification of 0.75 volume equivalents of MAGwise. Prior to loading on a high sensitivity D5000 tapescreen, the library was diluted 1:10. Average fragment size with 0.75X cleanups is ~800 bp using a region analysis of 200-1500).

**Note:** purePlex library preparation kits sometimes produce fragments >1000 bp that appear to constitute a large portion of the library on instruments such as the Agilent Bioanalyzer, Tapestation, or Fragment Analyzer. However, these fragments do not appreciably contribute to the library concentration and do not cluster on standard Illumina flow cells. As such it is not necessary to remove them from the library.

If instruments mentioned above are unavailable, run 3 µl of purified library and 5 µl of unpurified, diluted, library on a 2% Agarose E-gel EX alongside the 1 kb plus ladder (NEB) to determine the median fragment length.

**qPCR assay:** Use 2 µl of the purified, multiplexed library for qPCR analysis. Follow kit and instrument documentation for appropriate conditions and dilutions. For KAPA Library Quantification kits, prepare a 1/100K dilution of the multiplexed library. Use the average fragment size determined during fragment analysis, above, for the size adjustment. Library concentrations are typically 80 – 100 nM.

## Sequencing Loading and Read Configuration

The purePlex™ Library Prep kit libraries are dual indexed using 10 nt indices. Index lists can be found in appendix A or in an excel document that can be requested at [support@seqwell.com](mailto:support@seqwell.com). The protocol is optimized to generate median inserts of >300 nt to fully utilize 300 cycle Illumina kits. plexWell libraries are identical in structure, as well as adapter and sequencing primer site sequences, to Nextera® libraries. Standard Illumina sequencing primers are utilized. Refer to Illumina technical documentation for specific instructions on how to denature and dilute a purified library on your model of sequencing system<sup>4</sup> and for additional information on setting up the read configuration.

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<sup>4</sup> The sequencing primers provided in TruSeq v3 Cluster kits are incompatible with Nextera-style libraries, including purePlex libraries. The TruSeq Dual Index Sequencing Primer Box from Illumina is required for sequencing plexWell libraries on older systems, such as the HiSeq 2500, HiSeq 2000, HiSeq 1500, GAIIx, and HiScanSQ.



## Appendix A: purePlex™ index information

The most up-to-date list of indices for [all seqWell kits](#) is available in the Resources section for each product. Click on the “purePlex™ Master index list” to download an excel workbook containing all i7 indices (in list and plate layout) and i5 indices. The purePlex master index list available on our website under *Resources > Product Information* is the easiest way to copy and paste index sequences into sample sheets or for demultiplexing.

### purePlex Index Set 1 (Part No. 301067)

Plate Layout - **Set 1**

i7 indices for UDI plates - i7-TR96-1

	1	2	3	4	5	6	7	8	9	10	11	12
A	GTCAGTCCA	CAACTAATC	ATAACCTGAC	CAGGTACTTC	AACCGAGCCA	CAACGTCATT	ATTGGTCAGA	ATGTTGCGGA	CACCAATAAC	TGTCCGTCTT	CGAAGGACTG	TAGTTATCGC
B	TATCTCTTC	GTACTGGATT	TGCGGTTCCA	CAGGAATATG	TGGATTCAAG	AAGAACGATG	AAGACCTGTT	AATGCTAAC	GCGTCCACAA	CATGAGTAAC	TCTACCGTCA	TGCAGGTGAT
C	CCGCGAAGAA	CATCGGAGGA	AATACTTGCC	AACGCACAAT	CAACAGATAC	CTGCAATTAC	CGCTAATGAA	GTAACACGTA	ATGTGCGCTT	CTGCGGAAT	GGTAATATCG	TGTGAAGCTA
D	CCTACTCGGA	ATTCTGATGG	AATCGCGGAA	ATTGAGAAGS	AAGSTAATC	CGCCGATGAT	CAGAGTGCAT	GATATACGGA	TATCTAGTGC	CTGTAGTATG	TGTGCGAGTT	CAACTCCTGA
E	TTCGTATCAC	TATCGTTACC	CAGAACCGGA	ATTGCACCTT	TGACAATACG	CGGACAAGAC	GGTAAGCTGA	ATCCGAGAGG	GCCTACAATG	GAGCCGTACA	TTATCGCTGA	TGTAGCAACG
F	TCCTCCATCC	TGGTCTGTTA	TGAACCAAGG	AAGTGGATAC	GATTGTGCAT	CGTACTCCTC	GTATTCAAGT	CAATTCACAC	AAGCTCAGTT	CATTCTTAGG	GGTTGAGTTC	CGGTAAACGA
G	ATAACATCGC	TGCCAACATG	TGTTAGTCAG	GATAAGATGC	GCACACCATT	TGCATGAGTT	GCGCTACGTT	TTGTCAGTTC	TACGCTTAGA	TGACCGACAA	ATTGGACGCC	GTAGCAGCAG
H	GATATGCGTT	TCATTACACG	CAGTAGGTAA	TACCTCGACA	TCGTTATTCC	AATCTGGAGC	TTGTCATAGC	ATACGCCATT	ATCCACTAGG	TGTGTAACCG	TATGTGTGTG	CTACAGCCGA

