CG000507 Rev A



USER GUIDE

Chromium Next GEM Automated Single Cell 5' Reagent Kits v2

with Feature Barcode technology for Cell Surface Protein & Immune Receptor Mapping

FOR USE WITH

Chromium Next GEM Automated Single Cell 5' Kit v2, 24 rxns PN-1000290 Chromium Next GEM Automated Single Cell 5' Kit v2, 4 rxns PN-1000298 Chromium Next GEM Automated Single Cell 5' Feature Barcode Library Construction Kit, 24 rxns PN-1000455 Chromium Automated Single Cell Human TCR Amplification & Library Construction Kit, 24 rxns PN-1000300 Chromium Automated Single Cell Mouse TCR Amplification & Library Construction Kit, 24 rxns PN-1000310 Chromium Automated Single Cell Human BCR Amplification & Library Construction Kit, 24 rxns PN-1000305 Chromium Automated Single Cell Human BCR Amplification & Library Construction Kit, 24 rxns PN-1000305 Chromium Automated Single Cell Mouse BCR Amplification & Library Construction Kit, 24 rxns PN-1000311 Chromium Next GEM Chip K Automated Single Cell Kit, 48 rxns PN-1000289 Chromium Next GEM Chip K Automated Single Cell Kit, 16 rxns PN-1000297 Dual Index Kit TT Set A, 96 rxns PN-1000215 Dual Index Kit TN Set A, 96 rxns PN-1000250



Notices

Document Number

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ТОС

Introduction

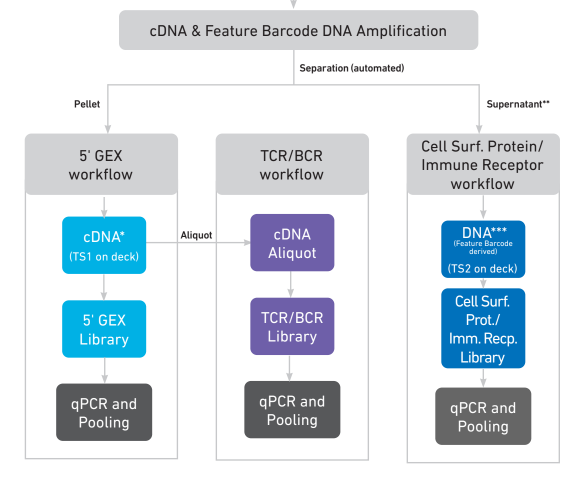
Chromium Next GEM Automated Single Cell 5' with Feature Barcode technology Workflow Additional Kits, Reagents & Equipment Recommended Thermal Cyclers Recommended Real Time qPCR System Protocol Steps & Timing Stepwise Objectives Chromium Next GEM Automated Single Cell 5' Reagent Kits v2

Chromium Automated Single Cell 5' Workflows

Single Cell Suspension

Cells labeled with:

- Feature Barcode oligonucleotide conjugated to antibodies for detecting cell surface protein expression or
- Feature Barcode oligonucleotide conjugated to multimeric MHC peptide complexes, such as a dCODE Dextramer[®] along with the Feature Barcode oligonucleotide conjugated antibody for mapping immune receptors



*For Automated Gene Expression flexible workflow with cDNA storage option, refer to Chromium Next GEM Single Cell 5' cDNA Kit v2 User Guide Supplement (CG000473) and Automated Library Construction User Guide (CG000474). If 5' Gene Expression libraries are not desired, stop the automated protocol after cDNA amplification and proceed directly to V(D)J amplification.

**Remove the supernatant from the instrument deck. It can be stored at 4°C or at -20°C for up to 5 weeks before proceeding to the next automated step (Feature Barcode DNA Supernatant – SPRIselect Cleanup).



***After the automated Feature Barcode DNA Supernatant – SPRIselect Cleanup step, the additional DNA volume that is not required for sample index PCR may be removed from the instrument based on the instrument touchscreen prompts. The remaining DNA volume can be stored at 4°C for up to 72 h or at –20°C for up to 4 weeks and may be used for additional Cell Surface Protein/Immune Receptor Mapping library construction using the manual workflow (cannot be used for automated library generation).

7

Additional Kits, Reagents & Equipment	The items in the table below have been validated by the Chromium Connect Automated Single Cell 5' pr listed materials.	
Supplier	Description	Part Number (US)
Plastics		
Hamilton	CO-RE Tips 50 μl Filtered Tips * CO-RE Tips 300 μl Filtered Tips* 60 ml Reagent Reservoir Self-Standing Hamilton PCR ComfortLid	235948 235903 194051 814300
	*CO-RE pipette tips will be phased out and replaced by new CO-RE II pip early 2022. CO-RE II tips include a new sealing surface to interface with with the current CO-RE stop disk is identical between the two tip designs	the CO-RE II stop disk. Geometry that interfaces
Eppendorf	96-well Full-Skirted Plate** 96-well Semi-Skirted Plate (Blue color listed; other colors are acceptable)	951020460 951020362
	**Alternatively, use Amplifyt 96-Well PCR Plates, Full-Skirted, Clear from Fisher Scientific (NC1959287 for 25/case, NC1959288 for 100/case)	n Thomas Scientific (485096/1149K05) or
Thermo Fisher Scientific	MicroAmp 8-Tube Strip, 0.2 ml MicroAmp 8-Cap Strip, clear	N8010580 4323032
Kits & Reagents		
Thermo Fisher Scientific	Nuclease-free Water	AM9937
Millipore Sigma	Ethanol, Pure (200 Proof, anhydrous)	E7023-500ML
Qiagen	Qiagen Buffer EB	19086
Equipment		
10x Genomics	10x Vortex Adapter Benchtop Vortex Benchtop Centrifuge Plate Centrifuge Benchtop Thermal Cycler	330002 standard lab equipment standard lab equipment standard lab equipment standard lab equipment
Additional materials ONLY for	r optional assays – qPCR and pooling	
Bio-Rad	10% Tween 20 96-well PCR Plates	1662404 HSP9665
Thermo Fisher Scientific	2 ml-Screw-cap Tubes, NonKnurl 0.5 ml-Screw-cap Tubes, NonKnurl	3488NK 3472NK
KAPA Biosystems	KAPA Library Quantification Kit for Illumina Platforms	KK4824
Qiagen	Qiagen Buffer EB	19086
Additional materials for Chro	mium Connect maintenance	
Thor Labs	Lens tissues	MC-5
VWR	Microcide SQ Broad Spectrum Disinfectant	25099
Contec	70% Isopropanol (alternative to VWR disinfectant)	SB167030IR

Use only indicated cleaning agents. DO NOT use bleach or organic oxidizers.

Additional Kits, Reagents & Equipment

Supplier	Description	Part Number (US)
Quantification & Quality Cont	trol	
Agilent	2100 Bioanalyzer Laptop Bundle (discontinued) (Replacement 2100 Bioanalyzer Instrument/ 2100 Expert Laptop Bundle	G2943CA G2939BA/G2953CA
Choose Bioanalyzer, TapeStation, LabChip, or Qubit based on availability & preference.	High Sensitivity DNA Kit 4200 TapeStation High Sensitivity D1000 ScreenTape/Reagents High Sensitivity D5000 ScreenTape/Reagents	5067-4626 G2991AA 5067-5592/ 5067-5593 5067-5584/ 5067-5585
Thermo Fisher Scientific	Qubit 4.0 Fluorometer Qubit dsDNA HS Assay Kit	Q33238 Q32854
PerkinElmer	LabChip GX Touch HT Nucleic Acid Analyzer DNA High Sensitivity Reagent Kit	CLS137031 CLS760672
Recommended	Thermal cyclers for off-deck use.	
Thermal Cyclers	Supplier Description Part Nu	ımber

Supplier	Description	Part Number
Bio-Rad	C1000 Touch Thermal Cycler with 96-Deep Well Reaction Module	1851197
Eppendorf	MasterCycler Pro (discontinued)	North America 950030010 International 6321 000.019
Thermo Fisher Scientific	Veriti 96-Well Thermal Cycler	4375786

Recommended Real Time qPCR System

Supplier	Description	Part Number
Bio-Rad	CFX96 Real-time System	1855096

The qPCR system should be compatible with Bio-Rad 96-well PCR Plates, P/N HSP9665 and with the KAPA Library Quantification Kit dye. Refer to manufacturer's recommendation.

Protocol Steps & Timing

		Steps	Timing
	MANUAL	Cell Preparation & Labeling (Dependent on cell type & labeling protocol) Gather & Load Reagents and Consumables	~1-2 h ~60 min
	AUTOMATED	 Master Mix Preparation Chromium Automated Controller Loading GEM Generation OPTIONAL Confirm GEM Generation (Manual, 5 min) ~45 min after starting Post GEM RT-Cleanup – Dynabead cDNA & Feature Barcode DNA Amplification cDNA Cleanup – SPRIselect 	~3.5 h Walk-away time
	MANUAL	Remove the amplified Feature Barcode DNA supernatant from the deck cDNA QC & Quantification	~60 min
	AUTOMATED	 5' Gene Expression Library Construction Fragmentation, End Repair & A-tailing Post Fragmentation, End Repair & A-tailing Double Sided Size Selection – SPRIselect Adaptor Ligation Post Ligation Cleanup – SPRIselect Sample Index PCR Post Sample Index PCR Double Sided Size Selection – SPRIselect 	~4.5 h Walk-away time
	MANUAL	Post Library Construction QC	~60 min
	MANUAL	V(D)J Amplification & Library Construction Gather & Load Reagents and Consumables	~45 min
12 h	AUTOMATED	 V(D)J Amplification 1 V(D)J Amplification 1 Double Sided Size Selection – SPRIselect V(D)J Amplification 2 V(D)J Amplification 2 Double Sided Size Selection – SPRIselect 	~3.5 h Walk-away time
plus	MANUAL	V(D)J Amplification QC & Quantification	~60 min
	AUTOMATED	 Fragmentation, End Repair & A-tailing Adaptor Ligation Post Ligation Cleanup – SPRIselect Sample Index PCR Post Sample Index PCR Cleanup – SPRIselect 	~4.5 h Walk-away time
	MANUAL	Post Library Construction QC	~60 min
	MANUAL	Cell Surface Protein/Immune Receptor Mapping Library Construction Gather & Load Reagents and Consumables	~45 min
	AUTOMATED	 Amplified Feature Barcode DNA – SPRIselect cleanup* Sample Index PCR Post Sample Index PCR Cleanup – SPRIselect 	~2.5 h Walk-away time
	MANUAL	Post Library Construction QC	~60 min
		OPTIONAL Library Quantification gPCR & Library Pooling	

OPTIONAL Library Quantification qPCR & Library Pooling *After the Amplified Feature Barcode DNA Supernatant – SPRIselect Cleanup step, the additional DNA volume that is not required for sample index PCR may be removed from the instrument based on the instrument touchscreen prompts . The remaining DNA volume can be stored at 4°C for up to 72 h or at –20°C for up to 4 weeks and may be used for additional Cell Surface Protein/Immune Receptor Mapping library construction using the manual workflow (cannot be used for automated library generation).

Stepwise Objectives

Chromium Connect automates the preparation of sequencing-ready, single cell libraries from input samples with walk-away convenience. Generation of Chromium Single Cell 5' Gene Expression, V(D)J, and Cell Surface Protein/Immune Receptor Mapping libraries on the Chromium Connect instrument includes automated Gel Beads-in-emulsion (GEM) generation, barcoding, and library preparation from single cell suspensions, along with additional functionalities for library quantification and pooling.

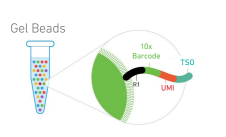
The Chromium Connect platform for 5' digital gene expression profiles 500-10,000 individual cells per sample. A pool of ~750,000 10x Barcodes is sampled separately to index each cell's transcriptome. It is done by partitioning thousands of cells into nanoliter-scale GEMs, where all generated cDNA share a common 10x Barcode. Libraries are generated and sequenced from the cDNA and 10x Barcodes are used to associate individual reads back to the individual partitions.

