plexWell™ WGS 24 Library Preparation Kit
for Illumina® Sequencing Platforms
(Part No. WGS24)

User Guide
v20220421
Introduction

This user guide provides detailed instructions for preparing 24 genomic DNA samples for Illumina Sequencing using the plexWell WGS 24 Library Preparation Kit. The intended use of this plexWell kit is for generating high quality, high complexity libraries from purified human, plant and animal DNA (For Research Use Only).

The primary advantages and benefits of using the plexWell WGS 24 Library Preparation Kit are a truly multiplexed library preparation workflow, superior sequencing performance, and significant labor/consumable cost savings.

This plexWell WGS 24 User Guide is optimized for 200 ng of high quality genomic DNA per sample and typically generates 750 – 1,500 fmoles of purified, size-selected multiplexed library output, ranging from 400 – 1,500 bp in library fragment length.

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1 Although optimized for the NovaSeq 6000, plexWell WGS multiplexed libraries are compatible with a range of Illumina sequencing platforms. Refer to Illumina technical documentation for specific instructions on how to set up a sequencing run for a Nextera-style single-indexed or dual-indexed library prior to loading a plexWell library on your particular model of sequencing system.
# Table of Contents

**Molecular Diagram** .................................................................................................................. 3  
**Workflow Diagram** .................................................................................................................. 4  
**plexWell Kit Components** ....................................................................................................... 5  

**Considerations before you begin**
- **Required Equipment, consumables, and reagents** ................................................................. 6  
- **Reagent handling** .................................................................................................................... 7  
  
**plexWell WGS 24 DNA Library Prep detailed protocol** ............................................................. 8  
  - **Sample-Barcoding (SB) Reaction Set-up** .............................................................................. 8  
  - **SB Reaction Stop** ................................................................................................................. 8  
  - **SB Reaction Pooling** ........................................................................................................... 9  
  - **SB Pool Purification** ............................................................................................................ 9  
  - **Pool-Barcoding (PB) Reaction Setup** ................................................................................ 11  
  - **PB Reaction Stop** .............................................................................................................. 11  
  - **PB Reaction Purification** ................................................................................................... 11  
  - **Library Amplification** ......................................................................................................... 12  
  - **Library Purification** ........................................................................................................... 13  
  
**Library QC** ................................................................................................................................. 15  

**Appendices**
- **A: plexWell WGS-24 index information** ............................................................................... 17  
- **B: plexWell Library Preparation Intermediates** ................................................................. 18  
- **C: Sample Sheet and Sequencer Loading Guidelines** ....................................................... 19  

**Experienced User Checklist** ..................................................................................................... 20
plexWell WGS-24 Library Preparation

**Molecular Diagram**

**SAMPLE BARCODING:**
- Disperse DNA into plexWell™ SB plate
- Sample 1
- Sample 2
- Up to 96 samples
- Sample barcode transposome

**POOL BARCODING:**
- Pool into single tube
- Pooled barcode transposome

**FILL-IN AND LIBRARY AMPLIFICATION:**
- Sequencer-ready library
plexWell WGS-24 Library Preparation

Workflow Diagram

Sample-Barcoding Module

1. Sample barcode (SB) reaction
2. SB reaction stop
3. SB pooling
4. SB pool purification

Pool-Barcoding Module

5. Pool barcode (PB) reaction
6. PB reaction stop
7. PB reaction purification

Library Generation

8. Library amplification
9. Final library purification

56 minutes (20 minutes hands-on)

50 minutes (15 minutes hands-on)

55 minutes (15 minutes hands-on)
Table 1. Components of plexWell WGS 24 Library Preparation Kit