i5 indices for UDI plates - i5-TR96-1

	1	2	3	4	5	6	7	8	9	10	11	12
A	GTAACACAGA	CAAGAGCGTG	ATCTCCACGG	TTACTGCTGC	CCGAGGTTAG	CTGATTAGGA	TGGAGCGATG	TCATACGTTT	TTGTCTTCC	TCTCAACAAG	CGTGATACTT	CACACAGTCG
B	TAAGTTGTGG	AATACTCGGC	CTCGCTGCTT	TTGTTCTGCG	CCGTAATCGA	CAATGCGGAG	TTGTTGTCGA	TTGTGTCCGA	GGAGTTCTAT	TCGCATGACC	TACAATCCGG	TACCATCTAG
C	CACCTACCTC	GTATCTTAGG	CTCAGTAGAC	TCGCGGACAT	CCATTCACTT	ATTCCGTTA	CCTCTGAACA	CTTATCCAGG	ATCTTAACCG	TACGAAGCAA	TAACCATTCG	TTGTAGCATC
D	ATATCAGTCC	TGGTCTGAGG	CTTGCTCTTT	ATCGAATTGC	TCCGACTACC	CTGACAACGG	TCCACCATAG	CGGTAACCTC	GGAGCTTGTT	CCAACGCTTA	TGCTACTGTT	TTCTTCCGTA
E	GGTTAAGTAG	CAAGGACATT	CGTCCAAGAG	TGTAGGATAG	TAGAACCCTGA	GCATGTTAAG	CAAGCATGAA	TAGGACCATG	GTTACAGGTT	CAGTTAGCTC	CGTCAATTCG	TCTAGCGTTG
F	TGGTGATAT	CGGTTGAGAA	GTATGGAACA	ATGTTGCTGA	ATATGTTGCA	TCGAGAAGCG	TCAGTTGGTT	CACAATTACG	CAACCAACAT	GTATGCTAAC	TTACGTTTGC	TCTAAGTCTG
G	TAGCTGGCAC	TACCTCGCTA	AATCGGTACG	TGAGGATGAA	TTGAGTATGG	TGCAGATGGA	CCATTATACC	AACGCGAATA	ATGTTCCGTT	GTTGAGGCAC	GGCTTGAATT	CTCATATCCA
H	TTACCGTTGG	TAAGCTCGTT	CTAATGTCGA	TGCTCTCCAA	ATTCTGGACA	TACACAACCG	AAGTGCAGG	TCACCATGAT	CGAGGAACAG	AACAGAACGG	GTTCTTCTTT	TGGAGGTATA

**purePlex Index Set 2B (Part No. 301068) – Lot #20220727 only**Plate Layout - **Set 2/B**

i7 indices for UDI plates - i7-TR96-B												
	1	2	3	4	5	6	7	8	9	10	11	12
A	CACACATCTC	GATCGACGTA	TCTTCAACGG	TCTTCGCTCA	AAGTGCAGCG	TCCGGTCATA	AAGCTGACGG	TCATAGGACG	GCCTTCTTCG	TTCCAGCCAG	CTCCACTGAA	CTACTACTTG
B	TGTGCTTGAA	CTCTGGTTCC	CGTAACTAGC	AACACCATGC	ATCCTACGCC	CCGTCGCTAT	CCGTTCTACC	GGAACATGAC	AACAACCGCA	GACTCAGGTT	CATAGTCGCC	ATGAGGACTA
C	CTAGTGAAGA	CGCGAATACG	TCAATCGAAG	ATCCAGCATG	CGCGAGCATT	ATAACATCCG	CAATCCGCAG	AATGTGCAGG	TCAGCGTAGA	AAGACCTCAA	GATACAAGCG	CCTAACCCGA
D	CCACCGATAT	AACCACGTAC	ATTGCGGTGG	CCGCTTAGGA	TTCTTGACTG	CAGCATATAG	GTCAACACAC	TGTCAATCTG	CACCTTCTAA	ATTCTAGCTC	TGAGAGGACC	GACAACCTGG
E	CTCTCGCGTA	TTGCCTACTG	CTTGAGTCCA	GCCTAGTGA	CCGTACTTGC	CGTTACATTC	AATACCGCTT	CCACGGTCTA	TACTCATGTG	CAGAATCAAG	CACACGTTGC	CCACTCATCG
F	AACCATCCGG	AACCTCGATT	CACTGACATA	AAGATTGAGG	CAAGCGCGTT	GGTCGAGTGA	TATGATCCTG	TGGAATACTC	CCAGAGTTAA	TCGCACGCTA	GTCTTAGCAG	TAAGCCACCG
G	TGCGTTATAG	CTGGAAGTGA	TATGGACTGA	ATCAATCACC	AACCTCAAACA	CTGTACCTAA	TTCCGTAGTA	ATCCAGACCG	GCTAACTTGC	CATACGGAAT	AACAGGAAGG	GCGATGAATT
H	GTATTCGCAA	TACTGCACGG	GATTCTGTAC	CTGCCTGTGA	TTAGCGGTTA	CCTTATATGG	CACATAATCG	CATCAAGTAG	TGCCATCACC	CTCGTGATCC	GACTATCAGA	CTTGTGCGTA