This document outlines the key automated protocol steps for generating the following libraries:

- Single Cell V(D)J libraries from V(D)J-amplified cDNA derived from poly-adenylated mRNA
- Single Cell 5' Gene Expression libraries from amplified cDNA derived from polyadenylated mRNA
- Single Cell 5' Cell Surface Protein libraries (include immune receptor mapping when cells are also labeled with multimeric MHC peptide complexes, such as Dextramer reagents) from amplified DNA derived from Feature Barcode

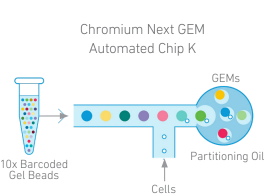
Single Cell 5' Gel Beads

The Single Cell VDJ 5' Gel Beads primer enables the production of barcoded, full-length cDNA from poly-adenylated mRNA, for generating Single Cell 5' Gene Expression and V(D)J, as well as Cell Surface Protein library DNA.



Automated GEM Generation & Barcoding

Automated GEM generation is done by combining barcoded Single Cell VDJ 5'Gel Beads, a Master Mix containing cells and enzymes, and Partitioning Oil onto Chromium Next GEM Automated Chip K. To achieve single cell resolution, cells are delivered at a limiting dilution, such that the majority (~90-99%) of generated GEMs contain no cell, while the remainder largely contain a single cell.



Enzyme

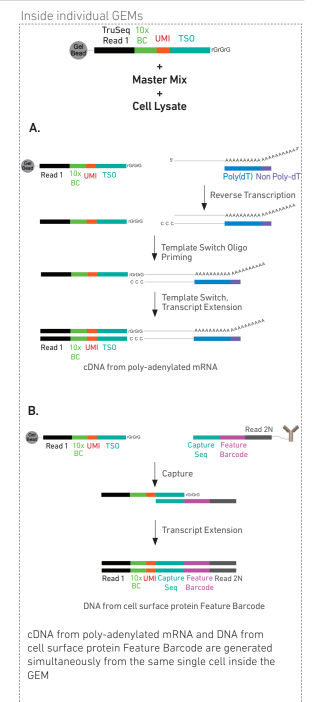
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Automated GEM Generation & Barcoding

Immediately following GEM generation, the Gel Bead is dissolved and any co-partitioned cell is lysed. Gel Bead primers containing (i) an Illumina TruSeq Read 1 sequence (read 1 sequencing primer), (ii) a 16 nt 10x Barcode, (iii) a 10 nt unique molecular identifier (UMI), and (iv) 13 nt template switch oligo (TSO) are released and mixed with the cell lysate and a Master Mix containing reverse transcription (RT) reagents and poly(dT) primers.

A. The cell lysate and the released Gel Bead primer incubated with the Master Mix containing RT reagents, produce 10x Barcoded, full-length cDNA from polyadenylated mRNA.

B. Simultaneously in the same partition, the Gel Bead primer captures the cell surface protein Feature Barcode conjugated to the antibody or to antibody and antigen containing (i) a Nextera Read 2 (Read 2N), (ii) a 15 nt Feature Barcode, and (iii) Capture Sequence. Incubation of the GEMs with the Master Mix containing RT reagents produces 10x Barcoded DNA from the cell surface protein Feature Barcode.



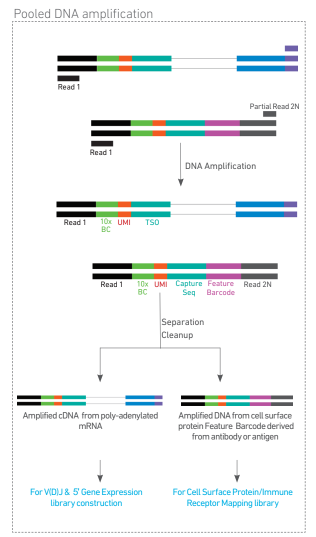
Automated Post GEM-RT Cleanup & DNA Amplification

GEMs are broken and pooled after GEM-RT reaction mixtures are recovered. Silane magnetic beads are used to purify the 10x Barcoded first-strand cDNA from polyadenylated mRNA and DNA from cell surface protein/antigen specificity Feature Barcode from the post GEM-RT reaction mixture, which includes leftover biochemical reagents and primers.

10x Barcoded, full-length cDNA is amplified via PCR with primers against common 5' and 3' ends added during GEM-RT. Amplification generates sufficient material to construct multiple libraries from the same cells, e.g. both T cell and/or B cell libraries and 5' Gene Expression libraries.

If 5' Gene Expression libraries are not desired, stop the automated protocol after cDNA amplification and proceed directly to V(D)J amplification. Unused 5' Gene Expression library construction reagents will be lost, if the Gene Expression Library Construction is not performed as the next step. A more suitable option in this case would be to use the Chromium Next GEM Automated Single Cell 5' cDNA Kit (PN-1000425) for cDNA generation.

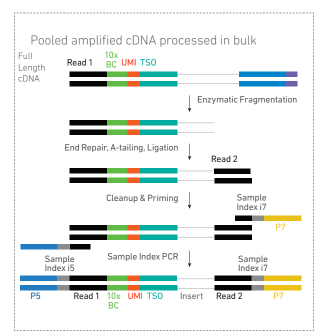
After separation and cleanup, supernatant, the containing the amplified DNA derived from the Feature Barcode associated with the cell surface protein, is removed from the instrument deck. It can be stored at 4°C or -20°C for up to 5 weeks before proceeding to automated Cell Surface Protein Library Construction.



Refer to Chromium Next GEM Automated Single Cell 5' cDNA Kit v2 User Guide Supplement (CG000473) and Automated Gene Expression Library Construction User Guide (CG000474) for more details.

Automated 5' Gene Expression Library Construction

Amplified full-length cDNA from poly-adenylated mRNA is used to generate 5' Gene Expression library. Enzymatic fragmentation and size selection are used to optimize the cDNA amplicon size prior to 5' gene expression library construction. P5, P7, i5 and i7 sample indexes, and Illumina R2 sequence (read 2 primer sequence) are added via End Repair, A-tailing, Adaptor Ligation, and Sample Index PCR. The final libraries contain the P5 and P7 priming sites used in Illumina sequencers.

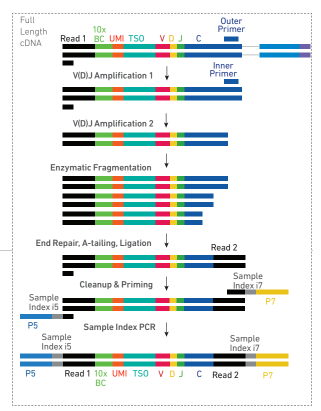


Automated V(D)J Amplification from cDNA

Amplified full-length cDNA from poly-adenylated mRNA is used to amplify full-length V(D)J segments (10x Barcoded) via PCR amplification with primers specific to either the TCR or BCR constant regions. If both T and B cells are expected to be present in the partitioned cell population, TCR and BCR transcripts can be amplified in separate reactions from the same amplified cDNA material.

Automated V(D)J Library Construction

Enzymatic fragmentation and size selection are used to generate variable length fragments that collectively span the V(D)J segments of the amplified TCR or BCR transcripts prior to library construction.

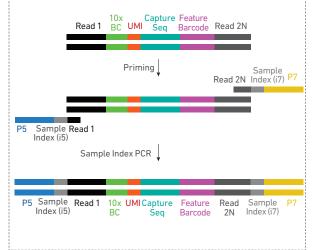


P5, P7, i5 and i7 sample indexes, and an Illumina R2 sequence (read 2 primer sequence) are added via End Repair, A-tailing, Adaptor Ligation, and Sample Index PCR. The final libraries contain the P5 and P7 priming sites used in Illumina sequencing.

Cell Surface Protein/ Immune Receptor Mapping Library Construction

Amplified DNA from the cell surface protein Feature Barcodes derived from the antibody or multimeric MHC peptide complexes, such as Dextramer reagents is used to construct the Cell Surface Protein library. A Cell Surface Protein library also detects antigen specificity if cells were labeled with both antibody and antigen.

P5, P7, i5 and i7 sample indexes, and Nextera Read 2 (Read 2N primer sequence) are added via Sample Index PCR.



Pooled amplified DNA processed in bulk

The final libraries contain the P5 and P7 priming sites used in Illumina sequencers.

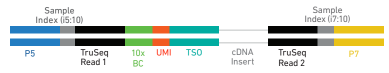
Sequencing

Illumina-ready dual index libraries can be sequenced at the recommended depth & run parameters. Illumina sequencer compatibility, sample indices, library loading and pooling for sequencing are summarized in the Sequencing chapter.

Chromium Single Cell V(D)J Dual Index Library



Chromium Single Cell 5' Gene Expression Dual Index Library



Chromium Single Cell 5' Cell Surface Protein Dual Index Library*



*Detects antigen specificity in cells labeled with antibodies and antigen

See Appendix for Oligonucleotide Sequences

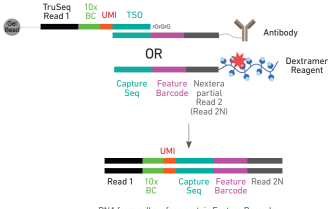
Cell Labeling Guidelines

Overview

Protein/s on the surface of a cell can be labeled with:

- A Feature Barcode oligonucleotide conjugated to a specific protein binding molecule, such as an antibody for detecting cell surface protein expression
- A Feature Barcode oligonucleotide conjugated to multimeric MHC peptide complexes, such as a dCODE Dextramer along with the Feature Barcode oligonucleotide conjugated antibody for mapping immune receptors and simultaneously detecting cell surface protein expression

The Feature Barcode conjugated molecule bound to the cell surface protein can be directly captured by the Gel Bead inside a GEM during GEM generation and amplified (see Stepwise Objectives for assay scheme specifics). The amplified DNA generated from the Feature Barcode can be used for Cell Surface Protein/Immune Receptor Mapping Library Construction.



DNA from cell surface protein Feature Barcode

Demonstrated Protocols for cell labeling

- Demonstrated Protocol "Cell Surface Protein Labeling for Single Cell RNA Sequencing Protocols with Feature Barcode technology (Document CG000149)".
- Demonstrated Protocol "Cell Labeling with dCODE Dextramer[®] Reagents for Single Cell RNA Sequencing Protocols with Feature Barcode technology (Document CG000203)".

Cell Surface Protein Library:

Amplified DNA from the cell surface protein Feature Barcode derived from the antibody or antibody and antigen is used to construct the Cell Surface Protein library. If cells were labeled with both antibody and antigen, the cell surface protein library will also map immune receptor.



Failure to label cell surface proteins with a Feature Barcode conjugated to a specific protein binding molecule prior to using the cells for GEM Generation & Barcoding will preclude generation of Cell Surface Protein library.

Chromium Next GEM Automated Single Cell 5' Reagent Kits v2

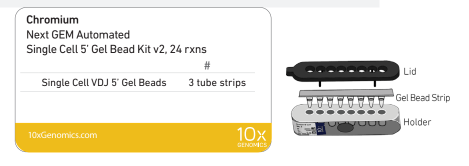
All reagent tube strips & tubes are for one time use only. DO NOT reuse.

Chromium Next GEM Automated Single Cell 5' Kit v2, 24 rxns PN-1000290

Reagent volumes and colors are different in each of the module types.

		\ \
Chromium Next GEM Automated Single Cell 5' v2, Module 1		
Module 1	# 24 tube strips	Module 1
Dynabeads™ MyOne™ SIL/	ANE 6 tubes (PN-2000048)	
10xGenomics.com		
omium Next GEM Auto Module 2, 24 rxns PN-	-	
	1000273 (Store at -	20 C)
Chromium Next GEM Automated Single Cell 5' v2, Module 2	24 5055	
Module 2	# 24 tube strips	Module 2
Module 2	#	Module 2
Module 2	#	Module 2
	#	Module 2
	# 24 tube strips	5' Kit v2,
10xGenomics.com omium Next GEM Auto Module 3, 24 rxns PN-7 Chromium	# 24 tube strips	5' Kit v2,
10xGenomics.com omium Next GEM Auto Module 3, 24 rxns PN-7	# 24 tube strips Look emated Single Cell 1000294 (store at -	5' Kit v2,
10xGenomics.com omium Next GEM Auto Module 3, 24 rxns PN-7 Chromium Next GEM Automated Single Cell 5' v2, Module 3,	# 24 tube strips 1000294 (store at - , 24 rxns #	5' Kit v2,
10xGenomics.com omium Next GEM Auto Module 3, 24 rxns PN-7 Chromium Next GEM Automated	# 24 tube strips Look emated Single Cell 1000294 (store at -	5' Kit v2, 20°C)

Chromium Next GEM Automated Single Cell 5' Gel Bead Kit v2, 24 rxns PN-1000291 (store at -80°C)



Chromium Next GEM Automated Single Cell 5' Kit v2, 4 rxns PN-1000298

Reagent volumes and colors are different in each of the module types.