<table>
<thead>
<tr>
<th>Box Ref.</th>
<th>Component</th>
<th>P/N</th>
<th>Description</th>
<th>Storage</th>
<th>Qty</th>
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<tbody>
<tr>
<td>WGS24-1</td>
<td>Sample Barcode Plate</td>
<td>SBW24</td>
<td>SBW24 Plate: assay ready SB reagent in low profile, fully-skirted 96-well green PCR plate (columns 1 – 3)</td>
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<td>WGS24-2</td>
<td>Coding Buffer (3X)</td>
<td>CB0384</td>
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<td>X Solution</td>
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<td>MAGwise™ Paramagnetic Beads</td>
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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 and 2.0 ml, DNA LoBind Tubes)
- PCR plate seals (must be evaporation-resistant)
- 96-well thermal cycler (compatible with low profile fully-skirted 96-well PCR plates, Biorad HSP 9641)
- 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 Amp4:**
  - 72°C for 10 minutes (fill-in)
  - 95°C for 3 minutes (initial denaturation)
  - 98°C for 30 seconds
  - 64°C for 15 seconds 4 cycles
  - 72°C for 30 seconds
  - 72°C for 3 minutes (final extension)
  - 4°C hold
Before starting the procedure:

Adjust input DNA concentration. Assay the DNA concentration of each sample (n=24) using a PicoGreen assay. Before adding DNA to the PicoGreen assay, we recommend serially-diluting a small aliquot of genomic DNA in 10 mM Tris-HCl, because highly concentrated DNA is notoriously difficult to quantify accurately. This protocol requires approximately 200 ng of purified genomic DNA input per sample. Before starting library preparation, adjust each input DNA concentration to 25 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 SBW24 Plate 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. If stored cold, warm at 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 if necessary). 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 dissolve or dilute input DNA because 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 procedure, review the adjusting sample concentration guidelines on the previous page to ensure that each sample has been adjusted to 25 ng/µl using 10 mM Tris-HCl, pH 8.0.

1. Sample- Barcoding (SB) Reaction Set-up

   a. After labeling the assay-ready Sample-Barcode Plate (SBW24 Plate), pulse-spin the SBW24 Plate in a centrifuge. After centrifugation, visually inspect columns 1 - 3 to confirm that the volume of sample-barcode reagent in all 24 wells appears uniform, and then carefully remove the plate seal from the SBW24 Plate. Set-up SB reactions at room temperature.

   b. Add 8 µl of input genomic DNA (25 ng/µl) to all wells (one sample per well). Mix the DNA thoroughly with the sample-barcode reagent in each well by pipetting up and down (10 times at 8 µl), being careful not to introduce excessive bubbles. Use clean tips for each sample.

   c. Next, carefully pipette 8 µl of Coding Buffer (3X) to each well (n=24) of the SBW24 Plate, using new tips for each transfer. Mix thoroughly and slowly by pipetting up and down (20 times at 8 µl), being careful not to introduce excessive bubbles. 

      **Useful Tip:** Aliquot 30 µl of Coding Buffer (3X) into a PCR 8-tube strip, and then use a multichannel pipettor to dispense 8 µl into the SBW24 Plate and to mix.

   d. Seal the SBW24 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. SB Reaction Stop

   a. Confirm there is no precipitate in the X Solution. Slowly pipette contents of tube 5 times to ensure everything is mixed.

      **Note:** This solution contains SDS and vigorous mixing will cause it to foam. Pipetting slowly and under the surface of the solution will give the best results.

   b. Pulse-spin SBW24 Plate and then remove seal.
c. Add 12 µl of X Solution to each well of the SBW24 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 45 µl of X Solution into a PCR 8-tube strip, and then use a multichannel pipettor to transfer 12 µl from the strip into columns 1 - 3 of the SBW24 Plate and to mix.

d. Securely reseal and pulse-spin the SBW24 Plate, pulse-fuge, and transfer to a thermal cycler and run the STOP program, below, with lid heating on:
   68°C for 10 minutes
   25°C hold

3. SB Reaction Pooling

a. Pulse-spin the SBW24 Plate and then remove seal. Using a P200 pipettor, transfer 26 µl of stopped SB reaction from every well (n=24) into a 2 ml LoBind tube. The total volume of the SB reaction pool will be approximately 624 µl. Note: It is important to transfer an equal volume from every SB reaction. After pooling, check the volume remaining in the wells of the SBW24 Plate to verify that none of the wells were missed during pooling. Do not pool together SB reactions from different SBW24 Plates.

