Takara Bio USA, Inc.

SMART-Seq® ICELL8® cx Application Kit User Manual

Cat. Nos. 640222, 640223, 640224 for ICELL8 cx CELLSTUDIO[™] v2.5 Software (051021)

Takara Bio USA, Inc.1290 Terra Bella Avenue, Mountain View, CA 94043, USAU.S. Technical Support: technical_support@takarabio.com

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I. Introduction

High-throughput full-length analysis of single cells

The SMART-Seq ICELL8 cx Application Kit protocol enables Illumina® sequencing and full-length transcriptome analysis of single cells isolated on the ICELL8 cx Single-Cell System.

The kit workflow (Figure 1, below) begins with staining and dilution of cell samples and the preparation of positive and negative controls, followed by dispensing of the cells and controls into the 5,184 nanowells of the ICELL8 350v Chip using the ICELL8 cx Single-Cell System and the ICELL8 cx CELLSTUDIO[™] v2.5 Software. As many as eight different samples can be analyzed in a single run, and the dispensing process is completed in approximately 15 min. During this time, cells are maintained in the humidity- and temperature-controlled environment provided by the ICELL8 cx Single-Cell System.

Following the dispensing of the samples and controls, the imaging component of the ICELL8 cx system is used to acquire images of the 5,184 nanowells with both blue and red wavelength filters, enabling visualization of Hoechst 33342 and propidium iodide staining, respectively. (Hoechst staining is used to visualize and identify candidate cells based on nuclei size/morphology criteria, while propidium iodide staining is used to further differentiate candidate from non-candidate cells.) Following image acquisition, images are analyzed either automatically or manually using ICELL8 cx CellSelect® v2.5 Software, which identifies nanowells containing single, viable cells that meet user-defined criteria. Upon identification and selection of candidate nanowells, CellSelect Software generates a file that will be used to direct the dispensing of RT-PCR reagents to selected nanowells for downstream cDNA synthesis and amplification steps.

Following freezing of the ICELL8 350v Chip, the chip is returned to the ICELL8 cx Single-Cell System and RT-PCR reagents are distributed to nanowells selected by the software. The chip is then transferred to the ICELL8 cx Thermal Cycler (referred to hereafter as thermal cycler), which is programmed to perform first-strand synthesis and amplification of cDNA in a single run. First-strand cDNA synthesis is initiated by the oligo dT primer (SMART-Seq ICELL8 CDS), followed by template switching (mediated by the template-switching oligo, SMART-Seq ICELL8 Oligonucleotide) and addition of an adapter to the 3' ends of full-length cDNAs. The first-strand cDNA is used as template for 2nd-strand cDNA synthesis, in which the added adapter serves as a priming site, allowing for unbiased amplification of full-length cDNA. The full-length cDNA is tagmented by TDE1 (Illumina Tagment DNA Enzyme) and the tagmented cDNA is amplified using forward and reverse indexing primers. The resulting library is extracted, purified, and amplified to yield the final sequencing-ready library. Following subsequent purification and validation steps, libraries are ready for sequencing on Illumina platforms.

By leveraging the capabilities of the ICELL8 cx Single-Cell System along with SMART® (Switching Mechanism At 5' end of RNA Template) technology, this protocol provides an efficient, cost-effective solution for high-throughput full-length transcriptome analysis of single cells.

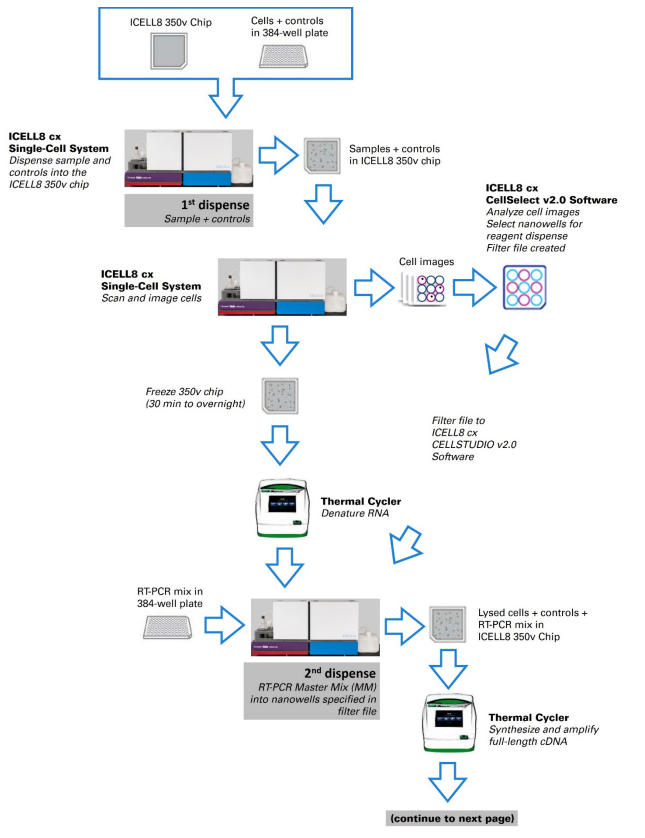


Figure 1. Complete SMART-Seq ICELL8 cx Application Kit workflow (continues on next page).

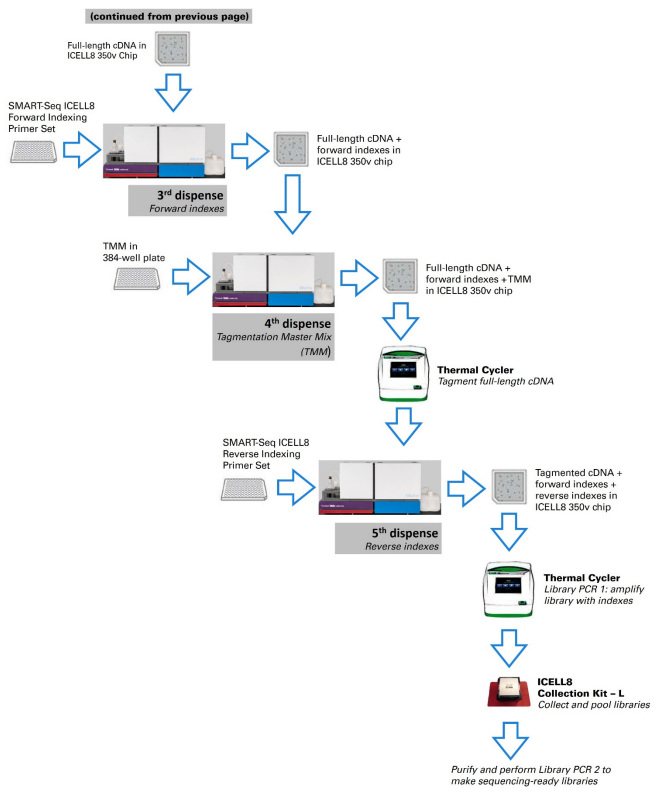


Figure 1. Complete SMART-Seq ICELL8 cx Application Kit workflow (continued from previous page).

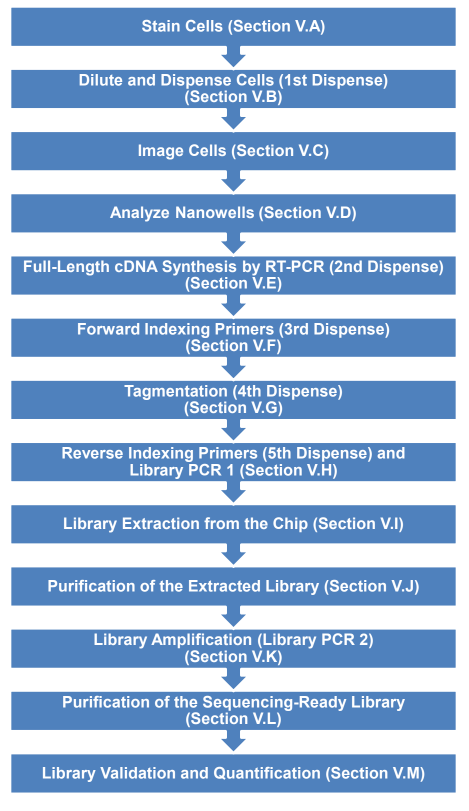


Figure 2. Protocols in the SMART-Seq ICELL8 cx Application Kit workflow.

II. List of Components

The SMART-Seq ICELL8 cx Application workflow requires use of an ICELL8 cx Single-Cell System (Cat. No. 640188 or 640189) and the SMART-Seq ICELL8 cx Application Kit:

- SMART-Seq ICELL8 cx Application Kit 1 Chip (Cat. No. 640222)
- SMART-Seq ICELL8 cx Application Kit 3 Chip (Cat. No. 640223)
- SMART-Seq ICELL8 cx Application Kit 5 Chip (Cat. No. 640224)

Each kit contains the necessary chip(s), reagents, and consumables required to run a total of 1, 3, or 5 experiments (respectively) of single-cell, full-length transcriptome analysis.

Table 1. Available SMART-Seq ICELL8 cx Application Kit contents.

SMART-Seq ICELL8 cx application kits	640222 (1 Chip)	640223 (3 Chip)	640224 (5 Chip)
Box 1 (ships at room temperature)			
ICELL8 350v Chip* (Cat. No. 640019)	1	3	5
ICELL8 Collection Kit – L* (Cat. No. 640212)	1	3	5
ICELL8 Loading Kit – B* (Cat. No. 640206)	1	3	5
Box 2 (ships at –70°C)			
SMART-Seq ICELL8 Reagent Kit* (Cat. No. 640202)	1		
SMART-Seq ICELL8 Reagent Kit* (Cat. No. 640203)		1	
SMART-Seq ICELL8 Reagent Kit* (Cat. No. 640204)			1
SMART-Seq ICELL8 Indexing Primer Set – A* (Cat. No. 640205)	1	2	3
SMART-Seq ICELL8 Indexing Primer Set – B* (Cat. No. 640218)		1	2

*Component can also be purchased separately.

III. Additional Materials Required

The following reagents and materials are required but not supplied with the SMART-Seq ICELL8 cx Application Kits or the ICELL8 cx Single-Cell System:

Required general lab supplies

- Personal protective equipment (PPE): powder-free gloves, safety glasses, lab coat, sleeve protectors, etc.
- Film Sealing Roller for PCR Plates ("film sealing roller") (Bio-Rad, Cat. No. MSR0001)
- PCR thermal cycler compatible with 0.2-ml tubes
- Minicentrifuges for 1.5-ml tubes and 0.2-ml tubes or strips
- 384-well plate orbital shaker
- Vortex mixer
- Centrifuges, rotors, and adapters. Recommended:
 - Eppendorf 5810R with swinging plate buckets, \geq 2,600g, and room temperature & 4°C operation
 - Kubota 3740 with rotor SF-240 for cell preparation
 - MICRO-TITER PLT BCKT PLT PK4 (VWR, Cat. No. 53513-874)
- ICELL8 384-Well Source Plate and Seal (Takara Bio, Cat. Nos. 640192, 640018, or 640037)
- Nuclease-free, non-stick 0.2-ml PCR tubes
- Nuclease-free LoBind 1.5-ml microcentrifuge tubes (Eppendorf)
- Conical tubes, 50-ml and 15-ml sizes
- 5-ml flip-cap tubes
- Single-channel pipettes: 2 μ l, 10 μ l, 20 μ l, 200 μ l, and 1,000 μ l

- Filter pipette tips: $2 \mu l$, $20 \mu l$, $200 \mu l$, and $1,000 \mu l$
- Wide-bore pipette tips: 200 µl and 1,000 µl
- Serological pipets and controller
- Nuclease-decontamination solution
- Exhaust hood system with UV

For staining and dispensing cells

- ICELL8 Chip Holder (Takara Bio, Cat. No. 640008); two chip holders are included with the instrument, and additional chip holders can be ordered separately.
- 1X PBS (no Ca²⁺, Mg²⁺, phenol red, or serum, pH 7.4; Thermo Fisher Scientific, Cat. No. 10010-023)
- ReadyProbes Cell Viability Imaging Kit, Blue/Red (contains Hoechst 33342 and propidium iodide; Thermo Fisher Scientific, Cat. No. R37610)
- Appropriate cell culture medium*
- TrypLE Express (Thermo Fisher Scientific, Cat. No. 12604-021)* *Required only if performing dissociation of adherent cells.

For tagmentation

• Illumina Tagment DNA TDE1 Enzyme and Buffer Small Kit (Illumina, Cat. No. 20034197) or Illumina Tagment DNA TDE1 Enzyme and Buffer Large Kit (Illumina, Cat. No. 20034198).

NOTES:

- Illumina Tagment DNA TDE1 Enzyme and Buffer Kits include Tagment DNA Enzyme (TDE1) and Tagment DNA Buffer (TD).
 - The Small Kit includes 0.17 ml (1 tube) of TDE1 and 1.24 ml (1 tube) of TD.
 - The Large Kit includes 0.65 ml (1 tube) of TDE1 and 1.24 ml (2 tubes) of TD.
- SMART-Seq ICELL8 cx Application Kit requires only TDE1. Do not use TD.

For library purification and validation

- SMARTer-Seq® Magnetic Separator PCR Strip (Takara Bio, Cat. No. 635011)
- Agencourt AMPure XP PCR purification system (5 ml; Beckman Coulter, Cat. No. A63880)

NOTES:

- Agencourt AMPure XP beads need to come to room temperature before the container is opened. Therefore, we strongly recommend aliquoting the beads into 1.5-ml tubes upon receipt, and then refrigerating the aliquots. Individual tubes can be removed for each experiment, allowing them to come to room temperature more quickly (~30 min). This aliquoting process is also essential for minimizing the chances of bead contamination.
- Immediately prior to use, vortex the beads until they are well dispersed. The color of the liquid should appear homogeneous. Confirm that there is no remaining pellet of beads at the bottom of the tube. Mix well to disperse before adding the beads to your reactions. The beads are viscous, so pipette them slowly.
- 80% ethanol: prepared fresh from anhydrous ethanol for each experiment
- Bioanalyzer instrument (Agilent) or similar

• Agilent High Sensitivity DNA Kit (110 samples; Agilent, Cat. No. 5067-4626)

NOTE: Alternatively, the libraries can be quantified by qPCR using the Library Quantification Kit (Takara Bio, Cat No. 638324).