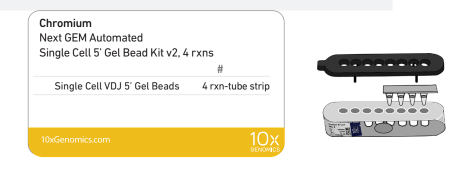
Chromium Next GEM Automated Single Cell 5' Kit v2, Module 1, 4 rxns PN-1000295 (store at 4°C)

Chromium Next GEM Automated Single Cell 5' v2, Module 1, 4 r>	kns	
	#	Module 1
O Module 1	4 tube strips	
O Dynabeads [™] MyOne [™] SILANE	2 tubes (PN-2000048)	
10xGenomics.com		

Chromium Next GEM Automated Single Cell 5' Kit v2, Modules 2 & 3, 4 rxns PN-1000296 (store at -20°C)

Chromium Next GEM Automated Single Cell 5' v2, Modules	2 & 3, 4 rxns	Module 2
	#	
O Module 2	4 tube strips	
O Module 3	4 tube strips	Module 3
Poly-dT RT Primer	2 tubes (PN-2000007)	
10xGenomics.com	10× Genomics	

Chromium Next GEM Automated Single Cell 5' Gel Bead Kit v2, 4 rxns PN-1000299 (store at -80°C)



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Chromium Automated Single Cell 5' Feature Barcode Library Construction Kit, 24 rxns PN-1000455

Reagent volumes and colors are different in each of the module types.

Automated Feature Bard Library Construction Kit Module 1, 24 rxns	#	Feature Barcode Library Constructio Module 1
Module 1	24 tube strips	
10xGenomics.com	10× genomics	
and the stars Design		· 1/1
	ode Library Construct -1000453 (store at -20	
Automated Feature Bard Library Construction Kit Module 2, 24 rxns		Feature Barcode Library Constructi Module 2
Module 2, 24 rxns	# 24 tube strips	
Module 2	24 tube strips	
10xGenomics.com	10×	
	GENOMICS	
	ture Barcode Primer I 00454 (store at -20°C)	-
	arcode Primer Kit,	
Automated 5' Feature Ba 24 rxns	#	
	# 4 6 tubes (PN-2000277)	

Chromium Automated Single Cell Human TCR Amplification & Library Construction Kit, 24 rxns PN-1000300

Reagent volumes and colors are different in each of the module types.

Chromium Automated TCR Amplification & Li V(D)J Module 1, 24 rxns PN	brary Constructio	n,
Chromium Automated Single Cell Hun Amplification & Library Co V(D)J Module 1, 24 rxns	nan TCR	
V(D)J Module 1	24 tube strips	V(D)J Module 1
10xGenomics.com		

Chromium Automated Single Cell Human TCR Amplification & Library Construction, V(D)J Module 2, 24 rxns PN-1000302 (store at -20°C)

Chromium Automated Single Cell Human TC Amplification & Library Construct V(D)J Module 2, 24 rxns		
V(D)J Module 2	24 tube strips	
Human T Cell Primer Mix 1 v2	6 tubes (PN-2000242)	
Human T Cell Primer Mix 2 v2	6 tubes (PN-2000246)	V(D)J Module 2
10xGenomics.com	10x	

Chromium Automated Single Cell Mouse TCR Amplification & Library Construction Kit, 24 rxns PN-1000310

Chromium Automated Single Cell Mouse TCR Amplification & Library Construction, V(D)J Module 1, 24 rxns PN-1000303 (store at 4°C)

Chromium Automated Single Cell Mouse Amplification & Library Cons V(D)J Module 1, 24 rxns	
V(D)J Module 1	24 tube strips
10xGenomics.com	10 genom

Chromium Automated Single Cell Mouse TCR Amplification & Library Construction, V(D)J Module 2, 24 rxns PN-1000304 (store at -20°C)

Chromium Automated Single Cell Mouse TC Amplification & Library Construc V(D)J Module 2, 24 rxns		
V(D)J Module 2	24 tube strips	
Mouse T Cell Primer Mix 1 v2	6 tubes (PN-2000256)	
Mouse T Cell Primer Mix 2 v2	6 tubes (PN-2000257)	V(D)J Module 2
	10x	

Chromium Automated Single Cell Human BCR Amplification & Library Construction Kit, 24 rxns PN-1000305

Chromium Automated Single Cell Human BCR Amplification & Library Construction, V(D)J Module 1, 24 rxns PN-1000306 (store at 4°C)

Chromium Automated Single Cell Huma Amplification & Library Cons V(D)J Module 1, 24 rxns	
V(D)J Module 1	24 tube strips
10xGenomics.com	1C geno

Chromium Automated Single Cell Human BCR Amplification & Library Construction, V(D)J Module 2, 24 rxns PN-1000307 (store at -20°C)

Chromium Automated Single Cell Human B Amplification & Library Constru- V(D)J Module 2, 24 rxns		
V(D)J Module 2	24 tube strips	
Human B Cell Primer Mix 1 v2	6 tubes (PN-2000254)	
Human B Cell Primer Mix 2 v2	6 tubes (PN-2000255)	V(D)J Module 2
	10	

Chromium Automated Single Cell Mouse BCR Amplification & Library Construction Kit, 24 rxns PN-1000311

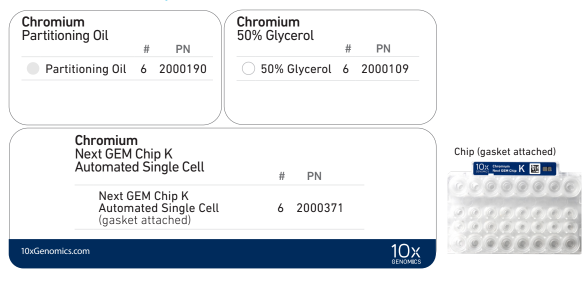
Chromium Automated Single Cell Mouse BCR Amplification & Library Construction Kit, V(D)J Module 1, 24 rxns PN-1000308 (store at 4°C)

Chromium Automated Single Cell Mous Amplification & Library Con V(D)J Module 1, 24 rxns	
V(D)J Module 1	24 tube strips
10xGenomics.com	10x genomics

Chromium Automated Single Cell Mouse BCR Amplification & Library Construction Kit, V(D)J Module 2, 24 rxns PN-1000309 (store at -20°C)

Chromium Automated Single Cell Mouse BC Amplification & Library Construct V(D)J Module 2, 24 rxns		
V(D)J Module 2	24 tube strips	
Mouse B Cell Primer Mix 1 v2	6 tubes (PN-2000258)	
Mouse B Cell Primer Mix 2 v2	6 tubes (PN-2000259)	V(D)J Module 2
10xGenomics.com	10×	

Chromium Next GEM Chip K Automated Single Cell Kit, 48 rxns PN-1000289 (store at ambient temperature)



Chromium Next GEM Chip K Automated Single Cell Kit, 16 rxns PN-1000297 (store at ambient temperature)

Chromium Partitioning Oil # PN	Chromium 50% Glycerol # PN	
Partitioning Oil 2 2000190	50% Glycerol 2 2000109	
Chromium Next GEM Chip K Automated Single Cell	# PN	Chip (gasket attached)
Next GEM Chip K Automated Single Cell (gasket attached)	2 2000371	
10xGenomics.com	10x genomics	

Dual Index Kit TT Set A, 96 rxns PN-1000215 (store at -20°C)

	# PN
Dual Index Plate TT Set A	1 3000431

Dual Index Kit TN Set A, 96 rxns PN-1000250 (store at -20°C)

	# PN
Dual Index Plate TN Set A	1 3000510

Quick Planner Card

Gather the listed items & reagents before running the assay. Follow the touchscreen prompts for detailed information.

G	ather indicated items prior to running the a	issay			
	Set thermal cycler to 37°C and lid to 50°C				Semi skirted plate, 96 well – 1 per run
	Nuclease free water – 10 ml				Full skirted plate, 96 well – 1 per run
	Ethanol, Pure (200 Proof, anhydrous) – 40 ml Combine 40 ml EtOH and 10 ml nuclease free v 80% EtOH	EtOH and 10 ml nuclease free water to prepare			 50 μl Black CO-RE/CO-RE II Pipette Tips, with filter 7-8 samples: 2 racks 4-6 samples: 2 racks
	Comfort lids – 6 per run				1-3 samples: 1 rack
	MicroAmp 8-tube strips, 0.2 ml – 3 per run				 300 µl Black CO-RE/ CO-RE II Pipette Tips, with filter 7-8 samples: 4 racks
	Reagent reservoirs, 60 ml – 3 per run				 4-6 samples: 3 racks 1-3 samples: 2 rack
1	0x Reagents	Storage	Pre	parat	on & Handling
	Next GEM Chip K Automated 1 per run	Room temp.	Set a deck		keep sealed. Follow the touchscreen prompts to load on
	Partitioning oil, 50% Glycerol 1 tube each per run	Room temp. (Chip box)	Keep capped. Follow the touchscreen prompts to remove the cap after cells are loaded on the deck.		
	Library Module 1 (black tube strip) 1 tube strip per sample	4°C	Use a thermal cycler (lid temp 50°C) to thaw for 30 min at 37°C . Vortex at 15 min and again at 30 min , centrifuge at 300 rcf for 1 min .		
	Dynabeads MyOne Silane - 1 tube per run DO NOT save excess 2 tubes/4rxn kit; 6 tubes/24rxn kit	4°C (Module 1 Box)	Equilibrate to room temperature. Vortex thoroughly (≥30 sec) immediately before use. Aspirate the full liquid volum with a pipette tip to verify that the beads have not settled in the bottom of the tube. If clumps are present, pipette mix to resuspend completely. DO NOT centrifuge before use. DO NOT remove cap until prompted on touchscreen.		
	Library Module 2 (grey tube strip) 1 tube strip per sample	-20°C	Thaw at room temperature for 30 min . Vortex, centrifuge at 300 rcf for 1 min .		
	Library Module 3 (white strip tube) 1 tube strip per sample	-20°C	Thav load	v at 4 ing, in	C or on ice. Maintain on ice until ready to load. Before vert mix (DO NOT vortex), centrifuge at 300 rcf for 1 min .
	Poly-dT RT Primer - 1 tube per run DO NOT save excess 2 tubes/4rxn kit; 6 tubes/24rxn kit	-20°C (Module 3 box)	Thav	w at ro	om temperature for 30 min . Vortex, centrifuge briefly.
	Dual Index Plate (SI Plate) 1 plate per run	-20°C	Thaw at room temperature for 30 min . Vortex, centrifuge briefly.		om temperature for 30 min . Vortex, centrifuge briefly.
	Gel Beads Strip(s)	-80°C	Thav	w at ro	om temperature ≥ 30 min . Vortex 30 sec , centrifuge 5 sec .
_	Feature cDNA Primers (as applicable) - 1 tube per run	-20°C	Tha	watro	om temperature for 30 min . Vortex, centrifuge briefly.

Tips & Best Practices

Consult the Chromium Connect User Guide (CG000180) and follow the Chromium Connect Touchscreen prompts for specifics of assay execution.

Consumables

• Use validated and recommended emulsion-safe plastic consumables as some plastics can destabilize GEMs.

Cell Concentration

- Resuspend samples in PBS+ 0.04% BSA (or alternative buffers specified in the relevant 10x Genomics Demonstrated Protocols - CG000053/ CG000149/CG000203). Total volume loaded onto the sample plate is 10 μl.
- Based on cell stock concentration, do sequential stock dilutions, if needed.
- Use 3 independent cell counts to determine cell concentration.
- The presence of dead cells in the suspension may also reduce the recovery rate. Consult the 10x Genomics Single Cell Protocols Cell Preparation Guide and the Guidelines for Optimal Sample Preparation flowchart (Documents CG000053 and CG000126 respectively) for more information on preparing cells.
- Refer to the 10x Genomics Support website for more information regarding cell type specific sample preparation.