i5 indices for UDI plates - i5-TR96-B												
	1	2	3	4	5	6	7	8	9	10	11	12
A	GAGGCTATGC	CGACACGTAT	CACGATTGTT	CTTCGACGTT	CAAGAGTACC	GAAGGACTAC	GGATGAACGT	CTCAATAGTG	TGATACTCGA	CGAGAGATTG	ATCTGCGAGG	CTACTCTTCC
B	CCGGTGTATT	TGCGCTTCTC	GTCCAGATGC	TTGTCAAGTAC	GGACCGTAAG	TAACGGCGGA	CGGTTGTCCA	TTCCACAGTA	CAACTTCGAG	TCAGGACCAC	TCCTGCAACC	TGTGCTCAGG
C	TGTGTAGGTT	CATCTATCGG	CACCTTGATT	CACAGACAAC	CAGCACGGTT	CCGTAATGTT	GTGGAACATT	TAGCTTGCGG	CGATAGCGTT	CGGTACGTGA	TTATCCGTGC	CCACTTGTGC
D	CATTACAGAG	TGAAGTGAAC	TCTGAGGATC	GAGAGTCGAG	GGACAATCTG	TAACAGCTTC	CAACCTCGGA	CTTGAGTTCG	CAGGCGGATA	TTAGCTGTTC	AAGCGCATGA	CCGCTTCTTT
E	CATTACTAGG	CTGCAAGGAT	CCATTAGTCA	AACAAGGATG	TTAAGCGGCG	CTATTCAACG	TACGGTGTGC	TACAATCGAC	TCCATCGGTT	TTCTGTGTCG	TCGTAAAGCA	CAGATTCCGG
F	TTCTCGATAG	GAAGTTACAG	GTTGAGCTAT	GCAACAACGT	TGGTCAAGAT	TACCACACGG	GCGCCTATAC	AACGGCATAG	CTCCTCGTGA	CTCGCTGCTT	CCGTGATTGC	CATATACAGC
G	TCTGAGGCGA	CGATGTACGA	GGATGCAAGC	GATCAAGACC	CCACGGTATG	GAGACTAATC	GAATGATGAG	CCAGAACCAG	TAGGAGGATT	CCACATAACC	TCAACCGTCA	GCTTCTTCAG
H	GGCACAGCAT	GTCCGCTAGA	CCAATACTAC	CCTAGTTGCA	CTGACCTGTA	GAGAATCCAT	GCAGCACCTA	TTGCACCAGA	CCGGTTAAGC	GAATAGTTGG	GCTTCTAACG	GGATACCTTA

**Important Note:** The first lot of Index set 2B (Lot #20220727) contained two barcodes (i7 plate well C6 and i5 plate well F10, highlighted in yellow above) that overlapped with indices in set 1. This has been corrected in all future lots of set 2 to remove the overlapping barcodes. See below or refer to the downloadable index list for the correct indices for all future lots of set 2.

**purePlex Index Set 2 (Part No. 301068) – All other lots except for 20220727**Plate Layout - **Set 2**

i7 indices for UDI plates - i7-TR96-2												
	1	2	3	4	5	6	7	8	9	10	11	12
A	CACACATCTC	GATCGACGTA	TCTTCAACGG	TCTTCGCTCA	AAGTGCAGCG	TCCGGTCATA	AAGCTGACGG	TCATAGGACG	GCCTTCTTCG	TTCCAGCCAG	CTCCACTGAA	CTACTACTTG
B	TGTGCTTGAA	CTCTGGTTCC	CGTAACTAGC	AACACCATGC	ATCCTACGCC	CCGTCGCTAT	CCGTTCTACC	GGAACATGAC	AACAACCGCA	GACTCAGGTT	CATAGTCGCC	ATGAGGACTA
C	CTAGTGAAGA	CGCGAATACG	TCAATCGAAG	ATCCAGCATG	CGCGAGCATT	<b>GACTCGATCC</b>	CAATCCGCAG	AATGTGCAGG	TCAGCGTAGA	AAGACCTCAA	GATACAAGCG	CCTAACCCGA
D	CCACCGATAT	AACCACGTAC	ATTGCGGTGG	CCGCTTAGGA	TTCTTGACTG	CAGCATATAG	GTCAACACAC	TGTCAATCTG	CACTTCTAAC	ATTCTAGCTC	TGAGAGGACC	GACAACCTGG
E	CTCTCGCGTA	TTGCCTACTG	CTTGAGTCCA	GCCTAGTGA	CCGTACTTGC	CGTTACATTC	AATACCGCTT	CCACGGTCTA	TACTCATGTG	CAGAATCAAG	CACACGTTGC	CCACTCATCG
F	AACCATCCGG	AACCTCGATT	CACTGACATA	AAGATTGAGG	CAAGCGCGTT	GGTCGAGTGA	TATGATCCTG	TGGAATACTC	CCAGAGTTAA	TCGCACGCTA	GTCTTAGGAC	TAAGCCACCG
G	TGCGTTATAG	CTGGAAGTGA	TATGGACTGA	ATCAATCACC	AACCTCAAAC	CTGTACCTAA	TTCCGTAGTA	ATCCAGACCG	GCTAACTTGC	CATACGGAAT	AACAGGAAGG	GCGATGAATT
H	GTATTCGCAA	TACTGCACGG	GATTCTGTAC	CTGCCTGTGA	TTAGCGGTTA	CCTTATATGG	CACATAATCG	CATCAAGTAG	TGCCATCACC	CTCGTGATCC	GACTATCAGA	CTTTCGCGTA