4. SB Pool Purification

a. Vortex (or vigorously pipette) room temperature MAGwise Paramagnetic Beads to ensure that the beads are fully resuspended.

b. Add 624 µl (approximately 1 volume equivalent) of MAGwise to the pooled SB reactions and mix thoroughly by pipetting. Incubate on bench for ≥5 minutes to allow DNA to bind.

c. Incubate in a tube rack on the bench for 5 minutes to allow the DNA to bind.

d. Place tube 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, add 1.5 ml of 80% ethanol without disturbing beads.

ii. After 30 seconds, slowly remove and discard supernatant, being careful not to disturb the pellet.

g. Wash beads with 80% ethanol a 2nd time.

i. With tube in the magnetic stand, add 1.5 ml of 80% ethanol without disturbing beads. If this volume is insufficient to cover the bead pellet, add a larger volume.

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 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 65 µl of 10 mM Tris to bead pellet, remove from magnetic stand and pipet 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 on the bench for at least 5 minutes to elute the purified SB 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 64 µl of DNA eluate to a PCR tube. The transferred eluate contains the purified SB reaction pool.

**Note:** Transfer of some beads with the purified SB reaction pool will not inhibit the Pool Barcode reaction.

**Optional QC:** Use 1 µl of the purified SB reaction pool for PicoGreen. The expected concentration is 18-36 ng/µl. If you are outside this range, contact support@seqwell.com

**SAFE STOPPING POINT**
Proceed immediately to the next step, or store the purified SB reaction pool at -20°C.
5. Pool-Barcode (PB) Reaction Setup

a. Add 5 µl of Pool-Barcode Reagent to the PCR tube containing the purified SB reaction pool from the last step. Mix thoroughly by pipetting.

b. Add 34.5 µl of Coding Buffer (3X), and mix the PB reaction thoroughly by vortexing for 5 seconds or pipetting ≥10 times at ≥ 75 µl.

c. Pulse-fuge the PCR tube containing the PB reaction then transfer to a thermal cycler, and run the TAG program, below, with lid-heating on:
   55°C for 15 minutes
   25°C Hold

6. PB Reaction Stop

a. Pulse-fuge PB reaction, then add 52 µl of X Solution to the PB reaction. Mix thoroughly by pipetting 10 times.

b. Recap the PB reaction tube, pulse-fuge, and transfer to a thermal cycler and run the STOP program, below, with lid heating on:
   68°C for 10 minutes
   25°C Hold

7. PB Reaction Purification

a. Pulse-spin stopped PB reaction, then transfer the entire volume of each stopped PB reaction to a 1.5 ml LoBind tube.

b. Vortex (or vigorously pipette) room temperature MAGwise Paramagnetic Beads to ensure they are completely resuspended. Add 156 µl (1 volume) of MAGwise beads to the stopped PB reaction and mix thoroughly by pipetting.

c. Incubate at room temperature on the bench for 5 minutes in a tube rack (non-magnetic) to allow the DNA to bind.

d. Incubate on bench for at least 5 minutes to allow DNA to bind.

e. Place tube in magnetic stand and let beads settle. A pellet should form on one side of the tube and the supernatant should be visibly cleared after 3 minutes.

f. Remove and discard supernatant with pipette. Be careful not to disturb the pellet.

g. Wash beads with 80% ethanol.
i. With tube in the magnetic stand, add 500 µl of 80% ethanol without disturbing beads.

ii. After ≥30 seconds, remove and discard supernatant, being careful not to disturb pellet.

h. Wash beads with 80% ethanol a 2\textsuperscript{nd} time.

i. With tube in the magnetic stand, add 500 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 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 101 µl of 10 mM Tris to bead pellet, remove from magnetic stand and pipet 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.

i. Incubate the resuspended beads on the bench for at least 5 minutes to elute the purified DNA from the beads.

j. Return tube to magnetic stand and allow bead pellet to form on the inner wall of the tube (~2 minutes).

k. When the supernatant has cleared completely, carefully transfer 100 µl of DNA eluate to a clean 1.5 ml tube. The transferred eluate contains the DNA purified from the PB reaction, and it is now ready for library amplification.