Multiplet Rate (%)	# of Cells Loaded	# of Cells Recovered
~0.4%	~870	~500
~0.8%	~1,700	~1,000
~1.6%	~3,500	~2,000
~2.3%	~5,300	~3,000
~3.1%	~7,000	~4,000
~3.9%	~8,700	~5,000
~4.6%	~10,500	~6,000
~5.4%	~12,200	~7,000
~6.1%	~14,000	~8,000
~6.9%	~15,700	~9,000
~7.6%	~17,400	~10,000

Cell Preparation

- Ensure cell counts are accurate.
- Based on cell stock concentration, do sequential stock dilutions, if needed. Refer to the Cell Suspension Volume Calculator Table for optimal pipetting volumes and concentrations.
- Load cell samples when prompted on the touchscreen.
- The cDNA amplification cycle number will be based on the targeted cell recovery. The cycle number chosen for one sample will apply to all the samples in a run.

Reagent Handling	 Fully thaw and thoroughly mix reagents before use.
	 Resuspend Dynabeads and Feature cDNA Primer 4 at the end of loading.
	 Ensure there are no air bubbles at the bottoms of reagent tubes.
	 Follow the prompts on the touchscreen for handling Library Modules during setup and use.
	 Follow the prompts on the touchscreen for handling V(D)J Modules 1 and 2.
	 Ensure correct reagent tube barcode orientation (on tubes and racks) as prompted by the touchscreen.
	 Prepare and dispense 80% ethanol off-deck to avoid spilling on consumables.
	 When indicated, promptly move reagents back to the recommended storage.
Chromium	 The automated chip includes a pre-installed gasket.
Automated Chip Handling	 Minimize exposure of reagents and chips to sources of particles and fibers, laboratory wipes, frequently opened flip-cap tubes, clothing that sheds fibers, and dusty surfaces.
	 Keep chip and gasket in sealed package until

Barcode Orientation



- led
- to y wipes, that
- Keep chip and gasket in sealed package until prompted to load.
- · After removing the chip from the sealed bag, use in ≤ **24 h**.
- Avoid contacting the bottom surface of the chip with gloved hands and other surfaces.
- DO NOT use chips or gaskets specific to other 10x Genomics protocols.

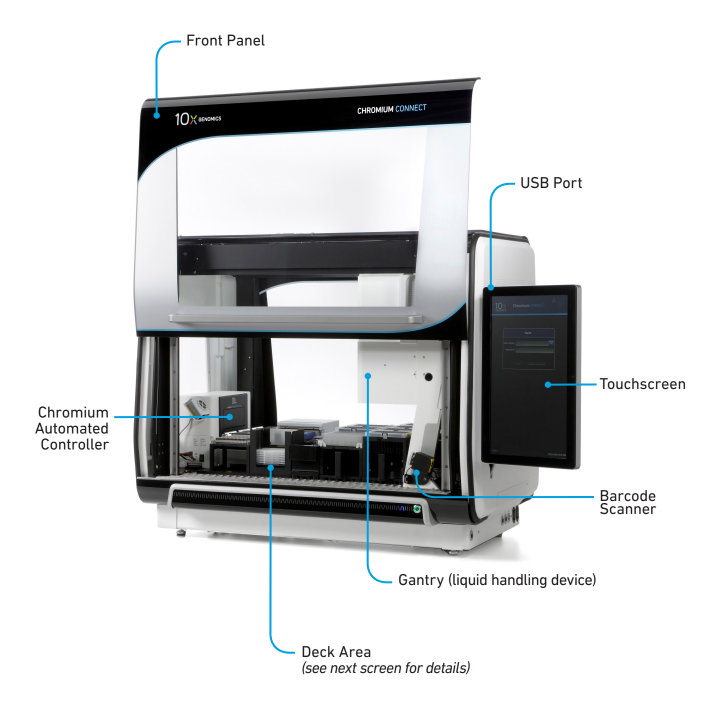


30

Chromium Connect

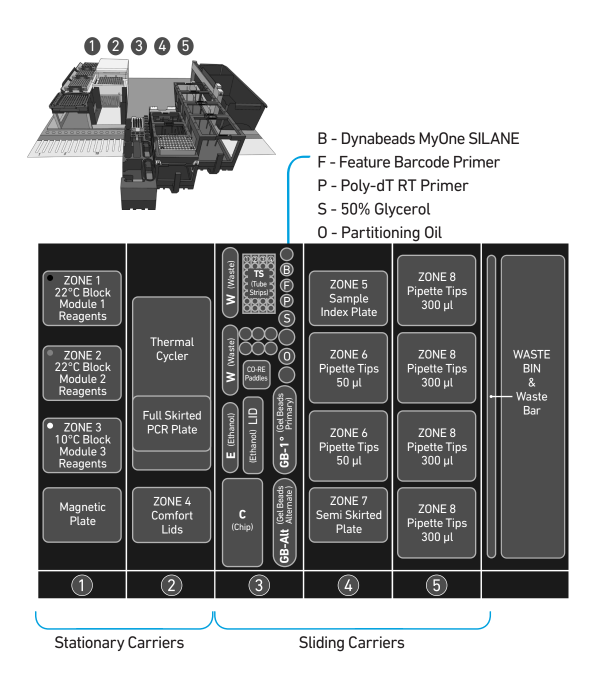
Instrument Orientation Deck Orientation CSV Setup

Instrument Orientation



Refer to the Chromium Connect Instrument User Guide (CG000180) and Quick Reference Cards (CG000256) for more information.

Deck Orientation



Refer to the Chromium Connect Instrument User Guide (CG000180) and Quick Reference Cards (CG000256) for more information.

			igents/Consumab	
ZONE 1 22°C Block Module 1 Reagents		Chromium Next GEN	I Automated Single Ce Zone	ell 5' Gene Expression v2 Assay Item
ZONE 2	Thermal Cycler		Zone 1 (Black)	22°C Block, Reagent Strips, Module 1
22°C Block Module 2 Reagents		1	Zone 2 (Gray)	22°C Block, Reagent Strips, Module 2
• ZONE 3 10°C Block	Full Skirted PCR Plate	l Stationary	Zone 3 (White)	10°C Block, Reagent Strips, Module 3
Module 3 Reagents			-	Magnetic Plate
Magnetic Plate	ZONE 4 Comfort Lids		-	Thermal Cycler
)		2 Stationary	-	Full Skirted PCR Plate (within Thermal Cycler)
1	2		Zone 4	Comfort Lids
			Position W	Waste Reservoirs
			Position TS	Tube Strips (positions 1, 2, & 4)
			Position B	Dynabeads [™] MyOne [™] SILANE
		3*	Position F	Feature Barcode Primer
			Position P	Poly-dT RT Primer
			Position S	50% Glycerol
		Sliding Deck Rails: 15-18	Position 0	Partitioning Oil
		Number of Lights: 4	Position CP	CO-RE Paddles
	3		Position E	Ethanol Reservoir
	ZONE 8		Position Lid	Lid for Ethanol Reservoir
ZONE 5 Sample Index Plate	Pipette Tips 300 µl		Position GB-1°	Gel Beads Primary
ZONE 6	ZONE 8	*Assay choices determine items	Position GB-Alt	Gel Beads Alternate
Pipette Tips 50 µl	Pipette Tips 300 µl	loaded in Carrier 3	Position C	Chip
ZONE 6	ZONE 8	4	Zone 5	Sample Index Plate
Pipette Tips 50 µl	Pipette Tips 300 µl	Sliding Deck Rails: 19-24	Zone 6	Pipette Tips 50 µl
ZONE 7 Semi Skirted	ZONE 8 Pipette Tips	Number of Lights: 6	Zone 7	Semi Skirted Plate
Plate	300 µl	5 Sliding Deck Rails: 25-30 Number of Lights: 6	Zone 8	Pipette Tips 300 μl

CSV Setup

Sample information can also be uploaded using a CSV file at the run setup screen. Use the folder icon to search a network file system or USB drive. Navigate to the appropriate CSV file and click "SELECT".

For 5' Gene Expression Library construction, use Chromium Connect Single Cell 5' Gene Expression Input File (CG000430) and for V(D)J Library construction, use Chromium Connect Single Cell 5' V(D)J Input File (CG000432). For 5' Cell Surface Protein Library construction, use Chromium Connect Single Cell 5' Feature Barcode Input File (CG000535). All the files are available on the 10x Genomics support website.

Alternatively, customer's CSV files can also be generated using the customer's LIMS system. If using a LIMS system to generate CSV files, use ChromiumConnect_SC5-GEX_InputSampleInfo_Template file (CG000429) and for V(D)J Library construction, use ChromiumConnect_SC5-VDJ_InputSampleInfo_Template file (CG000431). For 5' Cell Surface Protein Library construction, use Chromium Connect Single Cell 5' Feature Barcode InputSampleInfo_TemplateFile (CG000536).

SC5' GEX > Setup >	Load > Run	Complete		CHROMIUM CON	INECT 📃
Experiment Name Enter Experiment N	Name		SC3' GEX	Setup > Load >	Run > Complete
Instruction Level	Standard	~			
Run Steps	GEX			Select File	
GEM Check?	Yes	Y			
V(D)J Amplification?	No		■ Z:	Name	
qPCR Setup?	No	~			
Pooling?	No				
			File Name:		CSV files (*.csv) 🗸
					SELECT CANCEL
CANCEL					
			CANCEL	NEXT	

Run Setup Screen

Sample Input Files

Sample input files for Gene Expression, V(D)J Amplification, and Cell Surface Protein (Feature Barcode) are shown below. Refer to the relevant Chromium Connect SC5'-Input Sample Info Template (CG000430/CG000432/CG000535) for more information. The columns highlighted in blue are mandatory to start a run. Any missing fields/ corrections can be added during sample information setup. Final selections will be recorded in the final run report CSV file.



After entering assay i	information belo	w, click this butto	n to	autofill the	LIMS file	Export D	ata to ce
Run Parameters	Selection	Notes (included in run	logs	5)	Legend Blue		
ExperimentName	Sample Run					Enter info manually	
Instruction Level	Standard				Light Blue	Drop-down menu	
GEM Check?	Yes	-			Red	Invalid Entry	
Feature Barcode?	No						
FB Library Construction?	No						
V(D)J Amplification?	Yes						
qPCR Setup?	No						
Pooling?	No						
cDNA Cycles	13						
SI Cycles	14						
Number of Samples	8					1	
Sample Number	Sample ID	Sample Index		CellCount	ExpressionLevel		
ID1	aaa	A1		2001-6000	High		
ID2	bbb	B1		2001-6000	High		
ID3	ccc	C1		2001-6000	High		
ID4	ddd	D1		2001-6000	High		
ID5	eee	E1		2001-6000	High		
ID6	fff	F1		2001-6000	High		
ID7	999	G1		2001-6000	High		
ID8	hhh	H1		2001-6000	High		

V(D)J Sample Input File (CG000432)

After entering as	say information	below, click this b	utton	o autofill	the LIMS file	Export L	ata to csv
Run Parameters	Selection	Notes (included in run	logs)		Legend		
ExperimentName	Sample Run				Blue	Enter info manually	
Instruction Level	Standard				Light Blue	Drop-down menu	
Species-Cell Type?	Human-TCR				Red	Invalid Entry	
qPCR Setup?	No						
Pooling?	No						
Number of Samples	8					-	
Sample Number	Sample ID	Sample Index (A1-H12)	C	llCount	ExpressionLevel		
ID1	aaa	A1	20	01-6000	High		
ID2	bbb	B1	20	01-6000	High		
ID3	ccc	C1	20	01-6000	High		
ID4	ddd	D1	20	01-6000	High		
ID5	eee	E1	20	01-6000	High		
ID6	fff	F1	20	01-6000	High		
ID7	999	G1	20	01-6000	High		
ID8	hhh	H1	20	01-6000	High		
	≤ 32 characters/ symbols/spaces	Rows A-D not accessible for					

Feature Barcode Input File (for cell surface protein; CG000535)

fter entering assay	information bel	ow, click this butt	on	to autofill the	LIMS file	Export D	ata to csv
Run Parameters	Selection	Notes (included in run	log	1	Legend		
ExperimentName	Sample Run	Notes (meladed in run	loga	»/	Blue	Enter info manually	
Instruction Level	Standard				Light Blue	Drop-down menu	
FB DNA Aligout?	No				Red	Invalid Entry	
V(D)J Amplification?	No				Red	Invalid Entry	
qPCR Setup?	No						
Pooling?	No						
SI Cycles	8						
Number of Samples	8						
		Sample Index					
Sample Number	Sample ID	(A1-H12)		CellCount			
ID1	aaa	A1		2001-6000	1		
ID2	bbb	B1		2001-6000			
ID3	ccc	C1		2001-6000			
ID4	ddd	D1		2001-6000			
ID5	eee	E1		2001-6000			
ID6	fff	F1		2001-6000			
ID7	999	G1		2001-6000			
ID8	hhh	H1		2001-6000			
		Rows A-D not					

Sample Input Template Files

Sample input template files for Gene Expression, V(D)J Amplification, and Cell Surface Protein (Feature Barcode) are shown below. Refer to the relevant Chromium Connect SC5'-Input Sample Info Template (CG000429/CG000431/CG000536) for more information. The columns highlighted in blue are mandatory to start a run. Any missing fields/corrections can be added during sample information setup. Final selections will be recorded in the final run report CSV file.