i5 indices for UDI plates - i5-TR96-2												
	1	2	3	4	5	6	7	8	9	10	11	12
A	GAGGCTATGC	CGACACGTAT	CACGATTGTT	CTTCGACGTT	CAAGAGTACC	GAAGGACTAC	GGATGAACGT	CTCAATAGTG	TGATACTCGA	CGAGAGATTG	ATCTGCGAGG	CTACTCTTCC
B	CCGGTGTATT	TGCGCTTCTC	GTCCAGATGC	TTGTCAAGTAC	GGACCGTAAG	TAACGGCGGA	CGGTTGTCCA	TTCCACAGTA	CAACTTCGAG	TCAGGACCAC	TCCTGCAACC	TGTGCTCAGG
C	TGTGTAGGTT	CATCTATCGG	CACCTTGATT	CACAGACAAC	CAGCACGGTT	CCGTAATGTT	GTGGAACATT	TAGCTTGGCG	CGATAGCGTT	CGGTACGTGA	TTATCCGTGC	CCACTTGTGC
D	CATTACAGAG	TGAAGTGAAC	TCTGAGGATC	GAGAGTCGAG	GGACAATCTG	TAACAGCTTC	CAACCTCGGA	CTTGAGTTCG	CAGGCGGATA	TTAGCTGTTC	AAGCGCATGA	CCGCTTCTTT
E	CATTACTAGG	CTGCAAGGAT	CCATTAGTCA	AACAAGGATG	TTAAGCGGCG	CTATTCAACG	TACGGTGTGC	TACAATCGAC	TCCATCGGTT	TTCTGTGTCG	TCGTAAGACA	CAGATTCCGG
F	TTCTCGATAG	GAAGTTACAG	GTTGAGCTAT	GCAACAACGT	TGGTCAAGAT	TACCACACGG	GCGCTATATG	AACGGCATAG	CTCCTCGTGA	<b>AAGCCAGATG</b>	CCGTGATTGC	CCTATACAGC
G	TCTGAGGCGA	CGATGTACGA	GGATGCAAGC	GATCAAGACC	CCACGGTATG	GAAGCTAATC	GAATGATGAG	CCAGAACCAG	TAGGAGGATT	CCACATAACC	TCAACCGTCA	GCTTCTTCAG
H	GGCACAGCAT	GTCCGCTAGA	CCAATACTAC	CCTAGTTGCA	CTGACCTGTA	GAGAATCCAT	GCAGCACCTA	TTGCACCAGA	CCGGTTAAGC	GAATAGTTGG	GCTTCTAACG	GGATACCTTA