- Qubit fluorometer (Thermo Fisher Scientific) or similar
- Qubit dsDNA HS Assay Kit (100 assays; Thermo Fisher Scientific, Cat. No. Q32851)

For cell counting

• Recommended: Moxi Z Mini Automated Cell Counter Kit, U.S. Version (ORFLO, Cat. No. MXZ001) with Moxi Z Cell Count Cassettes, Type M (25 pack; ORFLO, Cat. No. MXC001) or Moxi Z Cell Count Cassettes, Type S (25 pack; ORFLO, Cat. No. MXC002)

NOTES:

- Alternatively, you may use a hemocytometer or any preferred cell counter with demonstrated, accurate cell counting.
- Refer to a Moxi Z user guide for guidance in selecting an appropriate cassette size for the cells being analyzed.

IV. General Considerations

A. Protocol Best Practices

- Two different chips can be multiplexed together in a single sequencing run if one chip is indexed with primers from SMART-Seq ICELL8 Indexing Primer Set A (Takara Bio, Cat. No. 640205) and the second is indexed with primers from SMART-Seq ICELL8 Indexing Primer Set B (Takara Bio, Cat. No. 640218).
- Perform all experimental procedures in sterile environments with the proper personal protective equipment (PPE). Use designated UV hoods with proper ventilation for manipulating cells and setting up molecular biology reactions. Decontaminate gloves with nuclease decontamination solution, water, and ethanol. Change gloves routinely.
- Minimize the exposure time of unsealed chips, reservoirs, reagents, and other consumables to the open air. Wearing laboratory sleeve protectors may reduce the likelihood of introducing contaminants from exposed hands and arms.
- The assay is very sensitive to variations in pipette volume. Please make sure that all pipettes are calibrated for reliable reagent delivery and that nothing adheres to the outsides of the tips when dispensing liquids.
- All lab supplies related to cDNA synthesis need to be stored in a DNA-free, closed cabinet. Ideally, reagents for cDNA synthesis should be stored in a freezer/refrigerator that has not previously been used to store PCR amplicons.
- Use nuclease-free, molecular biology- or PCR-grade reagents to set up all molecular biology reactions.
- Add enzymes to reaction mixtures last and thoroughly incorporate them by gently pipetting the reaction mixture up and down.
- DO NOT change the amount or concentration of any of the components in the reactions; they have been carefully optimized for the SMART-Seq ICELL8 cx Application Kit workflow.

- Because of the large volume or viscosity of mixtures subject to purification using AMPure beads, each round of purification requires a very strong magnet. Place the samples on the magnetic separation device for ~5 min or longer, until the liquid appears completely clear and there are no beads left in the supernatant.
- UV-treat reagent reservoirs, seals, pipettes, filter tips, and compatible reagents prior to use.

B. Safety

Refer to safety guidelines in the user manuals for all equipment used in this protocol.



WARNING: Perform all experimental procedures in sterile environments with the proper personal protective equipment (PPE). Use designated UV hoods with proper ventilation for manipulating cells and setting up molecular biology reactions. Decontaminate gloves with nuclease decontamination solution, water, and ethanol. Change gloves routinely.



WARNING: Use of equipment and reagents for cell preparation and isolation with the ICELL8 cx Single-Cell System may cause exposure to toxic or biohazardous chemicals, thereby presenting a hazard. Always wear appropriate personal protective equipment (PPE), which should at minimum include gloves, eye protection, and a lab coat, when handling equipment and reagents and operating instruments.



Note and heed all warning labels on the instruments used in this protocol.

C. ICELL8 cx System Application Notes

Refer to the <u>ICELL8 cx Single-Cell System User Manual</u> for full details. Included below are general reminders.

- All dispensing steps in the ICELL8 cx Single-Cell System Stage Module should be performed with a 384-Well Source Plate oriented with the A1 well positioned at the top-right corner of the 384-well plate nest (Figures 3 and 4, below). The source plate must be fully seated. This may be accomplished by pushing the source plate down after it has been placed on the plate nest.
- All dispensing steps in the ICELL8 cx Single-Cell System Stage Module should be performed with the chip oriented with the chamfered (beveled) corner positioned towards the bottom-right corner of the chip nest (Figures 3, next page).

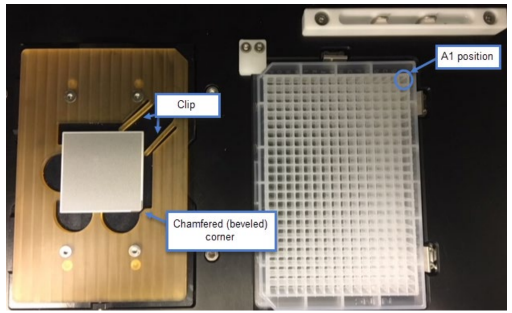


Figure 3. ICELL8 cx Single-Cell System Stage Module. (Left) chip nest. (Right) source plate nest.

D. ICELL8 350v Chip

Each ICELL8 350v Chip is engraved with a unique number (Figure 4). You can use this number to link your chip images and other experimental record files.

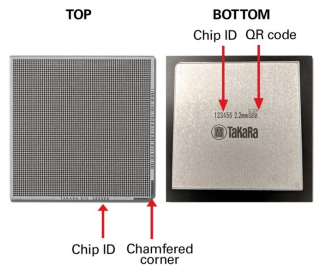


Figure 4. ICELL8 350v Chip features. (Left) Top view of the chip. Note the chamfered (beveled) corner at the bottom right. The "TaKaRa" logo and the chip ID (unique to each chip) are engraved on the chip border, near the chamfered corner. (**Right**) Bottom view of the chip. The chip ID is also engraved on the other side of the chip with a corresponding QR code that can be scanned by a barcode reader, allowing for the chip ID to be easily entered into the software.

E. Software

The instructions in this manual are written for use with CELLSTUDIO and CellSelect v 2.0 Software. Please refer to the <u>ICELL8 cx Single-Cell System User Manual</u> and the <u>ICELL8 cx CellSelect v2.5</u> <u>Software User Manual</u> for more detailed information.

V. Procedure

A. Protocol: Stain Cells

In this protocol, sample cells are stained with Hoechst 33342 and propidium iodide dyes that enable imaging, analysis, and selection of candidates suitable for downstream analysis following the cell dispense into the ICELL8 350v Chip. If starting from a nonadherent suspension culture, such as K-562 cells, use Procedure 1 (Section V.A.3); if starting from an adherent culture, such as 3T3 cells, use Procedure 2 (Section V.A.4), and then stain the trypsinized cell suspension using Procedure 1.

1. Cell and chip handling notes:

- This protocol requires several milliliters of healthy cell culture suspension with a cell density between 1 x 10⁵ and 7.5 x 10⁶ cells/ml. Some cell lines may require trypsinization to achieve a single-cell suspension.
- Keep cells at 37°C with 5% CO₂ in a cell culture incubator when not performing manipulations.
- Perform all wash steps in an exhaust UV hood. Avoid exposing the cell culture to ambient air to reduce the likelihood of contamination.
- Treat cells gently; do not vortex. Minimize bubble formation and frothing.
- Wear nitrile or powder-free gloves to reduce imaging artifacts.
- Centrifugation speed and time may need to be modified for different cell types.

2. Before you start:

- Perform a once-a-day warmup. Refer to the <u>ICELL8 cx Single-Cell System User Manual</u> (Section VIII) for more details.
- Prefreeze ICELL8 Chip Holder(s) (Figure 5) at -80°C.



Figure 5. ICELL8 Chip Holder.

- Set the chip centrifuge to 22°C.
- Prewarm 1X PBS (Ca²⁺ and Mg²⁺ free, pH 7.4), TrypLE Express*, and cell culture medium* at 37°C.

*Required only if performing dissociation of adherent cells.

- Dilute Control K-562 RNA (1 µg/µl; yellow cap) to 10 ng/µl for use in the next protocol (Section V.B, Table 2) as indicated in the following steps and keep the dilution on ice:
 - a) Dilute Control K-562 RNA to 50 ng/ μ l by mixing 38 μ l of Nuclease-Free Water (white cap) with 2 μ l of Control K-562 RNA (1 μ g/ μ l) in a sterile nuclease-free microcentrifuge tube.
 - b) Dilute Control K-562 RNA to 10 ng/μl by mixing 8 μl of Nuclease-Free Water with 2 μl of Control K-562 RNA (50 ng/μl) in a sterile nuclease-free microcentrifuge tube.

NOTES:

- Return Control K-562 RNA (50 ng/µl) stock solution to storage at -70°C.
- Diluted Control K-562 RNA should be kept on ice at all times.

3. Procedure 1: Staining cells in suspension

 Prepare a 1:1 mixture of Hoechst 33342 and propidium iodide. Combine 80 µl of each dye per ml of cells to be stained. Depending on your cell density and sample volume, scale as needed. An example using 2 ml of cells is described below (e.g., prepare 320 µl of premixed dye solution).

NOTE: Protect this mixture from light until ready for use.

- 2. Transfer \sim 2.1 ml of suspension cells to a fresh 5-ml tube.
- 3. Determine cell concentration using your preferred method or a Moxi automated cell counter and an appropriate Moxi Z cassette (e.g., use a Moxi Z cassette MF-M for K-562 cells; refer to a Moxi Z user manual for guidance in selecting an appropriate cassette size for the cells being analyzed). Use a pipette to transfer 75 µl of cell suspension to one end of the loading platform. Take a reading and record the cell concentration.
- 4. Add 320 μl of the premixed Hoechst 33342 and propidium iodide dye mix to the cells. Mix gently by inverting the tube 5 times. DO NOT vortex or overagitate the cells.
- 5. Incubate cells at 37°C for 20 min.
- 6. Add an equal volume of 1X PBS (prewarmed to 37°C) to stained cells. For the example described here, 2 ml of prewarmed 1X PBS is added to the 5-ml tube containing the stained cell suspension.
- 7. Mix the stained cell suspension and PBS by gently inverting the tube 5 times. DO NOT vortex or overagitate the cells.
- 8. Pellet the cells by centrifugation at room temperature. Avoid over-centrifugation or pelleting into a firm pellet or clump.

NOTE: Optimal centrifugation speed and time may vary depending on the cell type being analyzed. Examples:

- K-562 or 3T3 cells: 100g for 3 min.
- PBMCs or similarly sized cells: 500g for 3 min.
- 9. Gently remove tube from the centrifuge without disturbing the cell pellet.
- 10. Carefully decant the supernatant without disturbing the cell pellet.

- 11. Wipe remaining fluid from the tube top using a fresh Kimwipe such that the last remaining drop on the tilted tube top is gently removed.
- 12. Gently add 1 ml of 1X PBS (prewarmed to 37°C) to the tube side wall.
- 13. Use a wide-bore 1-ml pipette tip to gently mix the cell suspension by slowly pipetting up and down ~5 times. DO NOT vortex or overagitate the cells.
- 14. Count the cells using a Moxi automated cell counter and an appropriate cassette (or your preferred method). Take two readings for each stained cell sample and average the results. The recommended concentration range of stained cell suspension is $1.2-5.0 \times 10^5$ cells/ml. If the concentration is lower than 1.2×10^5 cells/ml, repellet the stained cell suspension and resuspend in a lower volume of 1X PBS to achieve a concentration in the recommended range.
- 15. Proceed to the next protocol (Section V.B).

4. Procedure 2: Preparing adherent cell types from a 75-cm culture flask*

*Adjust volumes accordingly for different-sized flasks.

- 1. Carefully remove culture media from a 75-cm flask containing adherent cells using a serological pipette.
- 2. Add 10 ml of 1X PBS prewarmed to 37°C by dispensing the PBS on the side walls of the flask. DO NOT pour PBS directly onto cells.
- 3. Wash the cells by tilting the flask gently. DO NOT mix by pipetting.
- 4. Remove the PBS from the cells using a serological pipette.
- 5. Add 3 ml of TrypLE Express prewarmed to 37°C to the flask to dissociate the cells.
- 6. The efficiency of cell dissociation from the flask surface may vary with cell type. Monitor the process visually using a microscope.
- When cell dissociation is sufficient, neutralize the trypsinization reaction by gently adding 7 ml of complete media (containing 10% serum), prewarmed to 37°C. DO NOT vortex or overagitate cells.
- 8. Follow the Procedure: Staining cells in suspension (Section V.A.1, above) starting from Step 1.

B. Protocol: Dilute and Dispense Cells (1st Dispense)

In this protocol, sample cells and controls are diluted and aliquoted into a 384-Well Source Plate and dispensed into the ICELL8 350v Chip using the ICELL8 cx Single-Cell System. Refer to the <u>ICELL8 cx</u> <u>Single-Cell System User Manual</u> for detailed information about instrument setup and operation.

1. Before you start

- Confirm that initialization and setup procedures for the ICELL8 cx Single-Cell System have been completed (see "Before you start" in Section V.A, above).
- Confirm the empty ICELL8 Chip Holder (Figure 5, above) is frozen at -80°C.
- Aliquot 300–500 μl of 1X PBS (Ca²⁺ and Mg²⁺ free, pH 7.4) on ice for positive and negative controls.

- This protocol requires a 384-Well Source Plate (with seal), a Plate Seal Applicator, a film sealing roller, and the following components: Control K-562 Total RNA (diluted to 10 ng/µl in the previous protocol), Second Diluent (100X; light blue cap), BSA (1%; red cap), RNase Inhibitor (white cap), SMART-Seq ICELL8 CDS (blue cap)—all from the SMART-Seq ICELL8 cx Reagent Kit—plus Blotting Paper and RC Film (from the ICELL8 Loading Kit B).
- Thaw Second Diluent (100X), BSA (1%), and SMART-Seq ICELL8 CDS on ice. Once thawed, keep on ice for the remainder of the protocol
- Use the concentration of stained cell suspension measured at the end of the previous protocol (Section V.A) and the information in Table 2 (below) to calculate the volumes of stained cell suspension and 1X PBS that should be combined for a final concentration of 1.4 cells/50 nl in a total volume of

1 ml.

2. Procedure

Prepare diluted stained cell suspension

- 1. Mix each component: Second Diluent, RNase Inhibitor, BSA (1%), and SMART-Seq ICELL8 CDS. Spin the tubes briefly to collect contents at the bottom.
- 2. In a 1.5-ml microcentrifuge tube, combine the volumes of Second Diluent, BSA (1%), RNase Inhibitor, SMART-Seq ICELL8 CDS, and prewarmed 1X PBS indicated in the corresponding column of Table 2 (below). Mix the combined reagents by vortexing, then spin the tubes briefly to collect contents at the bottom.

NOTE: The amount of 1X PBS added will depend on the starting concentration of stained cell suspension.