Gene Expression Sample Input Template File (CG000429)

SAMPLEP/ SAMPLEN/ SIINDE	X VOLUME	CellCount Expressior CYCLES	USERDEFI	IUSERDEFI	USERDEFI	USERDEFI	USERDEFI	USERDEFI	USERDEFI	USERDEFI	NED8
ID1											
ID2											
ID3											
ID4											
ID5											
ID6											
ID7											
ID8											
RUNPARA SELECTION											
runName											
Instruction Level											
GEM Check?											
Feature Barcode?											
FB Library Construction?											
V(D)J Amplification?											
qPCR Setup?											
Pooling?											
cDNA Cycles											
SI Cycles											
Notes											

V(D)J Sample Input Template File (CG000431)

SAMPLEP/ SAMPLEN/ SIIN	DEX VOLUME	CellCount Expression	or Cycles	USERDEFIL	JSERDEFII	USERDEFII	USERDEFII	USERDEFII	USERDEFI	USERDEFI	USERDEFIN	NED8
ID1												
ID2												
ID3												
ID4												
ID5												
ID6												
ID7												
ID8												
RUNPARAISELECTION												
runName												
Instruction Level												
Species-Cell Type?												
qPCR Setup?												
Pooling?												
Notes												

Feature Barcode Sample Input Template File (for Cell Surface Protein; CG000536)

V1.30													
SAMPLEP/SAMPLEN/SIINDEX	VOLUME	CellCount	Expressio	r Cycles	USERDEFI	NED8							
ID1													
ID2													
ID3													
ID4													
ID5													
ID6													
ID7													
ID8													
RUNPARAISELECTION													
runName													
Instruction Level													
FB DNA Aligout?													
V(D)J Amplification?													
qPCR Setup?													
Pooling?													
SI Cycles													
Notes													

Uploading Sample Information Using a Template File

The following tables provide specific guidelines on sample entry in the template file.

Sample Parameters	Information
Sample Name	Alphanumeric and up to 32 characters
SI Index	Location on sample index plate to be used for each sample during SI PCR
Expression Level	User defined field for tracking <u>Example:</u> High cell expression: Cell lines Low cell expression: PBMCs
Cell Count	User defined field for tracking (enter applicable option EXACTLY as shown below) 500-2000 2001-6000 6001-10000 DO NOT use commas. Space between symbol & number required.

Up to four user-defined fields (LIMS data) can be passed through the instrument for additional sample tracking.

Run Parameters	Information
Run Name	Alphanumeric and up to 32 characters
Instruction Level	Standard, Advanced, Expert Refer to the Chromium Connect Instrument User Guide (CG000180) for details
Run Steps	GEX/cDNA only
GEM Check	Opt-in for optional QC step: Yes/No
Feature Barcode	Opt-in for optional assay step: Yes/No
Feature Barcode Library Construction	Opt-in for optional assay step: Yes/No
V(D)J Amplification	Opt-in for optional assay step: Yes/No
qPCR Setup	Opt-in for optional assay step: Yes/No
Pooling	Opt-in for optional assay step: Yes/No

Items & Reagents for cDNA/DNA Amplification and 5' GEX Library Construction

Gather Items & Reagents

Follow prompts on the Chromium Connect touchscreen to gather the listed items and reagents for loading the Deck Carriers.

Gather the quantities specified for each of the items and reagents.

Item	Qty
Nuclease-free Water	10 ml
Ethanol, Pure (200 Proof, anhydrous)	40 ml
Hamilton	
Comfort Lids	6
50 µl CO-RE/CO-RE II Pipette Tips, with filter (Black, Conductive)	2 racks
300 µl CO-RE/ CO-RE II Pipette Tips, with filter (Black, Conductive)	4 racks
Reagent Reservoir, 60 ml	3
Eppendorf	
96-well Semi Skirted Plate	1
96-well Full Skirted Plate	1
Thermo Fisher Scientific	
MicroAmp 8-Tube Strip, 0.2 ml	2 (3 if generating Cell Surface Protein library)
10x Genomics	
Chromium Next GEM Chip K Automated Single Cell Kit (stored at room temperature) Partitioning Oil 50% Glycerol Chip K (keep chip sealed)	1
Chromium Next GEM Automated Single Cell 5' Gel Bead Kit v2 (stored at -80°C) Single Cell VDJ 5' Gel Bead v2	1 tube/sample
Chromium Next GEM Automated Single Cell 5' Kit v2	
Module 1 (stored at 4°C) Black tube strip Dynabeads	1 tube strip/sample 1 tube/run
Module 2 (stored at -20°C) Gray tube strip	1 tube strip/sample
Module 3 (stored at -20°C) White tube strip Poly-dT RT Primer	1 tube strip/sample 1 tube/run
Automated 5' Feature Barcode Kit (stored at -20°C) Feature cDNA Primer 4	1 tube/run
Dual Index Plate TT Set A (stored at -20°C) Verify name & PN	1 plate

See Additional Kits, Reagents & Equipment list for performing optional assays and/or QC.

Thaw & Prep Reagents

Follow prompts on the touchscreen to thaw and prepare reagents. Some important guidelines are highlighted below.

ACTION	GUIDELINES Follow touchscreen prompts for specifics and timing
Thaw Reagents	• Thaw reagents as indicated on the touchscreen. Verify no precipitate is present.
	 Ensure that the correct thawing locations and temperatures are used.
	 During reagent thaw load the consumables following touchscreen prompts.
Prepare Ethanol	• Prepare 50 ml 80% Ethanol in Nuclease-free water and dispense in Ethanol Reservoir when prompted.
Poly-dT RT Primer	• Vortex only when prompted on the touchscreen.
	Centrifuge briefly before loading.
Feature cDNA	• Vortex only when prompted on the touchscreen.
Primer 4	Centrifuge briefly before loading.
Dynabeads	• Vortex Dynabeads for \geq 30 sec .
	 Pipette mix to resuspend completely by using a 200 µl pipettor set to 150 µl; pipette mix at least 20 times until fully resuspended. DO NOT centrifuge.
	Confirm there are no bubbles at the bottom of the tube.
Library Modules	Thaw Library Modules as prompted on the touchscreen.
	 After reagent thaw, invert rack holding Module tube strips and vortex Library Modules 1 and 2 for 30 sec; verify no precipitate.
	 Confirm there are no bubbles at the bottoms of any module tubes
	 Centrifuge Library Modules 1 and 2 at 300 rcf for 1 min at 22°C.
	• Retrieve Library Module 3 from 4°C thaw. DO NOT vortex. Invert-mix and centrifuge at 300 rcf for 1 min at 22°C .



Confirm that there are no bubbles at the bottoms of any module tubes, Dual Index Plate wells, or Primer tubes.

Thaw & Prep Reagents

Follow prompts on the touchscreen to thaw and prepare reagents. Some important guidelines are highlighted below.

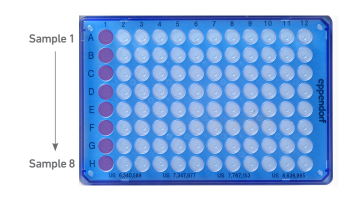
ACTION	GUIDELINES Follow touchscreen prompts for specifics and timing				
Prepare Gel Beads	 Equilibrate the Gel Beads for 30 min at room temperature before use. 				
	 Snap the tube strip holder with the Gel Bead strip into a 10x Vortex Adapter. Vortex 30 sec. 				
	 Centrifuge the Gel Bead strip for ~5 sec after removing from the holder. Confirm there are no bubbles at the bottom of the tubes and the liquid levels look even. 				
	 Place the Gel Bead strip back in the holder and secure the holder lid. 				
	Store unused Gel Beads at –80°C and avoid more than 12 freeze-thaw cycles. DO NOT leave Gel Beads at room temperature for >24 h .				
	 Remove Gel Beads from the Deck during any of the QCs and store the holder with the unused Gel Beads at -80°C. 				

Sample Preparation Guidelines

Sample Preparation Guidelines

- Resuspend samples in PBS + 0.04% BSA (or alternative buffers specified in the relevant 10x Genomics Demonstrated Protocols CG000053/CG000149/CG000203). Refer to the Cell Suspension Volume Calculator Table for the cell suspension and buffer volumes. Total volume loaded per sample onto the sample plate is 10 µl.
 - Based on cell stock concentration, do sequential stock dilutions, if needed.
 - It is recommended to use 3 independent cell counts to determine cell concentration.
 - The presence of dead cells in the suspension may also reduce the recovery rate. Consult the 10x Genomics Single Cell Protocols Cell Preparation Guide and the Guidelines for Optimal Sample Preparation flowchart (Documents CG000053 and CG000126, respectively) for more information on preparing cells.
 - The cell load impacts PCR cycle numbers for cDNA amplification and other downstream steps in the assay. Refer to Additional Protocol Guidelines chapter for more information.

	Manual	Automated
Sample Prep	Using 10x Genomics Demonstrated	Protocols for cell prep and QC
Sample Input Volume	Up to 38.7 µl	10 µl
Sample Loading	PCR strip tubes	96-well skirted plate
Samples per Chip	1-8	1-8
Samples Tested	Various	Human PBMCs, mouse PBMCs, mouse splenocytes, human melanoma



Samples are loaded in column 1, starting at A1. It is not necessary to add glycerol to unused sample wells when running <8 samples.

Cell Suspension Volume Calculator Table

(Chromium Connect Automated Single Cell 5' v2 protocol)

Volume of Cell Suspension Stock per reaction (µl) | Volume of PBS + 0.04% BSA (µl)

Cell Stock						ed Cell Re			4		
Concentration (cells/ µl)	500	1000	2000	3000	4000	5000	6000	7000	8000	9000	10000
100	8.50 1.50	n/a									
200	4.25 5.75	8.50 1.50	n/a								
300	2.83 7.17	5.67 4.33	n/a								
400	2.13 7.88	4.25 5.75	8.50 1.50	n/a							
500	1.70 8.30	3.40 6.60	6.80 3.20	n/a							
600	1.42 8.58	2.83 7.17	5.67 4.33	8.50 1.50	n/a						
700	1.21 8.79	2.43 7.57	4.86 5.14	7.29 2.71	9.71 0.29	n/a	n/a	n/a	n/a	n/a	n/a
800	1.06 8.94	2.13 7.88	4.25 5.75	6.38 3.63	8.50 1.50	n/a	n/a	n/a	n/a	n/a	n/a
900	0.94 9.06	1.89 8.11	3.78 6.22	5.67 4.33	7.56 2.44	9.44 0.56	n/a	n/a	n/a	n/a	n/a
1000	0.85 9.15	1.70 8.30	3.40 6.60	5.10 4.90	6.80 3.20	8.50 1.50	n/a	n/a	n/a	n/a	n/a
1100	0.77 9.23	1.55 8.45	3.09 6.91	4.64 5.36	6.18 3.82	7.73 2.27	9.27 0.73	n/a	n/a	n/a	n/a
1200	0.71 9.29	1.42 8.58	2.83 7.17	4.25 5.75	5.67 4.33	7.08 2.92	8.50 1.50	9.92 0.08	n/a	n/a	n/a
1300	0.65 9.35	1.31 8.69	2.62 7.38	3.92 6.08	5.23 4.77	6.54 3.46	7.85 2.15	9.15 0.85	n/a	n/a	n/a
1400	0.61 9.39	1.21 8.79	2.43 7.57	3.64 6.36	4.86 5.14	6.07 3.93	7.29 2.71	8.50 1.50	9.71 0.29	n/a	n/a
1500	0.57 9.43	1.13 8.87	2.27 7.73	3.40 6.60	4.53 5.47	5.67 4.33	6.80 3.20	7.93 2.07	9.07 0.93	n/a	n/a
1600	0.53 <mark>9.47</mark>	1.06 8.94	2.13 7.88	3.19 6.81	4.25 5.75	5.31 4.69	6.38 3.63	7.44 2.56	8.50 1.50	9.56 0.44	n/a
1700	0.50 9.50	1.00 9.00	2.00 8.00	3.00 7.00	4.00 6.00	5.00 5.00	6.00 4.00	7.00 3.00	8.00 2.00	9.00 1.00	10.00 0.00
1800	0.47 9.53	0.94 9.06	1.89 8.11	2.83 7.17	3.78 6.22	4.72 5.28	5.67 4.33	6.61 3.39	7.56 2.44	8.50 1.50	9.44 0.56
1900	0.45 9.55	0.89 9.11	1.79 8.21	2.68 7.32	3.58 6.42	4.47 5.53	5.37 4.63	6.26 3.74	7.16 2.84	8.05 1.95	8.95 1.05
2000	0.43 9.58	0.85 9.15	1.70 8.30	2.55 7.45	3.40 6.60	4.25 5.75	5.10 4.90	5.95 4.05	6.80 3.20	7.65 2.35	8.50 1.50