## purePlex Index Set 3 (Part No. 301069)

Plate Layout - **Set 3**

## i7 indices for UDI plates - i7-TR96-3

	1	2	3	4	5	6	7	8	9	10	11	12
A	AAGTTAGTCC	TTGAGCGCGA	CAGTGCCTT	CAGTATGTTG	ATTGCGCGAC	CATTGTACCA	CGGTGCAGAA	TTACCACCTT	AAGATGCGTG	GGAACCGGAT	GTGAGAACCG	CGGTCTACTA
B	GCCTCGATAT	TCGGAATAGA	ATAAGCCGCA	TATGCGAAGC	TTGAACACCG	CAGTACGTCC	CACAGGTATA	GAGATTAACG	TAGCGCCGAA	TTATTGGTGC	TTGTTGCTAC	CTGACACAAG
C	TGGACCTGTG	ATCCGCTACA	CGATAACCGG	ATGCCACTTC	TAAGTCATGG	TCTTCGAGTA	AAGGAGTACA	CAATAGTAGG	CAACATCAGA	CACCAGCTTC	AAGGTGGTCA	CCATTGCGGA
D	CCGTTCCAAAG	GGTCTTCCAC	ATGGAGCTAG	AACCTTCTTG	GACGCATGGA	AATAACGTCC	AATCATCGAC	TCTATTGTGG	GATAATTCCG	TTAGGTACGC	TAGTTGCCGG	ATGACTGGAA
E	GAACCATCAT	TTCGAGGACG	GAGGTGCCAT	CCGCATACAC	GCAACAGAGG	AACGTACTAG	AAGCCACATG	CGGACTCTGA	CCAAGCCTAC	CAACCTGCCA	GTGCTGCTTA	ATTGATAGCG
F	CATAAGGTCA	TGCAGTCAAT	CAACATGGTG	TTGAATCGCA	AACACGCCAC	TGTGGAATGG	GAGCAGCCTT	CATGTCCAAT	TTGCGTCATG	GGAATAGTCC	TTACACCAAC	TCCATTAAAG
G	GAATCATGCC	GTGATATGCA	GGAGTCTTAA	AAGAACACAG	GAAGTAACAG	AATAATGCGC	TTCAGTAACG	CTTAGGATAG	TTAGCCGCCA	GTGTCCAATA	CGTTCAGTAT	TTGGACGTAG
H	TCTACTGCTA	CATGAATCCG	AAGATGCGAA	CCTCCACACA	AACCTCGGTAC	CTAGTCTGAT	TAGACCAATC	GTTGCGGCTA	GATACCGCAC	CACGACCAAC	TTCCGCTCGA	CTTGCCACCA

## i5 indices for UDI plates - i5-TR96-3 (Type A Sequencers/Workflow A)

	1	2	3	4	5	6	7	8	9	10	11	12
A	CACAATCCTA	CGAATCAATG	TTCTAGTACC	ATCCAGCGCA	CATGCAATAG	CTCTTGTGTT	AACGAGCAGC	CCAATTTCGTT	TAGTGGTTAG	CCTAGTCTTA	CTAGCAACTT	ATCGACATCA
B	TGATTGAAGG	GTGAACCTAT	AATCGAGCTG	TCCAGACCTT	TTGAAGCTGC	CCGTCTTAT	GGTGTGCTTT	CACCTTACGG	TGCGGAGCTT	ATGGTCTTAG	GAGCTTCGGA	CCTTACCATT
C	TGGAGCCAGC	TATTCTCTCT	AAGACCTGTT	GTTAGGATCA	TGACTGAGGC	CGACCATTAC	GGTGAATAGG	TAGCGGCGTT	TTCGATCCAC	GAATGCGACC	GATACGGACC	GAAGGCAACA
D	CCATATTGAG	TGAAGCATTG	TTGAGTCTAC	CGCTTCTTCC	GGTGCATGAA	GATGTGACGC	GTACTCTTGA	ATTGCGCGAC	GCTGAATGCG	TTGGCGTTAG	CGTTGGTAAT	CACCTTGTA
E	CTTAGCATTA	TGGTATCGAG	TTATGGCGAG	CATGTGATAC	GGATATACGG	AACAGGTAGG	CACCTAAGAA	CCATTGCGAA	GATCAGCGAC	CCATCCGAAT	GTATTAGAGC	TGCGAAGTAG
F	AATCACACGC	GCATAAGCAG	GATCCGCTAT	GAGATTACTG	CATAATTCCG	ATAGGTTCCGG	TTCAACACCA	GCTACGATAC	TCCGCGGTAA	GTACACACAC	TAATAACCGG	TAAGGCGGTT
G	GAATTGTACC	CACTGATGAA	TTCTGGTTAC	TATCTTTGAC	TCCTAACCTA	CTGTAGCAAA	GCATAACTTG	ATCGCTCTGG	CTAATCAGAG	CCGATTGAA	GCTGCGTGAT	TATCGTACTC
H	CGGTAGATTG	CAGCGACTTC	GCATTACATC	GTAGTGCATT	GAACATCTTG	TTGCGGAACG	CCAGGTAGAT	CTAGACGCAG	TTCGCAACTG	GCAGTCTGAT	CAAGTAACCA	GAAGACCATC