To the 1.5-ml microcentrifuge tube from the previous step (containing Second Diluent, BSA (1%), RNase Inhibitor, SMART-Seq ICELL8 CDS, and 1X PBS), add the calculated volume of stained cell suspension that will yield a final concentration of 1.4 cells/50 nl in a total volume of 1 ml (refer to the ICELL8 Cell Dilution Tool spreadsheet provided by the Field Application Specialist during training). DO NOT use a pipette to mix at this step.

NOTES:

- Before obtaining the sample, mix the stained cell suspension gently by inverting the tube several times.
- Take the required volume of stained cell suspension from the center of the tube using a 20-µl or 200-µl pipette tip and add it slowly to the tube containing the other reagents.
- Work quickly to avoid settling of cells.

Prepare positive and negative controls

4. Prepare positive and negative controls in separate 1.5-ml microcentrifuge tubes using the volumes indicated in Table 2 (below).

NOTES:

- Keep the positive control sample on ice.
- Mix well, but do not vortex the positive and negative control samples.
- The final concentration of Control K-562 Total RNA should be 10 pg/50 nl (equivalent to the total RNA content from 1 cell).

Components	Cap color	Negative control	Positive control	Diluted stained cell suspension	For each sample (volume per source well)*
Second Diluent (100X)	Light blue	1.00 µl	1.00 µl	10.00 µl	1.00 µl
BSA (1%)	Red	1.00 µl	1.00 µl	10.00 µl	1.00 µl
RNase Inhibitor (40 U/µI)	White	1.00 µl	1.00 µl	10.00 µl	1.00 µl
SMART-Seq ICELL8 CDS	Blue	1.92 µl	1.92 µl	19.20 µl	1.92 µl
Control K-562 RNA (10 ng/µl)	-	-	2.00 µl	-	-
Stained cell suspension	_	_	-	Dilute to 1.4 cells/50 nl*	Dilute to 1.4 cells/50 nl*
1X PBS (Ca ²⁺ and Mg ²⁺ free)	-	95.08 µl	93.08 µl	Up to 1,000.00 µl	Up to 100.00 µl
Total		100.00 µl	100.00 µl	1,000.00 µl**	100.00 µl***

Table 2. Sample preparation guidelines.

*Sufficient stained-cell suspension should be included such that the final concentration in the 1,000-µl volume of diluted stained cell suspension is 1.4 cells/50 nl (i.e., 28,000 cells/ml).

**The 1,000-µl total volume of diluted stained cell suspension is sufficient for distributing 80 µl of cell suspension in each of eight source wells (see Figure 6, below).

***The 100-μl total volume of diluted stained cell suspension is sufficient for distributing 80 μl of cell suspension into a single source well. Multiply appropriately for the number of source wells used for each sample type.

Prepare cell dispense source plate

Refer to the <u>ICELL8 cx Single-Cell System User Manual</u> for detailed information about instrument setup and operation.

5. Using a wide-bore 1-ml pipette tip, gently mix the diluted stained cell suspension prepared in Step 3 of this protocol (above) by slowly pipetting up and down ~5 times. DO NOT vortex or overagitate the cells.

IMPORTANT: Proceed to the next step quickly to avoid settling of cells.

 Using a 200-μl pipette tip, slowly and carefully load 80 μl of cell suspension into wells A1, A2, B1, B2, C1, C2, D1, and D2 of a 384-Well Source Plate as indicated in Figure 6 (below).

NOTES:

- Make sure to take each aliquot from the center of the tube containing the diluted stained cell suspension.
- Be careful not to splash liquid into neighboring wells.
- Make sure not to introduce bubbles when adding the cell suspension to the 384-Well Source Plate.
- DO NOT vortex or spin down the 384-Well Source Plate.
- DO NOT tap the plate. If any bubbles are present, remove using a pipette tip.
- 7. Add positive control and negative control to the 384-Well Source Plate as indicated in the following steps and in Figure 6 (below):
 - Add 25 µl of prepared positive control to Well P24.
 - Add 25 µl of prepared negative control to Well A24.

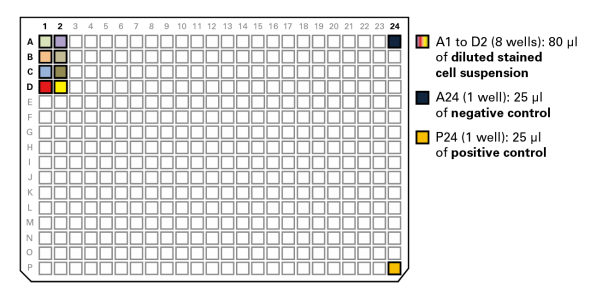


Figure 6. Setting up the 384-Well Source Plate for dispensing cell samples and controls.

- 8. Seal the 384-Well Source Plate with the 384-Well Source Plate Seal using the Plate Seal Applicator. DO NOT centrifuge the plate.
- 9. Open the packet containing the sealed ICELL8 350v Chip and remove the chip seal.

- 10. Place the chip on the chip nest in the ICELL8 cx Single-Cell System. The chamfered (beveled) corner of the chip should align with the chamfered corner of the chuck of the chip nest (refer to the <u>ICELL8</u> cx Single-Cell System User Manual, Section X.A).
- 11. Remove the seal on the 384-Well Source Plate and place in the ICELL8 cx Single-Cell System with the A1 corner positioned at the top-right corner of the plate nest. The beveled corners of the 384-Well Source Plate should be on the left side (refer to the <u>ICELL8 cx Single-Cell System User Manual</u>, Section X.B).
- 12. In CELLSTUDIO software, click the [Dispense cells and Controls (50 nl)] button (Figure 7).



Figure 7. Using CELLSTUDIO software to dispense cells into the ICELL8 350v Chip.

A "Select index set" *Workflow* window will display asking you to indicate which SMART-Seq Index Kit you are using, SMART-seq\SetA (Cat. no. 640205) or SMART-seq\SetB (Takara Bio, Cat. No. 640218). Choose the appropriate kit from the drop-down menu, and then click [Done].

ICELL8® cx CELLSTUDIO™ software	1											
Utilities Help												
DewPoint Chamber On Temp: RH:	:	Wash wate Available:	1 .	Vacuu	-		Firmware status Not connected					
Startup Advanced 3'DE - TCR SMA	RT-Seq											
72 x 72 : 350 n	Cells	PosCtrl	NegCtrl	R	T - P	CR m	nix	Tagn	nenta	tion	mix	
Chip ID	Norkflow							8	9	10	11	1
										145	161	
Dispense Cells and Co		Select	114	130	146	162						
	Index sets							115	131	147	163	
Scan chip	SMART-seq\SetA						\sim	115	131	147	105	
	SMART-seq\SetA SMART-seq\SetB							116	132	148	164	
				Done		Cance	el	117				
Dispense RT - PCR mix ¹ າວ	ommittered)	_										
		F			70	86	102	118				

Figure 8. Select index set workflow prompt. The display defaults to using Set A. Ensure the correct one is selected before clicking [Done].

13. Follow the subsequent software prompts and check the orientation of the source plate and ICELL8 350v Chip to ensure they are correctly loaded on the Dispensing Platform. Click [Done] to proceed.

Refer to the <u>ICELL8 cx Single-Cell System User Manual</u>, Section X.C "Dispense the Sample Cells and Experimental Controls into the Chip" for details about the dispense step.

- 14. After the sample and control dispense is completed, remove the chip from the chip nest and blot with blotting paper. Refer to the <u>ICELL8 cx Single-Cell System User Manual</u>, Section X.D "Blot and Centrifuge the Chip" for instructions to do that.
- 15. Seal the loaded chip with the RC Film (Figure 9, right)

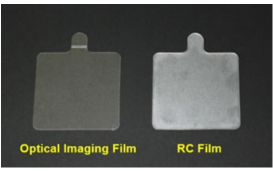


Figure 9. Imaging and RC films required for the dispenses. The Optical Imaging Film (left) has three layers. The RC Sealing Film (right) has a translucent backing. Please follow the instructions in each subprotocol regarding preparation and handling of the films.

Remove the liner from the RC Film and apply the exposed side of the film to the blotted chip (Figure 10, left, below).

16. Seal the blotted chip with the film using a film sealing roller (Figure 10, right, below).

NOTE: The RC Film is nonadhesive and can easily peel off. Be careful when handling.

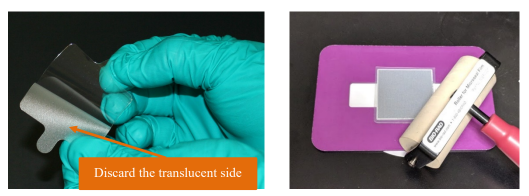


Figure 10. Preparing and adhering the RC film. RC film is composed of two layers. One layer is clear, and the other layer is translucent. **Plate A.** When sealing a chip, remove the translucent layer and discard it, and then put the remaining layer on the chip. **Plate B.** Tightly adhere the RC film using the film sealing roller. Make sure that the chip is securely sealed to avoid well-to-well contamination and evaporation.

17. Place the sealed chip on a centrifuge spin plate and centrifuge the sealed chip at 300g for 5 min at 22°C with full acceleration and full brake (Figure 11, below). If you have one chip, balance the centrifuge with the supplied Balance Chip or a blank chip. Proceed to the next protocol (Section V.C) once the centrifugation is complete.



Figure 11. ICELL8 cx Centrifuge Chip Spinner.

18. On the ICELL8 cx Single-Cell System, perform the [Tip Clean] procedure 3-4 times.

C. Protocol: Image Cells

In this protocol, images of all 5,184 nanowells of the ICELL8 350v Chip are acquired.

- 1. Refer to the <u>ICELL8 cx Single-Cell System User Manual</u>, Section X.E "Scan Chip for Single Cells and Freeze the Chip" for detailed information about this protocol, with the following guidelines:
 - a. Follow steps 1–5.
 - b. For step 6, the *New stack* info dialog window will display (Figure 12, below). The "Barcodes" field will be grayed out because the barcode file is pre-configured within the during the Dispense (Section V.B, Figure 8).

New stack info	
Chip ID	99995
Analysis settings	AnalysisSetting_350nL_chip.xml - Predefined V
Barcodes	<from file=""></from>
Chip comment	
	OK Cancel

Figure 12. New stack info dialog window.

NOTE: If an incorrect barcode file was selected during the Dispense step, it can be corrected during the next step.

Continue with the imaging procedure and then proceed to Section V.D.

- c. Continue with the procedure from steps 7–12.
- d. At Step 13, remove the liner from only one side of the Optical Imaging film and apply the exposed sticky side of the imaging film to the chip, sealing carefully with the Plate Seal Applicator (similar to Figure 10, above).
- Place the imaged chip into an empty ICELL8 Chip Holder that has been prechilled at -80°C (Figure 7). The chip holder should click closed and should close evenly, indicating a proper magnetic seal. Make sure that the imaging film is well sealed over the chip.
- 3. Freeze cells at -80°C for a minimum of 30 min before proceeding to RT-PCR (Section V.E, below).

SAFE STOPPING POINT: The chip can be frozen and stored in the chip holder at -80°C for subsequent processing.

D. Protocol: Analyze Nanowells with ICELL8 cx CellSelect Software (Optional)

If using cells which were imaged in the previous step, manually inspect the selected nanowells in CellSelect Software to exclude or include one or more candidate wells. Please refer to the <u>ICELL8 ex</u> <u>CellSelect v2.5 Software User Manual</u>, Section III, for more information about this process.

NOTE: If an incorrect barcode file was selected during Dispense, make sure to reference Section III.C ("Load a Different Barcode File") of the <u>ICELL8 cx CellSelect v2.5 Software User Manual</u>.

E. Protocol: Full-Length cDNA Synthesis by RT-PCR (2nd Dispense)

First-strand cDNA is synthesized by the oligo dT primer (SMART-Seq ICELL8 CDS) and followed by the template switching oligo (SMART-Seq ICELL8 Oligonucleotide) for template switching at the 5' end of transcripts. The first-strand cDNA is used as template for 2nd-strand cDNA synthesis and amplification during PCR.

1. Before you start

- If needed, perform a once-a-day warmup on the ICELL8 cx instrument.
- Set the centrifuge(s) used for spinning the chip and 384-Well Source Plate to 4°C.
- Preprogram the thermal cycler with Preheat (Step 1) and RT-PCR (Step 19) before the experiments.
- Run and hold the programs before the reactions.
- This protocol requires a 384-Well Source Plate (with seal), a Plate Seal Applicator, a film sealing roller, and the following components: Nuclease-Free Water (white cap), SMART-Seq ICELL8 RT-PCR Buffer (red cap), SMART-Seq ICELL8 Oligonucleotide (orange cap), RNase Inhibitor (white cap), Terra PCR Direct Polymerase Mix (pink cap), and SMARTScribe Reverse Transcriptase (purple cap)—all from the SMART-Seq ICELL8 Reagent Kit—plus Blotting Paper and RC Film (from the ICELL8 Loading Kit B).
- Thaw all reagents on ice except for the enzymes. Remove the enzymes from -20°C storage just prior to use and keep them on ice at all times. Gently mix and spin down all thawed reagents and enzymes.

2. Procedure

- 1. If necessary, remove the ICELL8 Chip Holder containing the ICELL8 350v Chip from the -80°C freezer. Thaw the chip in the chip holder until it reaches room temperature (about 10 min) to lyse cells.
- 2. Take the chip out of the chip holder, use a Kimwipe to dry any liquid on the chip surface, and then centrifuge the chip at 3,220g (minimum 2,600g) for 3 min at 4°C. Keep the chip on ice until ready for Preheat.
- 3. Run and hold the Preheat program below. Place the ICELL8 350v Chip into the thermal cycler that has been preheated to 76°C and a heated-lid temperature of 72°C. Resume the following Preheat program:

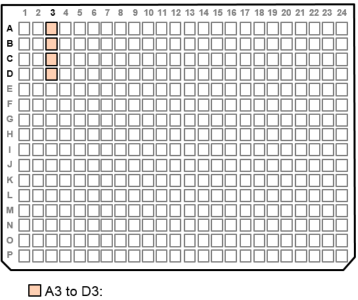
76°C	5 sec
72°C	3 min

- 4. After the 3 min incubation at 72°C, immediately place the chip on an ICELL8 cx Cold Block on ice for at least 2 min.
- 5. Centrifuge the chip at 3,220g (minimum 2,600g) for 3 min at 4°C, and keep the chip on the cold block on ice until ready for the RT-PCR dispense.
- 6. Run and hold the program for RT-PCR.
- 7. Mix all components except for SMARTScribe Reverse Transcriptase in the order listed in Step 7, and then vortex it briefly.