Grey boxes: Yellow boxes: Volumes that would exceed the allowable buffer volume in each reaction

Blue boxes:

Indicate a low transfer volume that may result in higher cell load variability

Optimal range of cell stock concentration to maximize the likelihood of achieving the desired cell recovery target

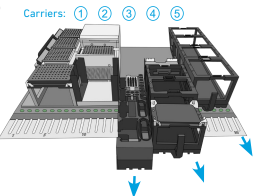
Carrier Loading Guidelines

Carrier Loading Guidelines

Follow the instructions on the touchscreen to load the carriers.

Carriers

- Handle the carriers as prompted.
- Ensure that Carriers 3, 4, and 5 are completely slid out and placed on an off-deck workspace before loading.
- Align the carriers to the corresponding Deck Rails when sliding them in or out of the deck.
- Ensure correct orientation of tube labels with barcodes to enable Barcode Scanning.



Barcode Orientation

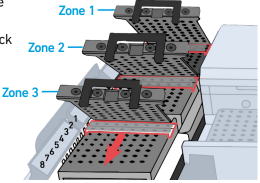


Modules

- Load one tube strip/sample of each of the indicated modules in the corresponding positions on the Carrier, starting from back to front row.
- DO NOT skip any rows when loading.
- Use pinhole alignment to place module tube strips in the correct orientation (as shown on the touchscreen).

Label Tube Strip Orientation

- The cDNA tube strip will be at Position 1 and the final library tube strip will be at Position 4 in the Tube Strip Holder.
- Amplified DNA derived from the Feature Barcode associated with the cell surface protein will be at Position 2.
- Label tube strip orientation for collecting DNA and final libraries.



Tube Strip (TS) Holder



Consult the Chromium Connect User Guide (CG000180) for more information.

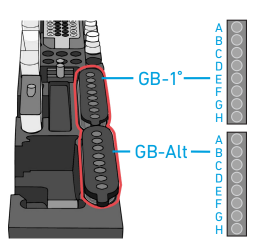
Carrier Loading Guidelines

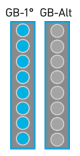
Load Gel Beads

- Up to 2 Gel Bead tube strips may be loaded in the primary (GB-1°)and alternate (GB-Alt) positions. One Gel Bead tube is required/sample.
- If only loading one tube strip, load in the primary position.
- Select the location of the loaded Gel Bead tube/s on the touchscreen.
- Examples of various Gel Bead loading combinations are illustrated below.

Example 1

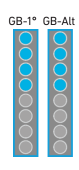
8 samples run with 1 Gel Bead tube strip loaded in GB-1° location.





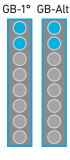
Example 2

8 samples run with 2 Gel Bead tube strips loaded in GB-1° and GB-Alt locations.



Example 3

4 samples run with 2 Gel Bead tube strips loaded in GB-1° and GB-Alt locations.



Additional Protocol Guidelines

Confirm GEM Generation cDNA Amplification Cycles cDNA QC & Quantification

50

Confirm GEM Generation

- Instrument will pause for **5 min** during GEM QC.
- Carefully remove Full Skirted Plate from the Thermal Cycler.
- Hold up the Full Skirted Plate and view the bottom of the wells in Column 3 to confirm GEM generation (shown below).

Column 3

Reload Full Skirted Plate in the Thermal Cycler.



cDNA Amplification Cycle Number

- cDNA amplification cycles are determined by target cell number.
- Recommended guidelines for selecting optimal amplification cycle numbers

	Recommended starting point for cy	cle number optimization.
Targeted Cell Recovery	<u>Low RNA</u> <u>Content Cells</u> <u>e.g., Primary Cells</u> Total Cycles	<u>High RNA Content Cells</u> <u>e.g., Cell Lines</u> Total Cycles
500-2,000	16	14
2,001-6,000	14	12
6,001-10,000	13	11

• The optimal number of cycles is a trade-off between generating sufficient final mass for library construction and minimizing PCR amplification artifacts. The number of cDNA cycles should also be reduced if large numbers of cells are sampled.



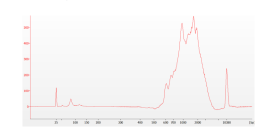
Cycle number selected for one sample will apply to all the other samples in the run.

cDNA QC & Quantification

- a. Follow the instruction on the touchscreen for cDNA QC & quantification.
- **b.** Run sample on an Agilent Bioanalyzer High Sensitivity chip.
 - Run 1 µl undiluted product for input cells with low RNA content (<1 pg total RNA/cell), and 1 µl of 1:10 diluted product for input cells with high RNA content.

Representative Trace for PBMCs

For V(D)J + GEX Library Construction proceed directly to GEX Library Construction first, followed by V(D)J Amplification and V(D)J Library Construction. If GEX library is not desired, proceed directly to V(D)J Amplification.

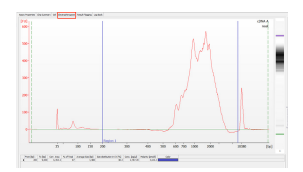


- c. If proceeding to 5' GEX Library Construction, determine cDNA yield for each sample. Example calculation below.
- **d.** Enter the cDNA concentration $(pg/\mu l)$ and the calculated input volume (μl) on the touchscreen to proceed with GEX library construction.

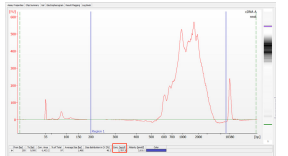
EXAMPLE CALCULATION

i. Select Region

Under the "Electropherogram" view choose the "Region Table". Manually select the region of $\sim 200 - \sim 9000$ bp



ii. Note Concentration [pg/µl]



Alternate Quantification Method:

LabChip (See Appendix for representative traces)

iii. <u>Calculate</u> Concentration: 2787.20 pg/μl Dilution Factor: 1 cDNA Conc. = <u>Conc. (pg/μl) x Dilution Factor</u> = <u>2787.20 x 1</u> =2.79 ng/μl 1000 (pg/ng) 1000

Example Calculation for Carrying Forward 60 ng Sample for 5' GEX Library Construction

Volume for 60 ng =
$$\frac{60 \text{ ng}}{2.79 (\text{ng}/\mu)}$$
 = 21.5 µl

- If the volume for 60 ng exceeds 22 μl , carry ONLY 22 $\mu l\,$ sample into library construction. The sample input volume should be in the 5-22 μl range.

Sample volume for = library construction

= 21.5 µl

If <60* ng available, carry forward 22 μl sample (2-60 ng) into 5' GEX Library Construction.

*Note that the intended sample amount differs from manual protocol to account for pipetting differences in automation.



DO NOT exceed a mass of 60 ng in the 22 μl carry forward volume.

5' Gene Expression (GEX) Library Construction Guidelines

Sample Index PCR Post Library Construction QC

Sample Index PCR

- The cycle numbers can be manually selected based on cDNA input.
- Recommended guidelines for selecting optimal Sample Index PCR cycle number.

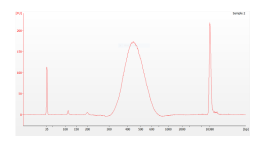
cDNA Input	Total Cycles
1-30 ng	16
31-60 ng	14



Cycle number selected will apply to all the samples in the run.

Post Library Construction Run **1** µl sample on an Agilent Bioanalyzer High Sensitivity chip. QC

Representative Trace



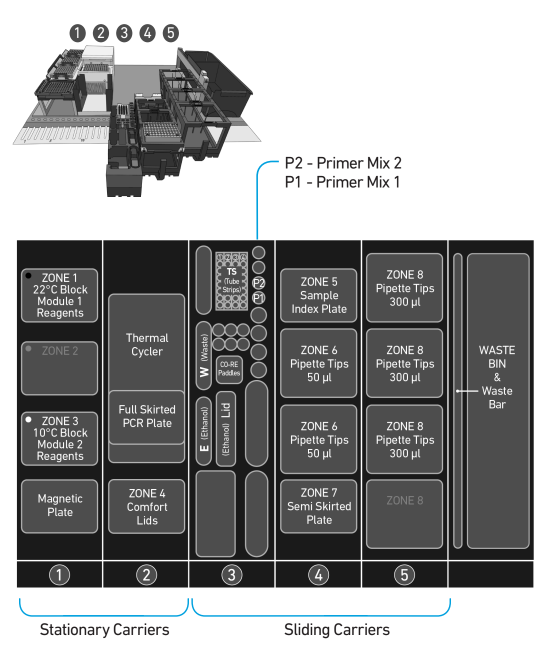
Determine the average fragment size from the Bioanalyzer trace. This will be used as the insert size for library quantification.

Alternate QC Method:

• LabChip (See Appendix for representative traces)

V(D)J Amplification & Library Construction Guidelines

Deck Orientation for V(D)J Amplification Gather Items & Reagents Thaw & Prep Reagents V(D)J Amplification Post Library Construction QC Deck Orientation – V(D)J Amplification & Library Construction



Refer to the Chromium Connect Instrument User Guide (CG000180) and Quick Reference Cards (CG000256) for more information.