**SAFE STOPPING POINT**

Proceed immediately to the next step, or, store the purified PB reaction at -20ºC.

8. **Library Amplification**

a. Add 16 µl of **Library Primer Mix** to the eluate in the 1.5 ml tube.

b. Add 100 µl of Kapa HiFi Hot Start ReadyMix (2X), and mix well by pipetting.
c. Divide the PCR mix evenly into 4 tubes of a PCR 8-tube strip (54 µl x 4 tubes).

d. Close the strip, pulse-spin and run the FILL_AMP4 program, below, with lid heating on:

- **Fill-in:** 72°C for 10 min
- **Initial denaturation:** 95°C for 3 min
- **4 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.

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

a. After PCR, pulse-spin and pool the four library amplification reactions together in a 1.5 ml LoBind tube. Measure the total pooled volume.

  **Note:** volumes normally change due to film-loss and evaporation during thermal cycling, so it is important to measure the pooled volume prior to purification below.

b. Retain 5 µL of the unpurified pooled library amplification reactions and set-aside on ice for electrophoretic analysis later.

c. Vortex (or vigorously pipette) room temperature **MAGwise Paramagnetic Beads** to ensure they are completely resuspended.

d. Add 0.70-0.75 volume equivalents of MAGwise beads to the pooled, unpurified multiplexed library. Mix thoroughly by pipetting up and down.

e. Incubate in a tube rack on the bench for 5 minutes to allow the DNA to bind.

f. 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.
g. 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.

h. 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 pellet.

i. Wash beads with 80% ethanol a 2nd 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 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 pipet 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.

j. Incubate for 5 minutes on the bench to elute the purified library from the magnetic beads.

k. Return tubes to magnetic stand and allow bead pellet to form on the inner wall of the tube (~2 minutes).

l. 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. The remaining 2 µl of eluate may be used for electrophoretic analysis.

**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 along with an aliquot of unpurified amplified library from step 9b through an 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). Alternatively, run 3 µl of purified library and 5 µl of unpurified, diluted, library on an 2% Agarose E-gel EX alongside the 1 kb plus ladder (NEB) to determine the median fragment length.

The majority of library fragments should be between 500 and 1,000 bp in length. However, plexWell 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.

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 WGS24 libraries should be diluted ~1:7 (unpurified library) and ~1:10 (final purified library) prior to loading them on a TapeStation High Sensitivity D5000 assay. For optimal sequencing results, use a region analysis for fragments of 200-1500 bp to determine the average cluster-able fragment length for size adjustment with SYBR based qPCR.

![Figure 1](image)

**Figure 1.** (Left) Representative plexWell library traces generated using a TapeStation 2200 with High Sensitivity DNA 5000 reagents and tapescreens. Libraries were prepared from 24 replicates of human genomic DNA (ATCC NA12878) using a final library purification of 0.7 (black) and 0.75 (blue) volume equivalents of MAGWise. (Right) Table of typical plexWell WGS24 library results. Data includes the concentration determined by qPCR using the Kapa Library Quantification kit and the average fragment length (200-1500) for the size adjustment. Insert size was generated by first sequencing the libraries using MiSeq v2 chemistry and aligning to the hg38 reference genome, determining the median insert for each of the 24 samples, then taking the average of the 24 values.
Figure 2. This data demonstrates that the same library may yield similar fragment size distribution on two different electrophoretic analysis. Libraries were prepared from 24 replicates of human genomic DNA (ATCC NA12878) using a final library purification of 0.7 volume equivalents of MAGwise. (Left) Representative plexWell WGS 24 library traces generated using an Agilent Bioanalyzer 2100 with High Sensitivity DNA reagents. (Right) Representative of the same plexWell WGS 24 library traces generated using a TapeStation 2200 with High Sensitivity DNA 5000 reagents and tapescreens.

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 in 11 a., above, for the size adjustment value used for calculating the purified, multiplexed library concentration. Library concentrations are typically 30 – 60 nM.