8. Add the SMARTScribe Reverse Transcriptase to the Master Mix immediately prior to use. Mix it by vortexing gently.

IMPORTANT: Remove the reverse transcriptase and PCR polymerase from the freezer, gently tap the tubes to mix, and add to the RT-PCR reaction mix. Mix by gently vortexing for 1–2 seconds and spin the tube briefly in a minicentrifuge to collect contents.

- 83.6 µl Nuclease-Free Water (white)
- 105.6 µl SMART-Seq ICELL8 RT-PCR Buffer (red)
 - 8.8 µl SMART-Seq ICELL8 Oligonucleotide (orange)
 - 4.4 µl RNase Inhibitor (white)
 - 8.8 µl Terra PCR Direct Polymerase Mix (pink)
 - 8.8 µl SMARTScribe Reverse Transcriptase (200 U/µl; purple)
- 220.0 µl Total volume
- 9. Bring the RT-PCR Master Mix to room temperature and pipette 50 µl of RT-PCR Master Mix into the 384-plate source wells (A3, B3, C3, and D3) highlighted in Figure 13, below.



add 50 μl of RT-PCR Master Mix per well

Figure 13. Aliquoting 50 µl of RT-PCR Master Mix into each of the orange source wells.

10. Seal the 384-Well Source Plate with a 384-Well Source Plate Seal using the Plate Seal Applicator.

- 11. Refer to the <u>ICELL8 cx Single-Cell System User Manual</u>, Section X.F "Dispense Reagents and/or Indexes into the Chip" for detailed information about this protocol, with the following guidelines:
 - a. Start in Section X.F, step 3, with loading the source plate into the plate nest.
 - b. At step 6, click the [Dispense RT PCR mix] button (Figure 14).

Startup Advanced 3'DE - TCR SMART-Seq
72 x 72 : 350 nl Chip ID
234914
Dispense Cells and Controls (50 nl)
Scan chip
Dispense RT - PCR mix (50 nl filtered)
Dispense Index 1 (50 nl filtered)
Dispense Tagmentation mix (50 nl filtered)
Dispense Index 2 (50 nl filtered)

Figure 14. Using CELLSTUDIO software to dispense the RT - PCR mix.

- c. Follow steps 7–13 as written in Section X.F.
- d. For step 14, centrifuge the sealed chip at 3,220g (minimum 2,600g) for 3 min at 4°C.
- e. Do step 15.
- 12. Place the chip into the thermal cycler with a heated-lid temperature of 72°C and the RT-PCR program held previously, at Step 5.

13. Resume the RT-PCR program:

42.3°C 36.6°C	5 sec 3 hr
99°C	9 sec
95.5°C	1 min
14 cycles*:	
100°C	5 sec
99°C	7 sec
59°C	5 sec
64°C	30 sec
69.5°C	5 sec
67.5°C	6 min
4°C	forever

*See Appendix B for cycling guidelines.

SAFE STOPPING POINT: The chip can be left in the thermal cycler at 4°C overnight.

F. Protocol: Forward Indexing Primers (3rd Dispense)

72 forward indexing primers are dispensed from the prealiquoted 384-well plate (stored at -70° C), which is used for Library PCR 1.

1. Before you start

- If needed, perform a once-a-day warmup -or- make sure that the [Tip Clean] procedure was done 3–4 times after the last dispense.
- Set the centrifuge(s) used for spinning the chip and 384-Well Source Plate to 4°C.
- This protocol requires the 384-well plate containing 72 forward indexing primers (stored at 70°C), a Plate Seal Applicator, a film sealing roller, and the following components from the ICELL8 Loading Kit B: Blotting Paper and RC Film.
- Prepare both 384-well plates containing 72 forward indexing primers and 72 reverse indexing primers (for plate maps, see Figures 15 and 32, respectively):

IMPORTANT: Steps a–e should be performed before you begin this protocol.

- a. Take the 384-well plates out of the -80°C freezer and seal them using the Plate Seal Applicator immediately before thawing the plates. (DO NOT open the sealing foil.)
- b. Without opening the sealing foil, thaw the frozen indexes at room temperature.
- c. Centrifuge the plates at 3,220g (minimum 2,600g) for 3 min between 4°C and room temperature.
- d. Shake the plates with a 384-well plate shaker for 1 min at room temperature.

e. Centrifuge the plates at 3,220g (minimum 2,600g) for 3 min between 4°C and room temperature. Keep the plate containing forward indexes at room temperature for use in this protocol, and store the plate containing reverse indexes at 4°C until Section V.H. If centrifuging or stored at 4°C, bring the plate back to room temperature prior to the dispense.

IMPORTANT: If you observe that the sealing foil is not securely sealed, please contact TBUSA Technical Support at <u>technical support@takarabio</u>.com or 800.662.2566.

	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24
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A5 to P8, A9 to D9, and A10 to D10: each well contains 20 μl of forward index

Figure 15. SMART-Seq ICELL8 forward indexing primers. The 72 forward indexes shown are prealiquoted into a 384-well plate (20 µl per well). See Appendix A for barcode sequences.

2. Procedure

- 1. Remove the chip from the thermal cycler (Section V.E, Step 14). Centrifuge the chip at 3,220g for 3 min at 4°C.
- 2. Remove the plate seal and place the 384-well plate containing forward indexing primers in the ICELL8 cx Single-Cell System with the A1 corner positioned at the top-right corner of the plate nest (see Figures 3 and 4).
- 3. Gently remove the RC Film from the chip and insert the chip into the chip nest.
- 4. Refer to the <u>ICELL8 cx Single-Cell System User Manual</u>, Section X.F "Dispense Reagents and/or Indexes into the Chip", with the following guidelines:
 - Start in Section X.F, step 5, by closing the front of the instrument.
 - At step 6, click the [Dispense Index 1] button (Figure 16).

Startup Advanced 3'DE - TCR SMART-Seq	
72 x 72 : 350 nl	
Chip ID 234914	
Dispense Cells and Controls (50 nl)	
Scan chip	
Dispense RT - PCR mix (50 nl filtered)	
Dispense Index 1 (50 nl filtered)	
Dispense Tagmentation mix (50 nl filtered)	
Dispense Index 2 (50 nl filtered)	

Figure 16. Using CELLSTUDIO software to dispense Index 1.

- Follow steps 7–13 as written in Section X.F.
- For step 14, centrifuge the sealed chip at 3,220g (minimum 2,600g) for 3 min at 4°C.
- Do step 15.

G. Protocol: Tagmentation (4th Dispense)

Synthesized full-length cDNAs are tagmented by Tagment DNA Enzyme (TDE1).

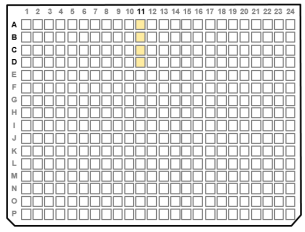
- 1. Before you start
 - Preprogram the thermal cycler with the Tagmentation program (Step 15) before the experiment.
 - Run and hold the program before the reaction.
 - This protocol requires a 384-Well Source Plate, a 384-Well Source Plate Seal, a Plate Seal Applicator, a film sealing roller, TDE1, and the following components: Nuclease-Free Water (white cap), Terra PCR Direct Polymerase Mix (pink cap), TRH (white cap), MgCl₂ (white cap) all from the SMART-Seq ICELL8 Reagent Kit—Blotting Paper, and RC Film (from ICELL8 Loading Kit – B).
 - Thaw Nuclease-Free Water and MgCl₂ and remove the enzymes from -20°C storage just prior to use. Keep them on ice at all times. Gently mix and spin down the thawed MgCl₂ and enzymes.

2. Procedure

1. Make the Tagmentation Master Mix by mixing all components in the order listed in the table below. Vortex briefly.

52.80 µl	Nuclease-Free Water (white)
139.04 µl	MgCl ₂ (white)
17.60 µl	TDE1
8.80 µl	Terra PCR Direct Polymerase Mix (pink)
1.76 µl	TRH (white)
220.0 µl	Total volume

2. Bring the Master Mix to room temperature and pipette 50 μl into the 384-plate source wells (A11, B11, C11, and D11) as highlighted in Figure 17, below.



A11 to D11: add 50 μl of Tagmentation Master Mix per well

Figure 17. Adding Tagmentation Master Mix to the source plate wells shown in yellow.

- 3. Seal the 384-Well Source Plate with the 384-Well Source Plate Seal using the Plate Seal Applicator.
- 4. Centrifuge the source plate at 3,220g (minimum 2,600g) for 3 min at 4°C.
- 5. Gently remove the RC Film from the chip and insert the chip into the chip nest.
- 6. Refer to the <u>ICELL8 cx Single-Cell System User Manual</u>, Section X.F "Dispense Reagents and/or Indexes into the Chip", with the following guidelines:
 - Start in Section X.F, step 5, by closing the front of the instrument.
 - At step 6, click the [Dispense Tagmentation mix] button (Figure 18).

Startup Advanced 3'DE - TCR SMART-Seq				
72 x 72 : 350 nl				
Chip ID 234914				
Dispense Cells and Controls (50 nl)				
Scan chip				
Dispense RT - PCR mix (50 nl filtered)				
Dispense Index 1 (50 nl filtered)				
Dispense Tagmentation mix (50 nl filtered)				
Dispense Index 2 (50 nl filtered)				

Figure 18. Using CELLSTUDIO software to dispense the Tagmentation mix.

- Follow steps 7–13 as written in Section X.F.
- For step 14, centrifuge the sealed chip at 3,220g (minimum 2,600g) for 3 min at 4°C.
- Do step 15.
- 7. Place the chip into the thermal cycler with a heated-lid temperature of 40°C and the tagmentation program held at the beginning of this protocol.
- 8. Resume the Tagmentation program:

42.3°C	4 sec
36.6°C	30 min
4°C	forever

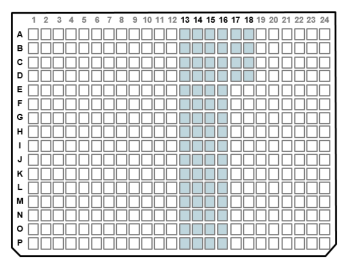
H. Protocol: Reverse Indexing Primers (5th Dispense) and Library PCR 1

Reverse indexing primers are dispensed from the prealiquoted 384-well plate (stored at -70° C), which is used for Library PCR 1.

1. Before you start

- Set the centrifuge(s) used for spinning the chip and 384-well index plate to 4°C.
- Preprogram the thermal cycler with the Library PCR 1 program (Step 12) before the experiment.
- Run and hold the program before the reaction.
- This protocol requires the 384-well plate containing 72 reverse indexing primers (stored at -70°C) and the following components from the ICELL8 Loading Kit B: Blotting Paper and RC Film.

• Bring to room temperature the 384-well plate containing 72 reverse indexing primers (prepared in Section V.F; for plate map, see Figure 19).



A13 to P16, A17 to D17, and A18 to D18: each well contains 20 μl of reverse index

Figure 19. SMART-Seq ICELL8 reverse indexing primers. The 72 reverse indexes shown in blue are prealiquoted into a 384-well plate (20 μ l per well). See Appendix A for barcode sequences.

2. Procedure

- 1. Remove the chip from the thermal cycler (Section V.G, Step 8). Centrifuge the chip at 3,220g for 3 min at 4°C.
- 2. Remove the seal and place the 384-well plate containing reverse indexing primers in the ICELL8 cx Single-Cell System with the A1 corner positioned at the top-right corner of the plate nest. The beveled corners of the 384-well index plate should be on the left side (see Figures 3 and 4).
- 9. Gently remove the RC Film from the chip and insert the chip into the chip nest.
- 10. Refer to the <u>ICELL8 cx Single-Cell System User Manual</u>, Section X.F "Dispense Reagents and/or Indexes into the Chip", with the following guidelines:
 - Start in Section X.F, step 5, by closing the front of the instrument.
 - At step 6, click the [Dispense Index 1] button (Figure 20).

Startup Advanced 3'DE - TCR SMART-Seq				
72 x 72 : 350 nl				
Chip ID 234914				
Dispense Cells and Controls (50 nl)				
Scan chip				
Dispense RT - PCR mix (50 nl filtered)				
Dispense Index 1 (50 nl filtered)				
Dispense Tagmentation mix (50 nl filtered)				
Dispense Index 2 (50 nl filtered)				

Figure 20. Using CELLSTUDIO software to dispense Index 2.

- Follow steps 7–13 as written in Section X.F.
- For step 14, centrifuge the sealed chip at 3,220g (minimum 2,600g) for 3 min at 4°C.
- Do step 15.
- 3. Place the chip into the thermal cycler with a heated-lid temperature of 72°C to perform Library PCR 1.
- 4. Resume the Library PCR 1 program:

77°C 72°C 99°C 95.5°C	12 sec 3 min 11 sec 1 min
6 cycles:	
100°C	20 sec
99°C	10 sec
53.3°C	5 sec
58°C	15 sec
71°C	5 sec
67.5°C	2 min
4°C	forever

SAFE STOPPING POINT: The chip can be left in the thermal cycler at 4°C overnight.

I. Protocol: Library Extraction from the Chip

This protocol extracts the library amplified by Library PCR 1 from the chip.

Refer to the <u>ICELL8 cx Single-Cell System User Manual</u>, Section XI "Protocol: Extract library from the chip" for the procedure. The collected volume should be no less than 60% of the Maximum Potential Volume.

SAFE STOPPING POINT: The eluate can be frozen at -20° C.

J. Protocol: Purification of the Extracted Library

In this protocol, the extracted library is purified twice using a 1:1 proportion of AMPure XP beads.

1. Before you start

• Equilibrate an aliquot of AMPure XP beads to room temperature for ~30 min prior to use.

NOTE: Immediately prior to use, vortex the beads until they are well dispersed. The color of the liquid should appear homogeneous. Confirm that there is no remaining pellet of beads at the bottom of the tube. Mix well to disperse before adding the beads to your reactions. The beads are viscous, so pipette them slowly.

- Prepare fresh 80% (v/v) ethanol before each cleanup (1 ml each). Use an anhydrous ethanol stock.
- This protocol requires Elution Buffer from the SMART-Seq ICELL8 Reagent Kit.
- Use only half of the volume of the extracted library in the procedure below. Store the remaining half volume in a freezer in case further analysis is needed.

2. Procedure

1. Transfer half of the volume of the extracted library to 0.2-ml PCR tube(s) and store the remaining half in a freezer.

NOTE: Add no more than 100 μ l of library per tube. If you have more than 100 μ l, split it into multiple tubes of less than 100 μ l each.