ZONE 1 22°C Block			agents/Consumab Automated Single Cell	les V(D)J Amplification & Library Construction
Module 1 Reagents Therm Cycle	Thermal	Carrier	Zone	Item
	Cycler	1 Stationary	Zone 1 (Black)	22°C Block, Reagent Strips, Module 1
 ZONE 3 	Full Skirted PCR Plate		Zone 3 (White)	10°C Block, Reagent Strips, Module 2
10°C Block Module 2 Reagents			-	Magnetic Plate
Magnetic	ZONE 4 Comfort	2	-	Thermal Cycler
Plate	Lids	Stationary	-	Full Skirted PCR Plate (within Thermal Cycler)
1	2		Zone 4	Comfort Lids
		3*	Position W	Waste Reservoirs
	(atsew) M (atsew) M (atsew	Sliding Deck Rails: 15-18	Position TS	Tube Strips (positions 1 & 4)
		*Assay choices determine items loaded in Carrier 3	Position P2	Primer Mix 2
	E (Ethanol) (Ethanol) Lid		Position P1	Primer Mix 1
	E		Position CP	CO-RE Paddles
			Position E	Ethanol Reservoir
	3		Position Lid	Lid for Ethanol Reservoir
ZONE 5 Sample	ZONE 8 Pipette Tips	4 Sliding Deck Rails: 19-24 Number of Lights: 6	Zone 5	Sample Index Plate
Index Plate	300 µl		Zone 6	Pipette Tips 50 µl
20NE 6 Pipette Tips 50 µl			Zone 7	Semi Skirted Plate
ZONE 6 Pipette Tips 50 µl ZONE 7 Semi Skirted Plate ZONE 8 Pipette Tips 300 µl ZONE 8 Pipette Tips 300 µl	Pipette Tips 300 µl	5 Sliding Deck Rails: 25-30 Number of Lights: 6	Zone 8	Pipette Tips 300 µl

Gather Items & Reagents for V(D)J Amplification and Library Construction

Follow prompts on the Chromium Connect touchscreen to gather the listed items and reagents for loading the Deck Carriers. Gather the quantities specified for each of the items and reagents.

ltem	Qty
Nuclease-free Water	10 ml
Ethanol, Pure (200 Proof, anhydrous)	40 ml
Hamilton	
Comfort Lids	6
50 µl CO-RE/CO-RE II Pipette Tips, with filter (Black, Conductive)	2 racks
300 µl CO-RE/CO-RE II Pipette Tips, with filter (Black, Conductive)	3 racks
Reagent Reservoir, 60 ml	2
Eppendorf	
96-well Semi Skirted Plate	1
96-well Full Skirted Plate	1
Thermo Fisher Scientific	
MicroAmp 8-Tube Strip, 0.2 ml	2
10x Genomics	
Chromium Automated Single Cell Human TCR Amplification & Library Construction v2	
V(D)J Module 1 (stored at 4°C) Black tube strip	1 tube strip/sample
V(D)J Module 2 (stored at -20°C) White tube strip Human T Cell Primer Mix 1 v2 Human T Cell Primer Mix 2 v2	1 tube strip/sample 1 tube/run 1 tube/run
Chromium Automated Single Cell Mouse TCR Amplification & Library Construction v2	
V(D)J Module 1 (stored at 4°C) Black tube strip	1 tube strip/sample
V(D)J Module 2 (stored at -20°C) White tube strip Mouse T Cell Primer Mix 1 v2 Mouse T Cell Primer Mix 2 v2	1 tube strip/sample 1 tube/run 1 tube/run
Chromium Automated Single Cell Human BCR Amplification & Library Construction v2	
V(D)J Module 1 (stored at 4° C) Black tube strip	1 tube strip/sample
V(D)J Module 2 (stored at –20°C) White tube strip Human B Cell Primer Mix 1 v2 Human B Cell Primer Mix 2 v2	1 tube strip/sample 1 tube/run 1 tube/run
Chromium Automated Single Cell Mouse BCR Amplification & Library Construction v2	
V(D)J Module 1(stored at 4°C) <i>Black tube strip</i>	1 tube strip/sample
V(D)J Module 2 (stored at -20°C) White tube strip Mouse B Cell Primer Mix 1 v2 Mouse B Cell Primer Mix 2 v2	1 tube strip/sample 1 tube/run 1 tube/run
Dual Index Plate TT Set A (stored at -20°C) Verify name & PN	1 plate

See Additional Kits, Reagents & Equipment

list for performing optional assays and/ or QC.

Thaw & Prep Reagents for V(D)J Amplification & Library Construction

Follow prompts on the touchscreen to thaw and prepare reagents. Some important guidelines are highlighted below.

ACTION	GUIDELINES Follow touchscreen prompts for specifics and timing
Thaw Reagents	 Thaw reagents as indicated on the touchscreen. Verify no precipitate is present.
	 Ensure that the correct thawing locations and temperatures are used.
	• During reagent thaw load the consumables following touchscreen prompts.
Prepare Ethanol	 Prepare 50 ml 80% Ethanol in Nuclease-free water and dispense in Ethanol Reservoir when prompted.
V(D)J Modules	• Thaw V(D)J Modules as prompted on the touchscreen.
	 After reagent thaw, invert rack holding Module tube strips and vortex V(D)J Modules 1 for 30 sec; verify no precipitate.
	 Confirm that there are no bubbles at the bottoms of any module tubes.
	 Centrifuge V(D)J Module 1 at 300 rcf for 1 min at 22°C.
	 Retrieve V(D)J Module 2 from 4°C thaw. DO NOT vortex. Invert-mix and centrifuge at 300 rcf for 1 min at 22°C.
Dual Index Plate TT Set A	 Vortex Dual Index Plate for 15 sec at maximum speed and centrifuge at 300 rcf for 1 min at 22°C.
Primer Mix 1 & 2	Vortex and centrifuge before loading.

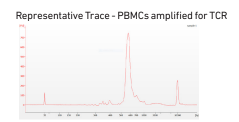


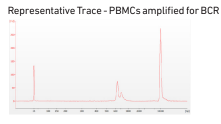
Confirm that there are no bubbles at the bottoms of any module tubes, Dual Index Plate wells, or Primer Mix tubes.

V(D)J Amplification QC & Quantification

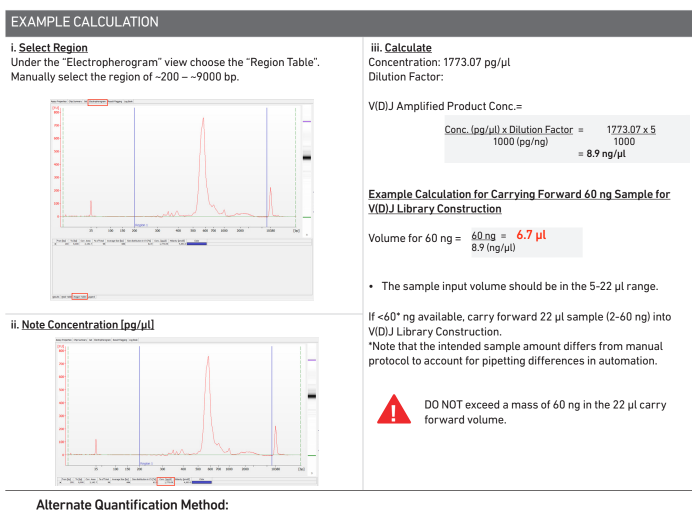
- a. Follow the instruction on the touchscreen for V(D)J Amplification QC & quantification.
- **b.** Run **1** μ**l** sample at **1:5 dilution** (Dilution Factor 5) on an Agilent Bioanalyzer High Sensitivity chip.

Samples of RNA-rich cells may require additional dilution in nuclease-free water. The number of distinct peaks may vary. Higher molecular weight product (2,000- 9,000 bp) may be present. This does not affect sequencing.





- c. Determine yield for each sample using the example calculation below.
- **d.** Enter the V(D)J amplified product concentration $(pg/\mu l)$ and the calculated input volume (μl) on the touchscreen to proceed with V(D)J library construction.



LabChip (See Appendix for representative traces)

Post Library ConstructionRun sample on an Agilent Bioanalyzer High Sensitivity chip.QC

Representative Trace - PBMCs amplified for TCR

Representative Trace

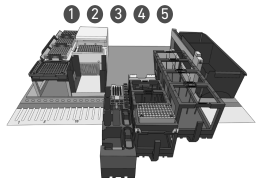
Determine the average fragment size from the Bioanalyzer trace. This will be used as the insert size for library

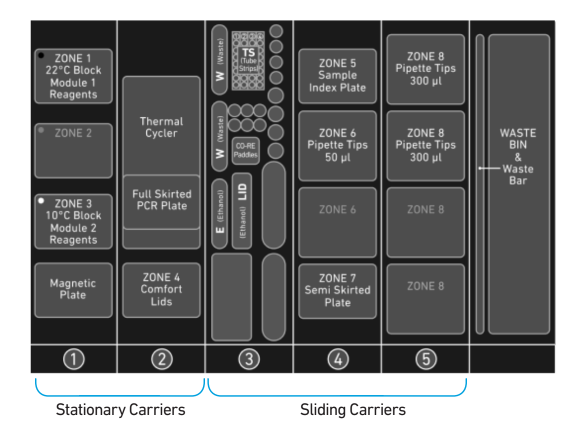
Alternate QC Methods

• LabChip (See Appendix for representative traces)

Cell Surface Protein/ Immune Receptor Mapping Library Construction Guidelines

Deck Orientation Gather Items & Reagents Thaw & Prep Reagents Automated DNA Supernatant – SPRIselect Cleanup Sample Index PCR Post Library Construction QC Deck Orientation – Cell Surface Protein/ Immune Receptor Library Construction





Refer to the Chromium Connect Instrument User Guide (CG000180) and Quick Reference Cards (CG000256) for more information.

ZONE 1 22°C Block		Deck Layout Reagents/Consumables Chromium Next GEM Automated Single Cell 5' Cell Surface Protein Library Construction		
Module 1 Reagents Thermal	Carrier	Zone	ltem	
ZONE 2 Cycler		Zone 1 (Black)	22°C Block, Reagent Strips, Module 1	
ZONE 3 Full Skirte PCR Plate		Zone 3 (White)	10°C Block, Reagent Strips, Module 2	
● ZONE 3 10°C Block Module 2 Reagents		-	Magnetic Plate	
Magnetic ZONE 4 Dista	2	-	Thermal Cycler	
Plate	Stationary	-	Full Skirted PCR Plate (within Thermal Cycler)	
1 2		Zone 4	Comfort Lids	
(Master) SL 200 C 21 C 21 C 200 C 20	3* Sliding	Position W	Waste Reservoirs	
	Deck Rails: 15-18 Number of Lights: 4	Position TS	Tube Strips (positions 1 & 4)	
asew Co-RE Paddes		Position CP	CO-RE Paddles	
E (Ethanol)		Position E	Ethanol Reservoir	
	\leq	Position Lid	Lid for Ethanol Reservoir	
	4	Zone 5	Sample Index Plate TN Set A Verify name & PN	
	Deck Rails: 19-24	Zone 6	Pipette Tips 50 µl	
ZONE 5 Sample Index Plate	Number of Lights: 6	Zone 7	Semi Skirted Plate	
ZONE 6 Pipette Tips 50 µl ZONE 6 Pipette Tips 50 µl ZONE 8 Pipette Tips 50 µl ZONE 8 Pipette Tips 300 µl	Sliding Deck Rails: 25-30	Zone 8	Pipette Tips 300 µl	
ZONE 7 Semi Skirted Plate				

Gather Items & Reagents for Cell Surface Protein/ Immune Receptor Mapping Library Construction Follow prompts on the Chromium Connect touchscreen to gather the listed items and reagents for loading the Deck Carriers. Gather the quantities specified for each of the items and reagents.

ltem	Qty
Nuclease-free Water	10 ml
Ethanol, Pure (200 Proof, anhydrous)	40 ml
Hamilton	
Comfort Lids	2
$50\mu l$ CO-RE/CO-RE II Pipette Tips, with filter (Black, Conductive)	1 rack
$300\ \mu l$ CO-RE/CO-RE II Pipette Tips, with filter (Black, Conductive)	1-2 racks
Reagent Reservoir, 60 ml	3
Eppendorf	
96-well Semi Skirted Plate	1
96-well Full Skirted Plate	1
Thermo Fisher Scientific	
MicroAmp 8-Tube Strip, 0.2 ml	2
10x Genomics	
Chromium Automated Single Cell 5' Feature Barcode Library Construction Kit	
Automated Feature Barcode Library Construction Kit, Module 1 (stored at 4°C) <i>Black tube strip</i>	1 tube strip/sample
Automated Feature Barcode Library Construction Kit, Module 2 (stored at -20°C) <i>White tube strip</i>	1 tube strip/sample
Dual Index Plate TN Set A (stored at -20°C) Verify name & PN	1 plate

See Additional Kits, Reagents & Equipment list for performing optional assays and/ or QC. Thaw & Prep Reagents for Cell Surface Protein/ Immune Receptor Mapping Library Construction Follow prompts on the touchscreen to thaw and prepare reagents. Some important guidelines are highlighted below.

ACTION	GUIDELINES Follow touchscreen prompts for specifics and timing
Thaw Reagents	 Thaw reagents as indicated on the touchscreen. Verify no precipitate is present.
	 Ensure that the correct thawing locations and temperatures are used.
	 During reagent thaw load the consumables following touchscreen prompts.
Prepare Ethanol	 Prepare 50 ml 80% Ethanol in Nuclease-free water and dispense in Ethanol Reservoir when prompted.
Feature Barcode Library Construction	Thaw Modules as prompted on the touchscreen.Confirm that there are no bubbles at the bottoms of any
Modules	module tubes.
Dual Index Plate TN Set A	Use the indicated plate. Verify name & PN.
Set A	 Vortex Dual Index Plate for 15 sec at maximum speed and centrifuge at 300 rcf for 1 min at 22°C.
	- hubbles at the betterne of any medule types and Duel Index

Confirm that there are no bubbles at the bottoms of any module tubes and Dual Index Plate wells.