Sequencing Loading and Read Configuration
plexWell WGS24 libraries are dual indexed using 8 nt indices. Each pool contains 24 unique sample-specific i7 indices (Appendix B) and single pool-specific i5 indices. These libraries can be sequenced in dual index (R1, i7, i5, R2) if multiplexed with other libraries or single index (R1, i7, R2) if sequenced alone. plexWell libraries are sequenced using the same primers as Nextera® libraries and contain the same adapter configuration. Refer to Illumina technical documentation for specific instructions on how to denature and dilute a purified library on your model of sequencing system and for additional information on setting up the read configuration.

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2 The primers provided in TruSeq v3 Cluster kits are incompatible with Nextera-style libraries, including plexWell 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, GA/Ix, and HiScanSQ.

3 The sequencing primers provided in TruSeq v3 Cluster kits are incompatible with Nextera-style libraries, including plexWell 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, GA/Ix, and HiScanSQ.
Appendix A: plexWell WGS-24 index information

The most up-to-date list of indices for all plexWell kits is available in the Resources section for this product. Click on the “plexWell Kit Master Index List” to download an excel workbook containing all i7 indices (in list and plate layout) and i5 indices. The master index list is the easiest way to copy and paste index sequences into sample sheets or for demultiplexing.

### I7 indices by well of SBW plate

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**PBW014 (i5) index sequence (5’ to 3’):** CCTATTGA
Appendix B: plexWell Library Preparation 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.

plexWell libraries use the Nextera Adapter sequences and are combinatorial dual-index libraries using 8 nt indices for both the i7 and i5 index sequences. plexWell libraries do NOT require custom sequencing primers.

All Illumina sequencers read the i7 index in the forward direction (as listed in Appendix A and in the plexWell index list available under the resources section of all plexWell products at seqWell.com. 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, enter the reverse complement of the i5 index (provided in the Illumina Workflow B column in the plexWell index list).

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
  • NovaSeq 6000 (v1.5 reagents)
  • NextSeq System
Experienced User Checklist

_Centrifuge all reagents prior to opening. Pulse-fuge all reactions before and after incubations._

Sample Barcoding

- Centrifuge **SBW24 plate**
- Add 8 µl sample (@25 ng/µl) to **SBW24 plate** and pipette 10x
- Add 8 µl coding buffer to **SBW24 plate** and pipette 10x
- Run TAG incubation
- Add 12 µl X-solution to SBW24 plate and pipette 10x
- Run STOP incubation
- Pool 26 µl from each SB reaction
- Complete SB purification (1 volume equivalent), eluting with 65 µl
- Transfer 64 µl of purified SB pool to 0.2 ml PCR tube (PB reaction tube)

Pool Barcode Reaction

- Add 5 µl PB Reagent to 64 µl of purified SB pool and pipette 5x
- Add 34.5 µl coding buffer to PB reaction tube and pipette 10x
- Run TAG incubation
- Add 52 µl X-solution to PB reaction and pipette 10x
- Run STOP incubation
- Transfer Stopped PB reaction to a 1.5 ml LoBind tube and complete PB purifications (1 volume equivalent), eluting with 101 µl of 10 mM Tris
- Transfer 100 µl of purified PB pool to LoBind tube

Library Amplification

- Add 16 µl Library Primer Mix to PB eluate
- Add 100 µl of KAPA HiFi HotStart ReadyMix and pipette to mix
- Aliquot 54 µl of amplification mix into each of 4 x 0.2 ml PCR tubes. Pulse-fuge
- Run Fill_Amp4 program
- Combine amplification reactions into a 1.5 ml LoBind tube.
- Remove 3 µl of reaction for QC analysis. Measure remaining volume.
- Complete Library Purification (0.7 volume equivalent), eluting with 32 µl of 10 mM Tris, pH 8.0
## Revision History

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<td>20220429</td>
<td>20210402</td>
<td>● Table of Contents added</td>
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<td>● Updated PB reaction mixing steps</td>
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<td>● Updated workflow diagram, purification steps, and amplification program name</td>
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Technical Assistance

For technical assistance, contact seqWell Technical Support.

Email: support@seqwell.com
Website: https://seqwell.com/product/wgs24/