- 2. Add 1:1 volume of well-vortexed AMPure XP Beads. (For example, add 100 µl of AMPure XP beads to 100 µl of the extracted library.)
- 3. Vortex the tube to mix well.
- 4. Incubate the tube at room temperature for 8 min to let the DNA bind to the beads.
- 5. Place the tube on the SMARTer-Seq Magnetic Separator PCR Strip for ~5 min or longer, until the liquid appears completely clear, and there are no beads left in the supernatant.

NOTE: During the ~5 min incubation, if there are beads not against the magnet, use the supernatant to resuspend them and pipet them onto the magnet with the rest of the beads.

6. While the tube is sitting on the magnetic separator, pipette out the supernatant.

- 7. Keep the tube on the magnetic separator. Add 200 µl of freshly made 80% ethanol to each sample without disturbing the beads. Wait for 30 seconds and carefully pipette out the supernatant containing contaminants. DNA will remain bound to the beads during the washing process.
- 8. Repeat Step 7 one more time.
- 9. Spin down the tube briefly to collect the liquid at the bottom of the well.
- 10. Place the tube on the magnetic separator for 30 seconds, and then remove all remaining ethanol by using pipette.

NOTE: It is important to make sure all ethanol is removed so the beads elute well and recovery is efficient.

11. Leave the tube at room temperature for ~5 min or longer until the pellet appears dry.

NOTE: If the beads are overdried, there will be cracks in the pellet. If this occurs, the DNA may not elute well from the beads and recovery may be reduced.

- 12. Once the beads are dried, add 50 μ l of Elution Buffer to cover the beads, with the following guidelines:
 - If using a single tube, add all 50 ul of Elution Buffer to the tube and proceed to step 13.
 - If the sample was split into multiple tubes at the beginning of this protocol, use a 50 μ l aliquot of Elution Buffer to elute all library tubes. For example, if you have 150 ul library and had split it into two tubes of 75 ul, add 25 ul of Elution Buffer to each tube.
- 13. Incubate the PCR tube(s) with the beads and Elution Buffer at room temperature for 2 min to rehydrate.
- 14. Mix the pellet by pipetting up and down 10 times to elute DNA from the beads, then put the tube back on the magnetic separator for 1 minute or longer until the solution is completely clear.

NOTE: During the incubation on the magnet, there may be a small population of beads not pelleting against the magnet. Use the supernatant to resuspend them by pipetting up and down, and pipet them onto the magnet with the rest of the beads. Continue incubation until there are no beads left in the supernatant.

- 15. Transfer clear supernatant containing purified library to a fresh 0.2-ml PCR tube. If the extracted library was split at the beginning of this protocol, pool the supernatant of all extraction tubes into a single tube.
- 16. Add 50 μ l of AMPure XP beads to the purified library.
- 17. Repeat Step 3–14 using fresh 80% ethanol at Step 7 and 14 μ l of Elution Buffer at Step 12.

K. Protocol: Library Amplification (Library PCR 2)

This PCR amplifies and yields the final sequencing-ready library.

1. Before you start

- Preprogram the PCR thermal cycler with the Library PCR 2 program (Step 3) before the experiment.
- Run and hold the program before the reaction.
- This protocol requires the following components from the SMART-Seq ICELL8 Reagent Kit: Terra PCR Direct Polymerase Mix (pink cap), 5X Primer Mix (green cap), and SeqAmp CB PCR Buffer (clear cap).

2. Procedure

- 1. Take out all reagents out from the freezer and thaw the 5X Primer Mix. Gently mix each reagent tube and spin down briefly. Store on ice.
- 2. Add all reagents in the order below to the purified library after Library PCR 1 (Section V.J, Step 16):

14 µl	Purified library after Library PCR 1
25 µl	SeqAmp CB PCR Buffer (clear)
10 µl	5X Primer Mix (green)
1 µl	Terra PCR Direct Polymerase Mix (pink)
50 µl	Total volume

3. Place the tube in a preheated PCR thermal cycler with a heated lid and run the following Library PCR 2 program:

98°(2 min		
7 cycles:			
	98°C	10 sec	
	60°C	15 sec	
	68°C	2 min	
4°(4°C		

SAFE STOPPING POINT: The tube can be stored at 4°C overnight.

L. Protocol: Purification of the Sequencing-Ready Library

In this protocol, the sequencing-ready library is purified using a 1:1 proportion of AMPure XP beads.

1. Before you start

• Equilibrate an aliquot of AMPure XP beads to room temperature for ~30 min prior to use.

NOTE: Immediately prior to use, vortex the beads until they are well dispersed. The color of the liquid should appear homogeneous. Confirm that there is no remaining pellet of beads at the bottom of the tube. Mix well to disperse before adding the beads to your reactions. The beads are viscous, so pipette them slowly.

- Prepare at least 400 μ l of fresh 80% (v/v) ethanol from a stock of anhydrous ethanol.
- This protocol requires Elution Buffer provided with the SMART-Seq ICELL8 Reagent Kit.

2. Procedure

- 1. Remove the PCR tube from the PCR thermal cycler (Section V.K, Step 3). Spin it down briefly.
- 2. Add 50 μ l (1:1 v/v) of well-vortexed AMPure XP beads to the tube.
- 3. Vortex the tube to mix well.
- 4. Incubate the tube at room temperature for 8 min to let the DNA bind to the beads.
- 5. Place the tube on Magnetic Stand for ~5 min or longer, until the liquid appears completely clear, and there are no beads left in the supernatant.

NOTE: During the ~5-minute incubation, if there are beads not against the magnet, use the supernatant to resuspend them and pipet them onto the magnet with the rest of the beads.

- 6. While the tube is sitting on the magnetic stand, pipette out the supernatant.
- 7. Keep the tube on the magnetic stand. Add 200 µl of freshly made 80% ethanol to each sample without disturbing the beads. Wait for 30 seconds and carefully pipette out the supernatant containing contaminants. DNA will remain bound to the beads during the washing process.
- 8. Repeat Step 7 one more time.
- 9. Spin down the tube briefly to collect the liquid at the bottom of the well.
- 10. Place the tube on the magnetic stand for 30 seconds, and then remove all remaining ethanol using a pipette.

NOTE: It is important to make sure all ethanol is removed so the beads elute well and recovery is efficient.

11. Leave the tube at room temperature for ~ 5 min or longer until the pellet appears dry.

NOTE: If beads were overdried, there will be cracks in the pellet. If this occurs, the DNA may not elute well from the beads and recovery may be reduced.

- 12. Once the beads are dried, add 17 μ l Elution Buffer to cover the beads and incubate it at room temperature for 2 min to rehydrate.
- 13. Mix the pellet by pipetting up and down 10 times to elute DNA from the beads, then put the tube back on the magnetic stand for 1 minute or longer until the solution is completely clear.

NOTE: During the incubation on the magnet, there may be a small population of beads not pelleting against the magnet. Use the supernatant to resuspend them by pipetting up and down, and pipet them onto the magnet with the rest of the beads. Continue incubation until there are no beads left in the supernatant.

14. Transfer clear supernatant containing purified library to a clean 1.5-ml PCR tube.

SAFE STOPPING POINT: The samples may be stored at –20°C.

M. Protocol: Library Validation and Quantification

To determine whether library production and purification were successful, we recommend analyzing and quantifying the final libraries using the Agilent 2100 Bioanalyzer and the High Sensitivity DNA Kit (Agilent, Cat. No. 5067-4626). Alternatively, the libraries can be quantified by qPCR using Takara Bio's NGS Library Quantification Kit (Cat. No. 638324). Please refer to the corresponding user manuals for detailed instructions.

Procedure

- Measure the concentration of the purified sequencing-ready library using 1 μl of the library, a Qubit fluorometer, and the Qubit dsDNA HS Assay Kit (Thermo Fisher Scientific, Cat. No. Q32851). Refer to the Qubit dsDNA HS assay kit user manual for sample prep instructions.
- 2. Based on the Qubit measurement, dilute the amplicon to 0.2 to $2.0 \text{ ng/}\mu\text{l}$ of library.

NOTE: We recommend using several concentrations for the Bioanalyzer (e.g., 0.2, 0.5, 1, and 2 ng/ μ l).

 Use 1 µl of each concentration to load the Agilent 2100 Bioanalyzer and the High Sensitivity DNA Chip from Agilent's High Sensitivity DNA Kit for validation. See the user manual for the Agilent High Sensitivity DNA Kit for instructions.

NOTE: Be careful not to transfer beads with your sample.

- 4. Use the Bioanalyzer results to determine library quality and average size. See Figure 21, below, for an example of a typical Bioanalyzer profile for an NGS library that has been successfully purified and size selected.
 - For libraries <850 base pairs as determined by the Bioanalyzer, use the Qubit concentration in combination with the average size of the library to calculate the molar library concentration.
 - For libraries >850 base pairs as determined by the Bioanalyzer, use qPCR to accurately measure the concentration of the sequencing library and ensure optimal loading on the sequencer. Use the Library Quantification Kit (Cat. No. 638324). Refer to the "Library Quantification Kit User Manual" on the <u>takarabio.com</u> website for instructions and use the average size as determined by the Bioanalyzer to calculate the molar library concentration.
- 5. Store the sequencing library at -20°C until ready for sequencing. Please review Appendix C for sequencing guidelines.
- 6. Please review Appendix D for data analysis guidelines.

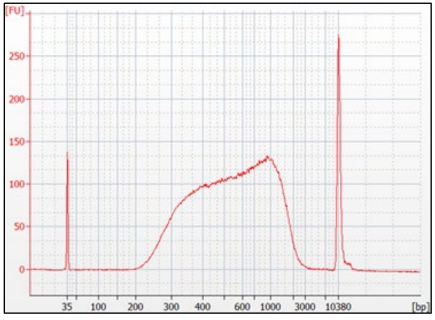


Figure 21. An electropherogram example from a library prepared from K-562 nuclei and quantified using the Agilent 2100 Bioanalyzer.

Appendix A: Forward and Reverse Indexing Primers

A shortened name is used for simplicity instead of the full name of each index. For example, "Forward Index 1" below stands for "SMART-Seq ICELL8 Forward Index 1". Sample sheets will be required if Illumina's bcl2fastq Conversion Software is used for demultiplexing the sequencing reads. Please refer to Appendix D for details.

Table 3. SMART-Seq ICELL8 Indexing Primer Set – A and SMART-Seq ICELL8 Indexing Primer Set – B forward indexing primers 1–72.

Forward indexing (i5) primers			Forward indexes (i5) on sample sheet	
Index number	Well position	Index sequence in primers	MiSeq®, HiSeq® 2000/2500, NovaSeq™	MiniSeq™, NextSeq®, HiSeq 3000/4000, HiSeq X
Forward Index 1	A5	AACCAACG	AACCAACG	CGTTGGTT
Forward Index 2	B5	AACGATAG	AACGATAG	CTATCGTT
Forward Index 3	C5	AAGAAGAC	AAGAAGAC	GTCTTCTT
Forward Index 4	D5	AGAGCCTA	AGAGCCTA	TAGGCTCT
Forward Index 5	E5	ATAGTCAA	ATAGTCAA	TTGACTAT
Forward Index 6	F5	CAACTGCA	CAACTGCA	TGCAGTTG
Forward Index 7	G5	CAGCATGA	CAGCATGA	TCATGCTG
Forward Index 8	H5	CCGCCTAA	CCGCCTAA	TTAGGCGG
Forward Index 9	15	CCTAGCGA	CCTAGCGA	TCGCTAGG
Forward Index 10	J5	CGCAACGG	CGCAACGG	CCGTTGCG
Forward Index 11	K5	CTTGGCCT	CTTGGCCT	AGGCCAAG
Forward Index 12	L5	GCGGTTCT	GCGGTTCT	AGAACCGC
Forward Index 13	M5	GCTTGATG	GCTTGATG	CATCAAGC
Forward Index 14	N5	GGCTCTCT	GGCTCTCT	AGAGAGCC
Forward Index 15	O5	TCAATGCT	TCAATGCT	AGCATTGA
Forward Index 16	P5	TGGTAATT	TGGTAATT	AATTACCA
Forward Index 17	A6	TTCTGAAC	TTCTGAAC	GTTCAGAA
Forward Index 18	B6	AACCAGAA	AACCAGAA	TTCTGGTT
Forward Index 19	C6	AACCGAAC	AACCGAAC	GTTCGGTT
Forward Index 20	D6	AACCGCCA	AACCGCCA	TGGCGGTT
Forward Index 21	E6	AATAAGGA	AATAAGGA	TCCTTATT
Forward Index 22	F6	ACCTTATT	ACCTTATT	AATAAGGT
Forward Index 23	G6	TGGTCCTG	TGGTCCTG	CAGGACCA
Forward Index 24	H6	CAACGAGG	CAACGAGG	CCTCGTTG
Forward Index 25	16	CCAATGGA	CCAATGGA	TCCATTGG
Forward Index 26	J6	CGCCTATG	CGCCTATG	CATAGGCG
Forward Index 27	K6	CTCTCCAA	CTCTCCAA	TTGGAGAG
Forward Index 28	L6	GGCTTGAA	GGCTTGAA	TTCAAGCC
Forward Index 29	M6	GTTAAGTT	GTTAAGTT	AACTTAAC
Forward Index 30	N6	TCAAGTAT	TCAAGTAT	ATACTTGA
Forward Index 31	O6	TCGCGGAT	TCGCGGAT	ATCCGCGA
Forward Index 32	P6	TGAGTCCT	TGAGTCCT	AGGACTCA
Forward Index 33	A7	AGTTGAAC	AGTTGAAC	GTTCAACT
Forward Index 34	B7	AACCTCAG	AACCTCAG	CTGAGGTT
Forward Index 35	C7	AACGGTCT	AACGGTCT	AGACCGTT