## purePlex Index Set 4 (Part No. 301070)

Plate Layout - **Set 4**

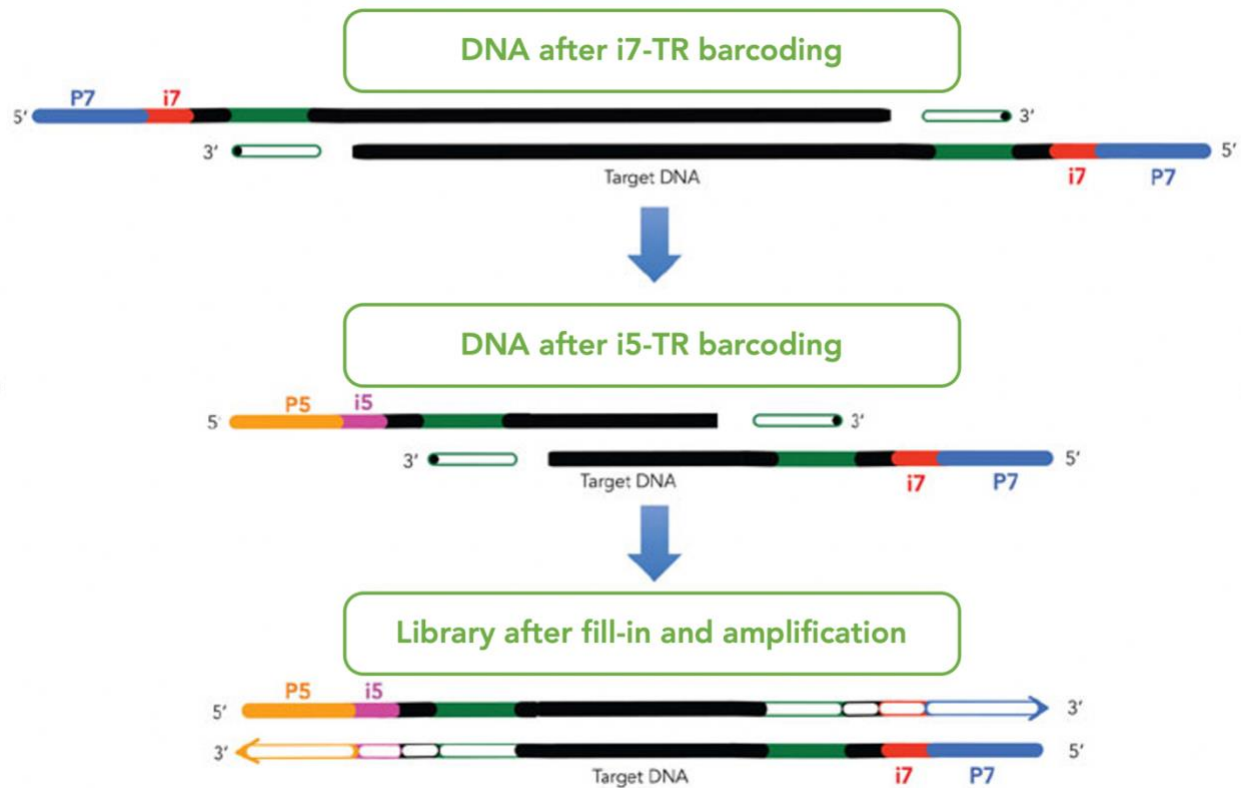
## i7 indices for UDI plates - i7-TR96-4

	1	2	3	4	5	6	7	8	9	10	11	12
A	TGATTGCGCC	GTTATCCGAA	ATCTGCGGAC	GGAACGCCTA	CACCTAAGAG	CTGCTAGGAA	CTGGTGCTTA	GGTAACGGTA	GGTACAGCCA	CCATGTACTA	GCTTCTAATG	TTGTTCCAGC
B	TCATATAGCG	TCCAACGACC	CATCCGTATA	ATAAGCAGTG	GCGCCATAGA	AATACCGAAG	GACCTTATAG	TAATACCAGG	TGCACTGACC	ATTCTCCAAG	CCTTGAGAAT	TTAAGCCCTC
C	ATCTAGGAAC	CGAGGTGTAA	GAATTGGAAC	CGCTATGGTA	GTACCACCGA	ATTGAACGTG	CGAAGCAGGA	GCGAAGCGTT	ATGCCGTAAC	GCACATTCTT	CAATATCTGG	GTGAAGTGTG
D	CGTCGATCAG	TCTCGGATAC	CTACGCGCTA	GAGTTCAGCA	CCAATCCAA	TTCCACATCA	GTACTGCTGC	ATGCGATTGG	ATGTTAGTGG	TGCTTGGAGA	CATACACCTA	CCTTATCGCC
E	CCGCCACATT	ATCGAGGTCA	CGGACGTAAC	TCCGACGCAT	AAGCGTGCAA	TCCACAGTTC	TGTAAGTTGC	TACCATGCAA	CGAGCTTGCA	GCTTGAAGTA	GTTCACGACG	ATCATTGTGC
F	ATTATCTGCC	CAAGAATGAC	TTCTCTAAGC	GGAATACAGA	CGACCATGTA	GGTAGTTGTC	GAAGGTTGCA	CGTCCTGAGA	CAACCAAGCA	ATAACGTAGG	CGTATCCAGG	CAAGCGTAAG
G	CCGAAGACAA	CAACCTTATC	AATAGCAGAG	CAATCGGTGA	AATAGSTTGG	CCGACAATAA	CATGTAAGAC	GGAACCTTATC	GCCGTCCTTA	AACTTCCGAG	GCACGATAAG	CACCTATGCC
H	CAAGTACGAG	TGAGCTTGTG	CAAGGATTAG	AATCCAGCGC	CTAATTCAAG	GCATAACCAA	GAGCGTGATC	GCTTCGACCA	CTGGAACAA	GTGCGGGAAT	CGGATAATAG	AACAACAACC