Forward Index 36D7AACTCAAGAACTCCAAGCTTGAGTTForward Index 37E7AACTCCGAAACTCCGATCGAGTTForward Index 38F7AAGGTTCAAAAGGTTCATGAACCTTForward Index 39G7AATTCGGTAATTCGGTACCGAAATTForward Index 40H7ACCAGACCACCAGACCGGTCTGGTForward Index 4117ACTTAGTAACTTAGTATACTAAGTForward Index 42J7ACCGGCAAAGGCGCAATTCGACCTForward Index 43K7AGGTCGAAAGGTCGAATTCGACCTForward Index 44L7AGTCTGGAAGGTCGAATCCAGACTForward Index 44K7AGGTCGAAAGGCGCAATCCAGACTForward Index 44N7ATCATTGATCATGGAATTCGGATForward Index 44N7ATCATTGATCATTGCAGATTGForward Index 46N7ATCATTGATGAATCTAGATTCATForward Index 48P7CAAGATTGCCGAATTGCAATTCGGForward Index 49A8CCGAATTGCCGAATTGCAATTCGGForward Index 50B8CCGAGTTCGGAGTTAATTCCGGForward Index 51C8CGTAAGGCCTCAGGCTTAAGGForward Index 53E8CGAAGACTCGGAGCTCGAGCTCCGForward Index 54F8CGTAGGCAGGCTAAGGCTTAGTCForward Index 55G8GAACTAAGGACTAAGGCTTAGTCForward Index 56H8GACTAAGGGATATGGCAATAGCForward Index 57I					
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Forward Index 48P7CAAGATTGCAAGATTGCAAGATTGForward Index 49A8CCGAATTGCCGAATTGCAATCTGGForward Index 50B8CCGGAGTTCCGGAGTTAACTCCGGForward Index 51C8CCTTCAGGCCTTCAGGCCTGAAGGForward Index 52D8CGAATATTCGAATATTAATATTCGForward Index 53E8CGGAGACTCGGAGACTAGTCTCCGForward Index 54F8CGTAGGCACGTAGGCATGCCTACGForward Index 55G8GAACTAAGGAACTAAGCTTAGTTCForward Index 56H8GAAGCTCGGAAGCTCGCGAGCTTCForward Index 57I8GACTATTGGACTATTGCAATAGTCForward Index 58J8GAGTAACGGCTCAAGGCCTTAGCGForward Index 59K8GCAGTCCAGCAGTCCATGGACTGCForward Index 60L8GCTCAAGGGCTCAAGGCCTTGAGCForward Index 61M8GGATATCGGGATATCGCGATATCCForward Index 62N8GTCAGATGTCAGATATCTGACCForward Index 63O8GTAAGAAGGTAAGAAGCTTCTACForward Index 64P8GTAGAAGTGTACTCTATAGATACForward Index 65A9GTCATCTAGTCAGCAATGCGGACForward Index 66B9GTCATCTAGTCAGCAATGCGGACForward Index 67C9GTCCGCAAGTCAGCAATGCGGACForward Index 68D9GTTCAATAGTTCAATATATTGAACForward Index 69	Forward Index 46	N7	ATCCATTG	ATCCATTG	CAATGGAT
Forward Index 49A8CCGAATTGCCGAATTGCAATTCGGForward Index 50B8CCGGAGTTCCGGAGTTAACTCCGGForward Index 51C8CCTTCAGGCCTTCAGGCCTGAAGGForward Index 52D8CGAATATTCGAATATTAATATTCGForward Index 53E8CGGAGACTCGGAGACTAGTCTCCGForward Index 53E8CGAAGGCACGTAGGCATGCCTACGForward Index 54F8CGTAGGCACGTAGGCATGCCTACGForward Index 55G8GAACTAAGGAACTAAGCTTAGTTCForward Index 56H8GAAGCTCGGAAGCTCGCGAGCTTCForward Index 5718GACTATTGGACTATTGCAATAGTCForward Index 5718GACTACGCGTTACTCForward Index 58J8GAGTAACGGAGTAACGGCTCAAGGCCTTGAGCForward Index 59K8GCAGTCCATGGACTGCForward Index 59K8GCAGTCCAGCAGTATCCForward Index 60L8GCTCAAGGCCTTGAGCForward Index 61M8GGATATCGGGATATCGCGATATCCForward Index 62N8GTCAGATGTCAGATATCTGACCForward Index 63O8GTAAGAAGGTAAGAAGCTTCTTACForward Index 64P8GTAGAAGTGTACTGATCAGATACForward Index 65A9GTATCTGAGTCAGCAATGCGGACForward Index 66B9GTCATCTAGTAGAAGTACTTCTACForward Index 67C9GTCCGCAAGTCAGCGCGACGTTA	Forward Index 47	07	ATGAATCT	ATGAATCT	AGATTCAT
Forward Index 50B8CCGGAGTTCCGGAGTTAACTCCGGForward Index 51C8CCTTCAGGCCTGAAGGForward Index 52D8CGAATATTCGAATATTAATATTCGForward Index 53E8CGGAGACTCGGAGACTAGTCTCCGForward Index 53E8CGGAGCACGTAGGCATGCCTACGForward Index 54F8CGTAGGCACGTAGGCATGCCTACGForward Index 55G8GAACTAAGGAACTAAGCTTAGTTCForward Index 56H8GAAGCTCGGAAGCTCGCGAGCTTCForward Index 5718GACTATTGGACTATTGCAATAGTCForward Index 58J8GAGTAACGGCTTAAGTCGACTACCForward Index 59K8GCAGTCCAGCAGTCCATGGACTGCForward Index 60L8GCTCAAGGCCTTGAGCCTTCTTACCForward Index 61M8GGATATCGGGATATCCCGATATCCForward Index 62N8GTCAGATGTCAGATACTTCTACForward Index 63O8GTAAGAAGGTAAGAAGCTTCTTACForward Index 64P8GTAGAAGTGTACAGATACTTCTACForward Index 65A9GTATCTGAGTAGAAGTACTTCTACForward Index 66B9GTCATCTAGTCAGCAATGCGGACForward Index 66B9GTCATCTAGTCAGCAATGCGGACForward Index 67C9GTCCGCAAGTCAGCGCGACGTTAForward Index 68D9GTCATCTAGACGCCForward Index 69Forward Index 69A10 </td <td>Forward Index 48</td> <td>P7</td> <td>CAAGATTG</td> <td>CAAGATTG</td> <td>CAATCTTG</td>	Forward Index 48	P7	CAAGATTG	CAAGATTG	CAATCTTG
Forward Index 51C8CCTTCAGGCCTTCAGGCCTGAAGGForward Index 52D8CGAATATTCGAATATTAATATTCGForward Index 53E8CGGAGACTCGGAGACTAGTCTCCGForward Index 53E8CGTAGGCACGTAGGCATGCCTACGForward Index 54F8CGTAGGCACGTAGGCATGCCTACGForward Index 55G8GAACTAAGGAACTAAGCTTAGTTCForward Index 56H8GAAGCTCGGAAGCTCGCGAGCTTCForward Index 57I8GACTATTGGACTATTGCAATAGTCForward Index 58J8GAGTAACGGAGTAACGCGTTACTCForward Index 59K8GCAGTCCAGCAGTCCATGGACTGCForward Index 60L8GCTCAAGGCCTTGAGCForward Index 61Forward Index 61M8GGATATCGGGATATCGCGATATCCForward Index 63O8GTAAGAAGGTAAGAAGCTTCTACForward Index 64P8GTAGAAGTGTAGAAGTACTTCTACForward Index 66B9GTCATCTAGTCATCTATAGATGACForward Index 67C9GTCCGCAAGTCCGCAATTGCGGACForward Index 68D9GTTCAATAGTCAGTCAGACGTCAGATForward Index 69A10TAACGTCGTAACGTCGCGACGTTAForward Index 69A10TAACGTCGTAACGTCGCGACGTTAForward Index 70B10TCGGAACGCGTTCCGAForward Index 71C10Forward Index 71C10CATTCTACCATTCTACGTAGAAT	Forward Index 49	A8	CCGAATTG	CCGAATTG	CAATTCGG
Forward Index 52D8CGAATATTCGAATATTAATATTCGForward Index 53E8CGGAGACTCGGAGACTAGTCTCCGForward Index 54F8CGTAGGCACGTAGGCATGCCTACGForward Index 55G8GAACTAAGGAACTAAGCTTAGTTCForward Index 56H8GAACTCGGAACTATGCGAGCTTCForward Index 57I8GACTATTGGACTATTGCAATAGTCForward Index 58J8GAGTAACGGAGTAACGCGTTACTCForward Index 59K8GCAGTCCAGCAGTCCATGGACTGCForward Index 59K8GCAGTCCAGCAGTCCATGGACTGCForward Index 60L8GCTCAAGGGCTCAAGGCCTTGAGCForward Index 61M8GGATATCGGGATATCGCGATATCCForward Index 63O8GTAAGAAGGTAAGAAGCTTCTTACForward Index 64P8GTAGAAGTGTAGAAGTACTTCTACForward Index 65A9GTATCTGAGTCAGCAATGCGGACForward Index 66B9GTCATCTAGTCAGCAATGCGGACForward Index 67C9GTCCGCAAGTCCGCAATGCGGACForward Index 68D9GTTCAATAGTTCAATATATTGAACForward Index 69A10TAACGTCGTAACGTCGCGACGTTAForward Index 70B10TCGGAACGTCGGAACGCGTCCGAForward Index 71C10CATTCTACCATTCTACGTAGAATG	Forward Index 50	B8	CCGGAGTT	CCGGAGTT	AACTCCGG
Forward Index 53E8CGGAGACTCGGAGACTAGTCTCCGForward Index 54F8CGTAGGCACGTAGGCATGCCTACGForward Index 55G8GAACTAAGGAACTAAGCTTAGTTCForward Index 56H8GAAGCTCGGAAGCTCGCGAGCTTCForward Index 57I8GACTATTGGACTATTGCAATAGTCForward Index 58J8GAGTAACGGAGTAACGCGTTACTCForward Index 59K8GCAGTCCAGCAGTCCATGGACTGCForward Index 59K8GCAGTCCAGCAGTCCATGGACTGCForward Index 60L8GCTCAAGGCCTTGAGCForward Index 61M8GGATATCGGGATATCGCGATATCCForward Index 61M8GGTCAGATGTCAGATATCTGACCForward Index 63O8GTAAGAAGGTAAGAAGCTTCTTACForward Index 64P8GTAGAAGTGTACGAAGTACTTCTACForward Index 65A9GTCATCTAGTAAGAAGTTCAGATACForward Index 66B9GTCATCTAGTCATCTATAGATGACForward Index 67C9GTCCACAAGTCCGCAATTGCGGACForward Index 68D9GTTCAATAGTTCAATATATTGAACForward Index 69A10TAACGTCGTAACGTCGCGACGTTAForward Index 70B10TCGGAACGTCGGAACGCGTCCGAAForward Index 71C10CATTCTACCATTCTACGTAGAATG	Forward Index 51	C8	CCTTCAGG	CCTTCAGG	CCTGAAGG
Forward Index 54F8CGTAGGCACGTAGGCATGCCTACGForward Index 55G8GAACTAAGGAACTAAGCTTAGTTCForward Index 56H8GAAGCTCGGAAGCTCGCGAGCTTCForward Index 57I8GACTATTGGACTATTGCAATAGTCForward Index 58J8GAGTAACGGAGTAACGCGTTACTCForward Index 59K8GCAGTCCAGCAGTCCATGGACTGCForward Index 59K8GCAGTCCAGCAGTCCATGGACTGCForward Index 60L8GCTCAAGGGCTCAAGGCCTTGAGCForward Index 61M8GGATATCGGGATATCGCGATATCCForward Index 62N8GGTCAGATGTCAGATATCTGACCForward Index 63O8GTAAGAAGGTAAGAAGCTTCTTACForward Index 64P8GTAGAAGTGTATCTGATCAGATACForward Index 65A9GTATCTGAGTATCTGATCAGATACForward Index 66B9GTCATCTAGTCCGCAATTGCGGACForward Index 67C9GTCCGCAAGTCCGCAATTGCGGACForward Index 68D9GTTCAATAGTTCAATATATTGAACForward Index 69A10TAACGTCGTAACGTCGCGACGTTAForward Index 70B10TCGGAACGTCGGAACGCGTCCGAAForward Index 71C10CATTCTACCATTCTACGTAGAATG	Forward Index 52	D8	CGAATATT	CGAATATT	AATATTCG
Forward Index 55G8GAACTAAGGAACTAAGCTTAGTTCForward Index 56H8GAAGCTCGGAAGCTCGCGAGCTTCForward Index 57I8GACTATTGGACTATTGCAATAGTCForward Index 58J8GAGTAACGGAGTAACGCGTTACTCForward Index 59K8GCAGTCCAGCAGTCCATGGACTGCForward Index 60L8GCTCAAGGGCTCAAGGCCTTGAGCForward Index 61M8GGATATCGGGATATCGCGATATCCForward Index 62N8GGTCAGATGGTCAGATATCTGACCForward Index 63O8GTAAGAAGGTAAGAAGCTTCTTACForward Index 64P8GTAGAAGTGTACGAAGTACTTCTACForward Index 65A9GTATCTGAGTCATCTATAGATACForward Index 66B9GTCATCTAGTCATCTATAGATGACForward Index 67C9GTCCGCAAGTCCGCAATTGCGGACForward Index 68D9GTTCAATAGTTCAATATATTGAACForward Index 69A10TAACGTCGTAACGTCGCGACGTTAForward Index 70B10TCGGAACGTCGGAACGCGTTCCGAForward Index 71C10CATTCTACCATTCTACGTAGAATG	Forward Index 53	E8	CGGAGACT	CGGAGACT	AGTCTCCG
Forward Index 56H8GAAGCTCGGAAGCTCGCGAGCTTCForward Index 57I8GACTATTGGACTATTGCAATAGTCForward Index 58J8GAGTAACGGAGTAACGCGTTACTCForward Index 59K8GCAGTCCAGCAGTCCATGGACTGCForward Index 60L8GCTCAAGGGCTCAAGGCCTTGAGCForward Index 61M8GGATATCGGGATATCGCGATATCCForward Index 62N8GGTCAGATGGTCAGATATCTGACCForward Index 63O8GTAAGAAGGTAAGAAGCTTCTTACForward Index 64P8GTAGAAGTGTAGAAGTACTTCTACForward Index 65A9GTATCTGAGTAATCTGATCAGATACForward Index 66B9GTCATCTAGTCATCTATAGATGACForward Index 67C9GTCCGCAAGTCCGCAATTGCGGACForward Index 68D9GTTCAATAGTTCAATATATTGAACForward Index 69A10TAACGTCGTAACGTCGCGACGTTAForward Index 70B10TCGGAACGCGTTCCGAGTAGAATGForward Index 71C10CATTCTACCATTCTACGTAGAATG	Forward Index 54	F8	CGTAGGCA	CGTAGGCA	TGCCTACG
Forward Index 57I8GACTATTGGACTATTGCAATAGTCForward Index 58J8GAGTAACGGAGTAACGCGTTACTCForward Index 59K8GCAGTCCAGCAGTCCATGGACTGCForward Index 60L8GCTCAAGGGCTCAAGGCCTTGAGCForward Index 61M8GGATATCGGGATATCGCGATATCCForward Index 62N8GGTCAGATGGTCAGATATCTGACCForward Index 63O8GTAAGAAGGTAAGAAGCTTCTTACForward Index 64P8GTAGAAGTGTATCTGATCAGATACForward Index 65A9GTATCTGAGTATCTGATCAGATACForward Index 66B9GTCATCTAGTCATCTATAGATGACForward Index 67C9GTCCGCAAGTCCGCAATTGCGGACForward Index 68D9GTTCAATAGTTCAATATATTGAACForward Index 69A10TAACGTCGTAACGTCGCGACGTTAForward Index 70B10TCGGAACGTCGGAACGCGTTCCGAForward Index 71C10CATTCTACCATTCTACGTAGAATG	Forward Index 55	G8	GAACTAAG	GAACTAAG	CTTAGTTC
Forward Index 58J8GAGTAACGGAGTAACGCGTTACTCForward Index 59K8GCAGTCCAGCAGTCCATGGACTGCForward Index 60L8GCTCAAGGGCTCAAGGCCTTGAGCForward Index 61M8GGATATCGGGATATCGCGATATCCForward Index 62N8GGTCAGATGGTCAGATATCTGACCForward Index 63O8GTAAGAAGGTAAGAAGCTTCTTACForward Index 64P8GTAGAAGTGTACTAGAAGTACTTCTACForward Index 65A9GTATCTGAGTATCTGATCAGATACForward Index 66B9GTCATCTAGTCATCTATAGATGACForward Index 67C9GTCCGCAAGTCCGCAATTGCGGACForward Index 68D9GTTCAATAGTTCAATATATTGAACForward Index 69A10TAACGTCGTAACGTCGCGACGTTAForward Index 70B10TCGGAACGTCGGAACGCGTTCCGAForward Index 71C10CATTCTACCATTCTACGTAGAATG	Forward Index 56	H8	GAAGCTCG	GAAGCTCG	CGAGCTTC
Forward Index 59K8GCAGTCCAGCAGTCCATGGACTGCForward Index 60L8GCTCAAGGGCTCAAGGCCTTGAGCForward Index 61M8GGATATCGGGATATCGCGATATCCForward Index 62N8GGTCAGATGGTCAGATATCTGACCForward Index 63O8GTAAGAAGGTAAGAAGCTTCTTACForward Index 64P8GTAGAAGTGTAGAAGTACTTCTACForward Index 65A9GTATCTGAGTATCTGATCAGATACForward Index 66B9GTCATCTAGTCATCTATAGATGACForward Index 67C9GTCCGCAAGTCCGCAATTGCGGACForward Index 68D9GTTCAATAGTTCAATATATTGAACForward Index 69A10TAACGTCGTAACGTCGCGACGTTAForward Index 70B10TCGGAACGTCGGAACGCGTTCCGAForward Index 71C10CATTCTACCATTCTACGTAGAATG	Forward Index 57	18	GACTATTG	GACTATTG	CAATAGTC
Forward Index 60L8GCTCAAGGGCTCAAGGCCTTGAGCForward Index 61M8GGATATCGGGATATCGCGATATCCForward Index 62N8GGTCAGATGGTCAGATATCTGACCForward Index 63O8GTAAGAAGGTAAGAAGCTTCTTACForward Index 64P8GTAGAAGTGTAGAAGTACTTCTACForward Index 65A9GTATCTGAGTATCTGATCAGATACForward Index 66B9GTCATCTAGTCATCTATAGATGACForward Index 67C9GTCCGCAAGTCCGCAATTGCGGACForward Index 68D9GTTCAATAGTTCAATATATTGAACForward Index 69A10TAACGTCGTAACGTCGCGACGTTAForward Index 70B10TCGGAACGTCGGAACGCGTTCCGAForward Index 71C10CATTCTACCATTCTACGTAGAATG	Forward Index 58	J8	GAGTAACG	GAGTAACG	CGTTACTC
Forward Index 61M8GGATATCGGGATATCGCGATATCCForward Index 62N8GGTCAGATGGTCAGATATCTGACCForward Index 63O8GTAAGAAGGTAAGAAGCTTCTTACForward Index 64P8GTAGAAGTGTAGAAGTACTTCTACForward Index 65A9GTATCTGAGTATCTGATCAGATACForward Index 66B9GTCATCTAGTCATCTATAGATGACForward Index 67C9GTCCGCAAGTCCGCAATTGCGGACForward Index 68D9GTTCAATAGTTCAATATATTGAACForward Index 69A10TAACGTCGTAACGTCGCGACGTTAForward Index 70B10TCGGAACGTCGGAACGCGTTCCGAForward Index 71C10CATTCTACCATTCTACGTAGAATG	Forward Index 59	K8	GCAGTCCA	GCAGTCCA	TGGACTGC
Forward Index 62N8GGTCAGATGGTCAGATATCTGACCForward Index 63O8GTAAGAAGGTAAGAAGCTTCTTACForward Index 64P8GTAGAAGTGTAGAAGTACTTCTACForward Index 65A9GTATCTGAGTATCTGATCAGATACForward Index 66B9GTCATCTAGTCATCTATAGATGACForward Index 67C9GTCCGCAAGTCCGCAATTGCGGACForward Index 68D9GTTCAATAGTTCAATATATTGAACForward Index 69A10TAACGTCGTAACGTCGCGACGTTAForward Index 70B10TCGGAACGTCGGAACGCGTTCCGAForward Index 71C10CATTCTACCATTCTACGTAGAATG	Forward Index 60	L8	GCTCAAGG	GCTCAAGG	CCTTGAGC
Forward Index 63O8GTAAGAAGGTAAGAAGCTTCTTACForward Index 64P8GTAGAAGTGTAGAAGTACTTCTACForward Index 65A9GTATCTGAGTATCTGATCAGATACForward Index 66B9GTCATCTAGTCATCTATAGATGACForward Index 67C9GTCCGCAAGTCCGCAATTGCGGACForward Index 68D9GTTCAATAGTTCAATATATTGAACForward Index 69A10TAACGTCGTAACGTCGCGACGTTAForward Index 70B10TCGGAACGTCGGAACGCGTTCCGAForward Index 71C10CATTCTACCATTCTACGTAGAATG	Forward Index 61	M8	GGATATCG	GGATATCG	CGATATCC
Forward Index 64P8GTAGAAGTGTAGAAGTACTTCTACForward Index 65A9GTATCTGAGTATCTGATCAGATACForward Index 66B9GTCATCTAGTCATCTATAGATGACForward Index 67C9GTCCGCAAGTCCGCAATTGCGGACForward Index 68D9GTTCAATAGTTCAATATATTGAACForward Index 69A10TAACGTCGTAACGTCGCGACGTTAForward Index 70B10TCGGAACGTCGGAACGCGTTCCGAForward Index 71C10CATTCTACCATTCTACGTAGAATG	Forward Index 62	N8	GGTCAGAT	GGTCAGAT	ATCTGACC
Forward Index 65A9GTATCTGAGTATCTGATCAGATACForward Index 66B9GTCATCTAGTCATCTATAGATGACForward Index 67C9GTCCGCAAGTCCGCAATTGCGGACForward Index 68D9GTTCAATAGTTCAATATATTGAACForward Index 69A10TAACGTCGTAACGTCGCGACGTTAForward Index 70B10TCGGAACGTCGGAACGCGTTCCGAForward Index 71C10CATTCTACCATTCTACGTAGAATG	Forward Index 63	O8	GTAAGAAG	GTAAGAAG	CTTCTTAC
Forward Index 66B9GTCATCTAGTCATCTATAGATGACForward Index 67C9GTCCGCAAGTCCGCAATTGCGGACForward Index 68D9GTTCAATAGTTCAATATATTGAACForward Index 69A10TAACGTCGTAACGTCGCGACGTTAForward Index 70B10TCGGAACGTCGGAACGCGTTCCGAForward Index 71C10CATTCTACCATTCTACGTAGAATG	Forward Index 64	P8	GTAGAAGT	GTAGAAGT	ACTTCTAC
Forward Index 67C9GTCCGCAAGTCCGCAATTGCGGACForward Index 68D9GTTCAATAGTTCAATATATTGAACForward Index 69A10TAACGTCGTAACGTCGCGACGTTAForward Index 70B10TCGGAACGTCGGAACGCGTTCCGAForward Index 71C10CATTCTACCATTCTACGTAGAATG	Forward Index 65	A9	GTATCTGA	GTATCTGA	TCAGATAC
Forward Index 68D9GTTCAATAGTTCAATATATTGAACForward Index 69A10TAACGTCGTAACGTCGCGACGTTAForward Index 70B10TCGGAACGTCGGAACGCGTTCCGAForward Index 71C10CATTCTACCATTCTACGTAGAATG	Forward Index 66	B9	GTCATCTA	GTCATCTA	TAGATGAC
Forward Index 69A10TAACGTCGTAACGTCGCGACGTTAForward Index 70B10TCGGAACGTCGGAACGCGTTCCGAForward Index 71C10CATTCTACCATTCTACGTAGAATG	Forward Index 67	C9	GTCCGCAA	GTCCGCAA	TTGCGGAC
Forward Index 70B10TCGGAACGTCGGAACGCGTTCCGAForward Index 71C10CATTCTACCATTCTACGTAGAATG	Forward Index 68	D9	GTTCAATA	GTTCAATA	TATTGAAC
Forward Index 71 C10 CATTCTAC CATTCTAC GTAGAATG	Forward Index 69	A10	TAACGTCG	TAACGTCG	CGACGTTA
	Forward Index 70	B10	TCGGAACG	TCGGAACG	CGTTCCGA
Forward Index 72 D10 TTACTTCT TTACTTCT AGAAGTAA	Forward Index 71	C10	CATTCTAC	CATTCTAC	GTAGAATG
	Forward Index 72	D10	TTACTTCT	TTACTTCT	AGAAGTAA