## i5 indices for UDI plates - i5-TR96-4 (Type A Sequencers/Workflow A)

	1	2	3	4	5	6	7	8	9	10	11	12
A	ATATCCTCCA	GTGTTAGGCG	GGCTGTGGTA	TTCTAGGCTT	GAGAGCAACC	TTGTTGGGAG	TGAGCGGTTA	TTCTGTACCC	CGCAAGGTTA	GGTCTGATTAC	GGTCTGAAAC	TTGCTTGGCC
B	GGCGGTGAAT	GCTATTCTGG	CTCTAGTTCG	TTGCTTGTCA	GACTATGTTT	CGAGGATGTT	ATTGACCTAC	CGTGGTTATT	TGTACTATGC	TAGGTAGACA	CCGCTATCAG	GTAGGACGTA
C	AAGAGCTGCA	ATCCGTTGTA	GAGTAGTCC	CATTCTTAAC	GATCATTCAG	AATCGTGCTA	TAACATCGCG	GTAGCTCTAG	CTTACATACC	GTAAGATGAC	GAGAGGAAGG	TACACCGCTG
D	CCGCTACCTT	CTCTCTCTCT	CATACAGAAA	GGTAAGGAAT	GATGTCGTTA	GACTGGAAAC	GAAGGTCATG	GCTGCACACA	TTCAAGCAGG	CTCAAGCTAT	TTACTTGGAC	TTCTTGCAT
E	TAACGGTAAC	GTTGAATGGC	GAAGATGTCG	TGTCGAGGTA	CGACTTAATC	CACAATATGG	GAGAAATCAG	GGACCAAGTA	TCAGGTACTA	GTGGAAGTGG	TTTGTGGCTG	CCTAATATGC
F	AAGTGGAGCA	AACTAATGGC	CCACGCTAAC	TGCGATCAGC	AATGGTTTAC	AATGGCAAGAA	ATCTCCGTGC	TGGAGCGTCA	GCTCCACTTA	CGCACTTCTA	TAACCAAGTA	TGAGCGGTAG
G	TCCAACCTCG	CACTTTCAGA	CAGTTATGAG	TAATGTCCTC	GAGGTAATCG	GGACAACAAT	AACGTCTAAG	TATGTGATGG	CGGATAGAA	GGTTTCATATC	TCGCGTCATC	CTTGTCTGGC
H	TGTACTGGAG	CATACACCAA	TTACCAAGGAG	CCACGGTTAA	CCACACATGC	CAGCTATGTA	AAGAAGCAGG	CTGTAAACGG	TTGAGTAACC	GTTAGTCAGC	CTAGCTGATA	TCAATCCGAA

## Appendix B: purePlex™ Library Intermediates



## Appendix C: Sample Sheet and Sequencer Loading Guidelines

Illumina sequencing systems differ in their use of sample sheets, availability of on-instrument demultiplexing, and run setup methods. If you have questions for your specific sequencer, contact Illumina tech support for the best guidance on setting up your run using your sequencing platform.

purePlex™ libraries use the Nextera Adapter sequences and are unique dual-index libraries using 10 nt indices for both the i7 and i5 index sequences. purePlex libraries do NOT require custom sequencing primers.

All Illumina sequencers read the i7 index in the forward direction (as listed in Appendix A or excel index list). The i5 index, however, is read differently on different sequencers depending on the version/chemistry of the sequencing kits. In this case, if using an Illumina sample sheet template, enter the i5 index in the forward direction as the sample sheet will auto-generate the reverse complement if needed. If demultiplexing using bcl2fastq and using a Workflow B sequencer, enter the reverse complement of the i5 index (provided in the excel under Workflow B).

As of January 1, 2021

The following Illumina sequencers use type A workflow (i5 read in forward direction):

- MiniSeq (rapid only)
- MiSeq
- HiSeq 2000/2500
- NovaSeq 6000 (v1 reagents)

The following Illumina sequencers use type B workflow (i5 read as reverse complement)

- iSeq 100
- MiniSeq
- HiSeq X
- HiSeq 3000/4000
- NextSeq 550
- NextSeq 1000/2000
- NovaSeq 6000 (v1.5 reagents)

## Appendix D: Sample pooling for plexities ≠24

The purePlex™ library preparation kit workflow has been optimized and validated using 24 samples per pool, with columns 1-3 in pool A, columns 4-6 in pool B, columns 7-9 in pool C and columns 10-12 in pool D. However, because both the i7 and i5 tagging steps are performed on individual samples, the purePlex kit allow flexibility in pooling and batch sizes. Alternate pooling strategies can be employed and recommendations for best practices are outlined below.