Table 4. SMART-Seq ICELL8 Indexing Primer Set – A reverse indexing primers 1–72.

Reverse indexing (i7) primers			Reverse indexes (i7) on sample sheet
Index number	Well position	Index sequence in primers	MiSeq, MiniSeq, NextSeq, HiSeq 2000/2500, HiSeq 3000/4000, HiSeq X, NovaSeq
Reverse Index 1	A13	AACCGGTT	AACCGGTT
Reverse Index 2	B13	AACCTAGA	TCTAGGTT

Reverse Index 3	C13	AAGACCAG	CTGGTCTT
Reverse Index 4	D13	AGAACGAC	GTCGTTCT
Reverse Index 5	E13	AGAACTCT	AGAGTTCT
Reverse Index 6	F13	ATTCAGCT	AGCTGAAT
Reverse Index 7	G13	CATACGTC	GACGTATG
Reverse Index 8	H13	CCATTATG	CATAATGG
Reverse Index 9	113	CCTTGAAT	ATTCAAGG
Reverse Index 10	J13	CGGAATCA	TGATTCCG
Reverse Index 11	K13	CTAGTTGC	GCAACTAG
Reverse Index 12	L13	CTCGCGTA	TACGCGAG
Reverse Index 13	M13	CTTGAGTC	GACTCAAG
Reverse Index 14	N13	GAACGTAT	ATACGTTC
Reverse Index 15	O13	GACTGCGG	CCGCAGTC
Reverse Index 16	P13	GCGTACGG	CCGTACGC
Reverse Index 17	A14	GCTTCTCC	GGAGAAGC
Reverse Index 18	B14	GGAGGCTC	GAGCCTCC
Reverse Index 19	C14	GGAGTATG	CATACTCC
Reverse Index 20	D14	GTCGCTAG	CTAGCGAC
Reverse Index 21	E14	TCGTTCGA	TCGAACGA
Reverse Index 22	F14	TCTCTACC	GGTAGAGA
Reverse Index 23	G14	TGGCGACG	CGTCGCCA
Reverse Index 24	H14	TTCGATGA	TCATCGAA
Reverse Index 25	114	TTGATCCA	TGGATCAA
Reverse Index 26	J14	AACCTGCC	GGCAGGTT
Reverse Index 27	K14	AACGCATC	GATGCGTT
Reverse Index 28	L14	AACGCCAT	ATGGCGTT
Reverse Index 29	 M14	AACGCGCA	TGCGCGTT
Reverse Index 30	N14	AAGAATGG	CCATTCTT
Reverse Index 31	014	AAGACGCT	AGCGTCTT
Reverse Index 32	P14	ACCAACCG	CGGTTGGT
Reverse Index 33	A15	ACCGAATG	CATTCGGT
Reverse Index 34	B15	ACTCGCTA	TAGCGAGT
Reverse Index 35	C15	AGAAGAGC	GCTCTTCT
Reverse Index 36	D15	AGAATCTC	GAGATTCT
Reverse Index 37	E15	ATGCTTAG	CTAAGCAT
Reverse Index 38	F15	CAGACCTT	AAGGTCTG
Reverse Index 39	G15	CCGCTAGG	CCTAGCGG
Reverse Index 40	H15	CCGGTTAG	CTAACCGG
Reverse Index 41	115	CCTCGACG	CGTCGAGG
Reverse Index 42	J15	CGAAGCTG	CAGCTTCG
Reverse Index 43	K15	CGACCGCG	CGCGGTCG
Reverse Index 44	L15	CGTCATAA	TTATGACG
Reverse Index 45	M15	CTAGGAGA	TCTCCTAG
Reverse Index 46	N15	CTATTCAT	ATGAATAG
Reverse Index 47	015	CTCTACTT	AAGTAGAG
Reverse Index 48	P15	CTGATTGA	TCAATCAG
	113	OTORTOR	TOATIONO

Reverse Index 49	A16	CTTCGTTA	TAACGAAG
Reverse Index 50	B16	GAAGCAGC	GCTGCTTC
Reverse Index 51	C16	GAATAGGC	GCCTATTC
Reverse Index 52	D16	GCTCTGCT	AGCAGAGC
Reverse Index 53	E16	GGAGCGCA	TGCGCTCC
Reverse Index 54	F16	GGCGGTAT	ATACCGCC
Reverse Index 55	G16	GGTAACGC	GCGTTACC
Reverse Index 56	H16	GGTACGCC	GGCGTACC
Reverse Index 57	I16	GGTAGAAT	ATTCTACC
Reverse Index 58	J16	GGTTAGTC	GACTAACC
Reverse Index 59	K16	GTCTCGCG	CGCGAGAC
Reverse Index 60	L16	GTTCTACG	CGTAGAAC
Reverse Index 61	M16	TAGTATCT	AGATACTA
Reverse Index 62	N16	TAGTTAGG	CCTAACTA
Reverse Index 63	O16	TATTGCGC	GCGCAATA
Reverse Index 64	P16	TCAGTTAA	TTAACTGA
Reverse Index 65	A17	TCCGTATA	TATACGGA
Reverse Index 66	B17	TCCTGAGA	TCTCAGGA
Reverse Index 67	C17	TCGTCGCC	GGCGACGA
Reverse Index 68	D17	TGGCGTTA	TAACGCCA
Reverse Index 69	A18	TGGTATGA	TCATACCA
Reverse Index 70	B18	TTAAGCGT	ACGCTTAA
Reverse Index 71	C18	TTCGCGAC	GTCGCGAA
Reverse Index 72	D18	TTGCATAT	ATATGCAA

Table 5. SMART-Seq ICELL8 Indexing Primer Set – B reverse indexing primers 73–144.