### Pooling >24 samples

For pooling of >24 samples, follow the user guide as written through tagging reaction stop. At step 4, tagging reaction pooling, pool all samples into a single tube. Be sure to choose an appropriate tube size to accommodate the total pool volume (32 µl per sample) plus MAGwise beads (see MAX volume in table below). After pooling all samples, add MAGwise (0.75 volume equivalents) and mix thoroughly. Divide total contents into 1.5 or 2.0 mL tubes for purification. Examples are provided below:

Total Samples	MAX volume with bead addition	# of tubes for purification	Tube type for purification	Elution Volume per pool
25-32	1792	1	2.0 mL	(N/24)*48
33-47	2632	2	1.5 mL	48
48	2688	2	1.5 mL	48
72	4032	3	1.5 mL	48
96	5376	4	1.5 mL	48

Make the following adjustments during pool purification.

- Ensure supernatant clears entirely before removal of supernatant or transfer of eluate.
- Ensure 80% ethanol washes completely cover the bead pellet
- Elute each pool as specified and transfer 2 µl less than total elution volume to its own PCR tube.

For amplification, add 4 µl of library primer mix to each PCR tube, then add 1 volume equivalent (transferred eluate volume + 4 µl) of KAPA HiFi Hotstart Readymix. Run amplification reaction.

Following the amplification, combine all reactions into a single tube. Remove 5 µl and set it aside for troubleshooting. Measure the remaining volume of combined amplified material. Add 0.75 volume equivalents of MAGwise for purification. Follow incubation times etc. for the purification and adjust Ethanol volumes to completely cover bead pellet. For  $\geq 48$  samples, increase elution volume to 50 µL.

### **Pooling <24 samples**

In some cases, it may be desirable to pool/process <24 samples at a time. Depending on pooling size, it is recommended to use higher inputs of DNA. Specifically, for plexes between 12-23 samples, it is recommended to use 10-50 ng of input DNA per sample and for plexes of <12, it is recommended to use 25-50 ng of input DNA per sample.

Follow steps 1-3 as written in the user guide. At step 4, reaction pooling, combine the samples to be pooled into a single tube and measure volume (32 µl per sample). Add 0.75 volume equivalents of MAGwise. Follow incubation times and reagent volumes as outlined in the user guide. Continue following the user guide for steps 6 and 7 (library amplification and purification). Note that for pools of <24 samples the library yield may be lower than what is typically expected for a 24-plex pool.

*For processing of <8 samples, additional PCR amplification cycles may be needed to achieve high enough concentrations for sequencer loading.*

## Experienced User Checklist

*Centrifuge all reagents prior to opening. Pulse-fuge all reactions before and after incubations.*

### i7 Tagging Reaction

- ☐ Use a new 96-well PCR plate (REACTION PLATE)
- ☐ Add 5 µl sample to REACTION PLATE and pipette 5x
- ☐ Pipette 5 µl from i7-TR plate to each well being used of the REACTION PLATE and pipette 5x
- ☐ Add 5 µl 3X coding buffer to the REACTION PLATE and pipette 10x
- ☐ Run TAG incubation

### i5 Tagging Reaction

- ☐ Add 5 µl Normalization Reagent and pipette 10x
- ☐ Pipette 5 µl from i5-TR plate (same wells used from i7-TR plate) to each well being used of the REACTION PLATE and pipette 10x
- ☐ Run TAG incubation
- ☐ Add 12.5 µl X-solution to each well of the REACTION PLATE and pipette 10x
- ☐ Run STOP incubation
- ☐ Pool 32 µl from each reaction for every 24 samples (4 tubes for 96 samples)
- ☐ Complete purification (0.75 volume equivalent), eluting with 48 µl of 10 mM Tris and transferring 46 µl to amplification tube.

### Library amplification

- ☐ Add 4 µl Library Primer Mix to amplification tube
- ☐ Add 50 µl of KAPA HiFi HotStart ReadyMix and pipette to mix
- ☐ Run Fill\_Amp8 program
- ☐ Dilute reaction to 125 µl with 10 mM Tris. Transfer 120 µl to purification tube. Retain remaining 5 µl for electrophoretic analysis.
- ☐ Complete library purification (0.75 volume equivalent), eluting with 32 µl of 10 mM Tris



Version	Release Date	Prior Version	Description of changes
V20230327	March 27, 2023	V20230105	<ul style="list-style-type: none"> <li>Updated to include barcode sets 3 and 4</li> </ul>
v20230105	Jan 5, 2023	v20220715	<ul style="list-style-type: none"> <li>Changed barcode set names from "1A" to "1" and "2B" to "2"</li> <li>Updated to correct barcodes in set 2 for all lots after 20220727.</li> <li>Updated plate names from "i7-TR96-A" to "i7-TR96-1", "i5-TR96-A" to "i5-TR96-1", "i7-TR96-B" to "i7-TR96-2", and "i5-TR96-B" to "i5-TR96-2"</li> </ul>
v20220715	July 18, 2022	v20220128	<ul style="list-style-type: none"> <li>Updated to include Set 2B index information</li> <li>Updated components table and workflow diagram</li> </ul>
v20220128	Jan 28, 2022	N/A	First version

## Technical Assistance

For technical assistance, contact seqWell Technical Support.

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