Reverse in	dexing (i7) p	orimers	Reverse indexes (i7) on sample sheet
Index number	Well position	Index sequence in primers	MiSeq, MiniSeq, NextSeq, HiSeq 2000/2500, HiSeq 3000/4000, HiSeq X, NovaSeq
Reverse Index 73	A13	AACTCTCC	GGAGAGTT
Reverse Index 74	B13	AACTGATA	TATCAGTT
Reverse Index 75	C13	AAGAGAAT	ATTCTCTT
Reverse Index 76	D13	AAGTTGGA	TCCAACTT
Reverse Index 77	E13	ACGAACTT	AAGTTCGT
Reverse Index 78	F13	ACGCAACC	GGTTGCGT
Reverse Index 79	G13	ACGGAGGA	TCCTCCGT
Reverse Index 80	H13	ACTTACGT	ACGTAAGT
Reverse Index 81	I13	ACTTCTAA	TTAGAAGT
Reverse Index 82	J13	AGACGGAA	TTCCGTCT
Reverse Index 83	K13	AGAGGTCC	GGACCTCT
Reverse Index 84	L13	AGATGCGA	TCGCATCT
Reverse Index 85	M13	AGCAAGGC	GCCTTGCT
Reverse Index 86	N13	AGGCCTTG	CAAGGCCT
Reverse Index 87	O13	AGGTTATG	CATAACCT

Reverse Index 88	P13	AGTATAGT	ACTATACT
Reverse Index 89	A14	ATGGTACT	AGTACCAT
Reverse Index 90	B14	ATTACGAA	TTCGTAAT
Reverse Index 91	C14	CATAACGT	ACGTTATG
Reverse Index 92	D14	CATTAGAA	TTCTAATG
Reverse Index 93	E14	CCAGGCAT	ATGCCTGG
Reverse Index 94	F14	CCGTACTA	TAGTACGG
Reverse Index 95	G14	CGCGCTCA	TGAGCGCG
Reverse Index 96	H14	CGCGGTTG	CAACCGCG
Reverse Index 97	114	CGCTCTGG	CCAGAGCG
Reverse Index 98	J14	CGGCTAAC	GTTAGCCG
Reverse Index 99	K14	CGTCCTCC	GGAGGACG
Reverse Index 100	L14	CGTTGCGG	CCGCAACG
Reverse Index 101	M14	CTACGTCC	GGACGTAG
Reverse Index 102	N14	CTATCAAG	CTTGATAG
Reverse Index 103	O14	CTCGAGGT	ACCTCGAG
Reverse Index 104	P14	CTCGTCCA	TGGACGAG
Reverse Index 105	A15	CTCTGGCC	GGCCAGAG
Reverse Index 106	B15	CTGCAATG	CATTGCAG
Reverse Index 107	C15	CTGCCTCG	CGAGGCAG
Reverse Index 108	D15	CTTCATGG	CCATGAAG
Reverse Index 109	E15	GAAGTCGT	ACGACTTC
Reverse Index 110	F15	GAATCATG	CATGATTC
Reverse Index 111	G15	GACGGATT	AATCCGTC
Reverse Index 112	H15	GACGTACG	CGTACGTC
Reverse Index 113	115	GAGGCCAA	TTGGCCTC
Reverse Index 114	J15	GATATATT	AATATATC
Reverse Index 115	K15	GCATTGGT	ACCAATGC
Reverse Index 116	L15	GCGAAGCA	TGCTTCGC
Reverse Index 117	M15	GCGCCTTC	GAAGGCGC
Reverse Index 118	N15	GCGCTCTT	AAGAGCGC
Reverse Index 119	015	GCTAAGAC	GTCTTAGC
Reverse Index 120	P15	GGAATTGG	CCAATTCC
Reverse Index 121	A16	GGCAGGAC	GTCCTGCC
Reverse Index 122	B16	GGTACCAA	TTGGTACC
Reverse Index 123	C16	GGTCCTAG	CTAGGACC
Reverse Index 124	D16	GTAATCCG	CGGATTAC
Reverse Index 125	E16	GTCCTAAC	GTTAGGAC
Reverse Index 126	F16	GTTCAGGC	GCCTGAAC
Reverse Index 127	G16	TAATACGT	ACGTATTA
Reverse Index 128	H16	TACGAGTT	AACTCGTA
Reverse Index 129	116	TACGGTAC	GTACCGTA
Reverse Index 130 Reverse Index 131	J16 K16	TATATGCC	GGCATATA
Reverse Index 131	L16	TATATTGA TCAGGCGA	
Reverse Index 132	M16	TCAGGCGA	TCGCCTGA CTTCATGA
NEVELSE INVEX 133	IVITO	ICAIGAAG	CITCATGA

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Reverse Index 134	N16	TCCGACCT	AGGTCGGA	
Reverse Index 135	O16	TCGAATAA	TTATTCGA	
Reverse Index 136	P16	TCGGTCAT	ATGACCGA	
Reverse Index 137	A17	TCTAGAGG	CCTCTAGA	
Reverse Index 138	B17	TCTCCGTC	GACGGAGA	
Reverse Index 139	C17	TGCGGACT	AGTCCGCA	
Reverse Index 140	D17	TTAACCAA	TTGGTTAA	
Reverse Index 141	A18	TTACCATT	AATGGTAA	
Reverse Index 142	B18	TTATCGTC	GACGATAA	
Reverse Index 143	C18	TTCATACG	CGTATGAA	
Reverse Index 144	D18	TTCCGGTC	GACCGGAA	

Appendix B: RT-PCR Cycling Guidelines

While 14 cycles of PCR during the RT-PCR step will work to provide good transcript identification and sensitivity for many cell types, different cell types may have different optimal cycle numbers during RT-PCR. Cells with high mRNA content (like cell lines such as K-562) may require as few as 8 cycles of PCR to provide high sensitivity, with higher cycle numbers decreasing sensitivity. In contrast, cells with low mRNA content (like primary cells such as PBMCs) may show their highest sensitivity at 14 cycles of PCR, with lower sensitivity at fewer cycles.

After bioinformatic analysis, if a cell type exhibits high percentages of exonic reads ($\geq 80\%$), the cycle number may be reduced. If instead a cell type exhibits low percentages of exonic reads (50–80%), then the cycle number may be increased.

Appendix C: Guidelines for Library Sequencing

A. Final Sequencing Library Structure

Libraries generated using the SMART-Seq full-length protocol on the ICELL8 cx system have standard Illumina adapters and indexes. The unique combinations of indexes (i5 and i7) are required to discriminate between cells from different wells. Therefore, dual indexes (2 x 8 nt) must be sequenced. This unique combination of i5 and i7 indexes per well in the nanochip is generated using 72 i5 (forward) and 72 i7 (reverse) indexes. The structure of a final sequencing library is shown in Figure 22, below. For the complete list of sequences of forward and reverse indexing primers, please refer to Appendix A.

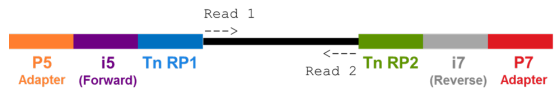


Figure 22. Structure of a final sequencing library. The final library is composed of Tn RP1 and Tn RP2 (Illumina Nextera read primer 1 and 2 sequences added by the TDE1 enzyme), i5 and i7 indexes, and P5 and P7 adapters.

B. Compatible Illumina Platforms

SMART-Seq full-length libraries generated with the ICELL8 cx system can be run on the following Illumina platforms with either single-end or paired-end sequencing (Takara Bio validated libraries with MiSeq, MiniSeq, NextSeq and NovaSeq):

- MiSeq
- MiniSeq
- NextSeq series

- HiSeq 2000/2500, 3000/4000
- HiSeq X series (Rapid Run and High Output)
- NovaSeq 6000

C. Recommendations for Library Sequencing

Samples should be pooled to a final concentration of 4 nM. Refer to Illumina documentation for instructions on denaturing and diluting libraries.

1. Loading Concentration and PhiX Recommendations

For libraries <850 base pairs as determined by the Bioanalyzer (Section V.M, Step 4), use the Qubit concentration in combination with the average size of the library to calculate the molar concentration of the library. For libraries >850 base pairs as determined by the Bioanalyzer, use the qPCR concentration (Section V.M, Step 4).

For NextSeq instruments, we recommend a loading concentration of 1.6 pM as a good starting point. Refer to information about individual Illumina instruments on Illumina's website for loading guidelines.

Libraries generated with this protocol do not require the inclusion of PhiX. However, particular sample types sometimes do not display the well-balanced nucleotide diversity required for base calling. If you are not sure about the performance of your sample, then please include PhiX. Follow Illumina guidelines on how to denature, dilute, and combine a PhiX control library with your own pool of libraries. Make sure to use a fresh and reliable stock of the PhiX control library.

2. Sequencing Depth Recommendations

The sequencing depth is dependent on the purpose of the study. Identification of cell heterogeneity can be achieved with shallow sequencing of about 50,000–100,000 reads/cell. However, identification of rare genes or alternative splicing transcript isoforms may need higher sequencing depth. We usually recommend more than 300,000 reads per cell. The following table provides some guidelines, for an example in which 1,000 single cells are selected in the experiment. Please refer to the "Illumina sequencing platforms" page for more details: <u>https://www.illumina.com/systems/sequencing-platforms.html</u>.

Platform	Maximum reads per run (millions)	Flow cell type	# of flow cells per run	# of lanes per flow cell	Maximum reads per lane (millions)	Reads per cell*; 1,000 cells (thousands)**	Platform recommendation***
MiSeq	25	_	1	1	-	25	No
MiniSeq	25	_	1	1	-	25	No
NextSeq Series (High output)	400	_	1	4	-	400	Yes
HiSeq 4000	5,000	_	1 or 2	8	312.5	312.5	Yes
HiSeq X Series	6,000	-	1 or 2	8	375	375	Yes
NovaSeq 6000	3,200	S1	1 or 2	2	800	800	Yes
NovaSeq 6000	8,200	S2	1 or 2	2	2,050	2,050	Yes
NovaSeq 6000	20,000	S4	1 or 2	4	2,500	2,500	Yes

Table 6. Sequencing depth recommendations.

*We recommend >300,000 reads per cell. However, this number depends on the goal of the experiment.

**Formula used: (# of reads per cell) = (maximum reads per run or lane) / (# of cells).

***The platform recommendation is based on our recommendation of 300,000 reads per cell. Therefore, it varies based on the number of single cells sequenced as well as on the purpose of the study.

3. Sequencing Run Parameters

SMART-Seq libraries generated with the ICELL8 cx system use standard Illumina sequencing primers and do not need custom primers. Dual indexes $(2 \times 8 \text{ cycles})$ *must* be sequenced and we recommend the number of cycles in Table 7, below. However, cycles of Read 1 and Read 2 can be flexible (e.g., 2×100 , 2×150 , and 2×250) depending on your experiments. Single-end sequencing can also be performed with SMART-Seq full-length libraries (e.g., 1×100 , 1×150 , and 1×250).

Table 7. Sequencing run parameters.

Sequencing read	Recommended # of cycles
Read 1	75
i7 Index	8
i5 Index	8
Read 2	75

Appendix D: Demultiplexing and Data Analysis

There are two options to demultiplex your sequencing data and generate the files.

1. Use CogentTM NGS Analysis Pipeline, Takara Bio's single-cell RNA-seq analysis program

Cogent NGS Analysis Pipeline (CogentAP) is an end-to-end analysis software that takes sequencing data as input and can perform de-multiplexing, alignment, counting, and report generation as an HTML-format report with the most commonly used statistics and plots in single-cell analysis. However, if desired, it can be used for demultiplexing alone. This choice can be made while launching the pipeline.

Significant advantages of using CogentAP:

- Perform end-to-end analysis or just de-multiplexing with ease
- Provide the well list generated during the experiment (using CellSelect cx software) as the input without the need to manually handle barcodes and sample information
- CogentAP automatically performs a reverse-complement check and correction of barcodes
- There is no limit on the number of barcodes one can process
- Output files generated by CogentAP can be loaded into Cogent NGS Discovery Software (CogentDS), our software for extended analysis, to create Uniform Manifold Approximation and Projection (UMAP) and t-Distributed Stochastic Neighbor Embedding (t-SNE) visualization popular in scRNA-seq.

Please make sure to keep your CogentAP and CogentDS versions up to date to ensure full compatibility with CELLSTUDIO software. Installation and upgrade information can be found in the user guides for both CogentAP and CogentDS, available through the ICELL8 software portal at <u>takarabio.com/icell8-software</u>.

2. Use Illumina's bcl2fastq Conversion Software

After installing bcl2fastq Conversion Software for demultiplexing, a custom sample sheet needs to be generated in order to use the software. Sample sheets can be created by Illumina Experiment Manager https://support.illumina.com/sequencing/sequencing_software/experiment_manager.html.

SMART-Seq ICELL8 indexes are unique sequences that are different from Illumina indexes. Before demultiplexing, the sample sheets created in the Illumina Experiment Manager (above) have to be customized with SMART-Seq ICELL8 forward and reverse indexes (see Appendix A).

Please note that forward indexes (i5) in the sample sheet are dependent on Illumina sequencers, and on some sequencers, the indexes must be entered in the sample sheet in the reverse-complement. The SMART-Seq indexes of the wells selected during an experiment can be obtained from the well-list file generated by the ICELL8 cx CellSelect Software on the ICELL8 cx instrument. Additionally, some versions of bcl2fastq software handle only ~400 barcodes at a time, requiring it to be run multiple times with multiple sample sheets, when more than 400 wells are selected. Please see Illumina's User Guide for more details

(https://support.illumina.com/sequencing/sequencing_software/bcl2fastq-conversion-software.html).

The resulting demultiplexed FASTQ files can then be analyzed by other compatible custom pipelines, such as the mappa software.

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This document has been reviewed and approved by the Quality Department.