Automated Feature Barcode DNA – SPRIselect Cleanup

- The first step of Cell Surface Protein/Immune Receptor Mapping Library Construction is SPRIselect Cleanup of the Amplified Feature Barcode DNA. After the SPRIselect Cleanup step, the excess DNA volume that is not required for automated Sample Index PCR may be removed from the instrument based on the instrument touchscreen prompts.
- The Feature Barcode DNA aliquot can be removed and stored at 4°C for up to 72 h or at -20°C for up to 4 weeks and may be used for additional Cell Surface Protein/ Immune Receptor Mapping library construction using the manual Chromium Next GEM Single Cell 5' Reagent Kits v2 with Feature Barcode technology for Cell Surface Protein & Immune Receptor Mapping workflow (CG000330).



- The Feature Barcode DNA aliquot cannot be used for generation of additional libraries using the automated workflow.
- The downstream automated Cell Surface Protein library construction steps are not impacted by whether or not the excess purified Feature Barcode DNA is removed from the instrument.

Sample Index PCR

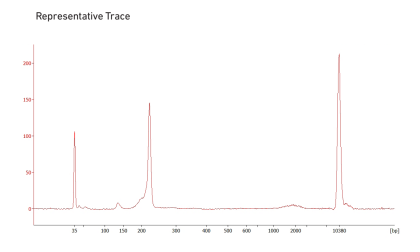
- The default cycle number for Sample Index PCR is 8 cycles.
- The cycle number may be manually changed (7-10 cycles) based on target protein expression levels and number of antibodies used for labeling.



Cycle number selected will apply to all the samples in the run.

Post Library Construction A

Run 1 μ l sample at 1:10 dilution on an Agilent Bioanalyzer High Sensitivity chip.



Determine the average fragment size from the Bioanalyzer trace. This will be used as the insert size for library quantification.

Alternate QC Method

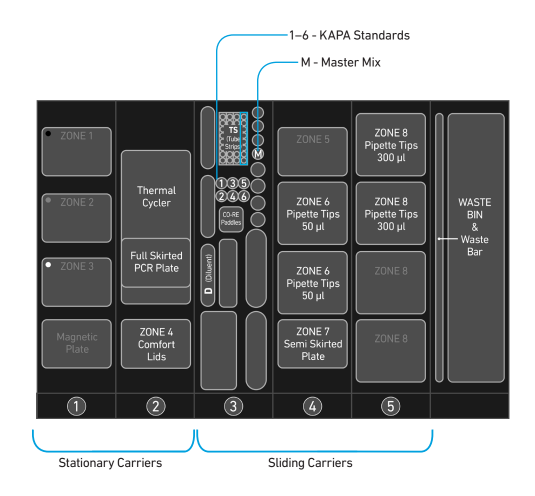
• LabChip (See Appendix for representative traces)

Post Library Construction Quantification & Pooling

Deck Orientation – Library Quantification Post Library Construction Quantification Deck Orientation – Library Pooling Library Pooling

Deck Orientation – Library Quantification

Library quantification using qPCR is recommended for accurate pooling and loading on sequencers. If the option is selected during gene expression run-setup, automated qPCR plate-setup can be run directly on Chromium Connect after library generation and final library QC. Alternatively, the option can be selected from the instrument home screen, at the user's convenience. Up to 8 samples can be quantified on a 96 well reaction plate, including duplicates for each sample. The minimum sample volume required is **25 µl**. Only **6 µl** of the sample will be used for qPCR plate setup.



Gather Items & Reagents

Follow prompts on the Chromium Connect touchscreen to gather the listed items and reagents for loading the Deck Carriers for Library Quantification.

Gather the quantities specified for each of the items and reagents.

ltem	Qty
Hamilton	
Comfort Lid	1
50 µl CO-RE/CO-RE II Pipette Tips, with filter (Black, Conductive)	2 racks
$300\ \mu l$ CO-RE/CO-RE II Pipette Tips, with filter (Black, Conductive)	2 racks
60-ml Reagent Reservoir	1
Eppendorf	
96-well Semi Skirted Plate	1
Thermo Fisher Scientific	
2-ml Tube with Screw Cap	1
Bio-Rad	
96-well Hard-Shell Full Skirted Plate	1
Reagent	Qty
Qiagen Buffer EB	50 ml
Nuclease-free Water	1 ml
10% Tween-20	250 µl
Libraries (in an 8-tube strip)	1-8
KAPA Library Quantification Kit, thawed	
SYBR FAST Master Mix Primer Mix Standards	5 ml 1 ml 6

Post Library Construction Quantification

- Prepare reagents as prompted on the touchscreen.
- Vortex and centrifuge KAPA standards and libraries before use.
- Retrieve previously prepared Master Mix + Primer Mix
 OR

Add 1 ml Primer Mix to 5 ml SYBR FAST Master Mix.

• Prepare specified Quantification Master Mix in the 2-ml tube using the guidance below.

# Sample	Master Mix + Primer Mix (µl)	Water (µl)	Total Vol (µl)
8	1305	435	1740
7	1200	400	1600
6	1095	365	1460
5	990	330	1320
4	885	295	1180
3	780	260	1040
2	675	225	900
1	570	190	760

Volumes listed take into account volume for 6 standards

- Follow the touchscreen prompts for loading, scanning, and executing the run.
- During the run, the following steps will be executed by the instrument:
 -KAPA Master Mix transfer to the 96-well Hard Shell Full Skirted Plate (layout below)
 -Diluent transfer to dilution plate
 - -Serial dilutions of libraries

-Addition of library dilutions, KAPA Standards, and negative controls to the plate

Total reaction volume (20 μl)= 16 μl Master Mix + 4 μl Library Dilution/ KAPA Standard/ Negative Control (NTC)	1 2 3 4 5 6 7 8 9 10 11 12 A B B C B C B B B B B B B B B B B B B B
<u>Dilutions:</u> 1:12,500 1:62,500 1:312,500 1:1,562,500	E C C C C C C C C C C C C C C C C C C C

- After the run is completed, follow the unloading instructions on the touchscreen.
- Cap and store libraries at 4°C ≤72 h or -20°C ≤4 weeks.
- Remove Full Skirted Plate. Seal plate and centrifuge at 300 rcf for 1 min at 22°C.
- Follow the manufacturer's recommendations for qPCR-based quantification. For library quantification for sequencer clustering, determine the concentration based on average size (bp) derived from the Bioanalyzer/TapeStation trace.

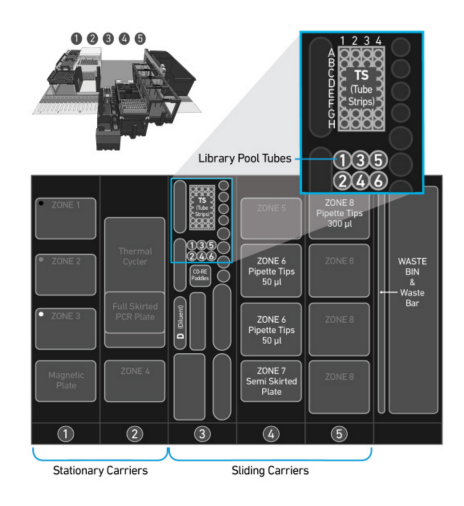
Step	Temperature	Run Time
1	95°C	00:05:00
2	95°C	00:00:30
3	60°C	00:00:45
4	Go to Step 2, 29X (Total 30 cycles)

• Resource:

Use the Chromium Connect Library Quantification Worksheet (CG000157) provided on the 10x Genomics Support website for calculating library concentrations.

Deck Orientation – Library Pooling

The libraries may be pooled on the Chromium Connect instrument and used for sequencing, taking into account the preferred cell numbers and per-cell read depth requirements for each library. Samples utilizing the same sample index should not be pooled together, or run on the same flow cell lane, as this would prevent correct sample demultiplexing. The Chromium Connect deck layout for Library Pooling is shown below.



Gather Items & Reagents

Follow prompts on the Chromium Connect touchscreen to gather the listed items and reagents for loading the Deck Carriers for Library Pooling.

Gather the quantities specified for each of the items and reagents.

ltem	Qty
Hamilton	
50 µl CO-RE/CO-RE II Pipette Tips, with filter (Black, Conductive)	1-2 rack
$300\ \mu l$ CO-RE/CO-RE II Pipette Tips, with filter (Black, Conductive)	1-2 rack
Reagent Reservoir, 60 ml	1
Eppendorf	
96-well Semi Skirted Plate	1
Thermo Fisher Scientific	
0.5-ml Tube with Screw Cap	6
MicroAmp 8-Tube Strip, 0.2 ml	1-4
Reagent	Qty
Qiagen Buffer EB	50 ml
Libraries (in up to four 8-tube strips)	up to 32 libraries

Library Pooling

- Follow the touchscreen prompts for loading, scanning, and executing the run.
- Briefly vortex and centrifuge libraries in the 8-tube strip.
- Confirm that there are no bubbles at the bottoms of any library tubes.
- Ensure a minimum 25 µl library volume is available in the tubes.
- After run is complete, follow touchscreen prompts to unload and store the libraries.
- Unload remaining items and clean as prompted on the touchscreen.
- Resource:

Use the Chromium Connect Library Pooling Worksheet (CG000466) provided on the 10x Genomics Support website to calculate volumes to be pooled. The calculated volumes can be input into the instrument either manually, or via the CSV file generated from this workbook.

Sequencing

Sequencing Libraries

Chromium Single Cell V(D)J, 5' Gene Expression, and Cell Surface Protein Dual Index libraries comprise standard Illumina paired-end constructs which begin with P5 and end with P7. These libraries include 16 bp 10x Barcodes encoded at the start of Read 1. Sample index sequences are incorporated as the i5 and i7 index read for V(D)J, 5' Gene Expression, and Cell Surface Protein libraries.

TruSeg Read 1, TruSeg Read 2, and Nextera Read 2 (Read 2N) are all standard Illumina sequencing primer sites. TruSeq Read 1 and TruSeq Read 2 are used in paired-end sequencing of V(D)J and 5' Gene Expression libraries. TruSeg Read 1 and Nextera Read 2 (Read 2N) are used for paired-end sequencing of Cell Surface Protein library. Sequencing these libraries produce a standard Illumina BCL data output folder.



Illumina Sequencer Compatibility	 The compatibility of the listed sequencers has been verified by 10x Genomics. Some variation in assay performance is expected based on sequencer choice. For more information about performance variation, visit the 10x Genomics Support website. MiSeq NextSeq 500/550/2000 HiSeq 2500 (Rapid Run) HiSeq 3000/4000 NovaSeq
Sample Indices	Each well of the Dual Index Kit TT Set A (PN-1000215) and Dual Index Kit TN Set A (PN-1000250) contains a mix of one unique i7 and one unique i5 sample index. If multiple samples are pooled in a sequence lane, the sample index name (i.e. the Dual Index plate well ID) is needed in the sample sheet used for generating FASTQs with "cellranger mkfastq". If multiple libraries are pooled in a sequence lane, a separate sample index is needed with each library (see Tips & Best Practices).
Click to TOC	Chromium Automated Single Cell 5' Reagent Kits v2 with Feature Barcode technology Cell Surface Protein • Rev A 75

Library Sequencing Depth & Run Parameters	Sequencing Depth	Minimum 5,000 read pairs per cell for V(D)J library Minimum 20,000 read pairs per cell for 5' Gene Expression library Minimum 5,000 read pairs per cell for Cell Surface Protein Dual Index library
	Sequencing Type	Paired-end, Dual indexing
	Sequencing Read	Read 1: 26 cycles i7 Index: 10 cycles i5 Index: 10 cycles Read 2: 90 cycles

Library Loading

Once quantified and normalized, V(D)J, 5' Gene Expression, and Cell Surface Protein libraries should be denatured and diluted as recommended for Illumina sequencing platforms. Refer to Illumina documentation for denaturing and diluting libraries. Refer to the 10x Genomics Support website for more information.

Instrument	Loading Concentration (pM)	PhiX (%)	
MiSeq	10	1	
NextSeq 500	1.5	1	
NovaSeq	150*/300	1	
NextSeq 2000	650	1	

* Use 150 pM loading concentration for Illumina XP workflow.

Library Pooling

V(D)J, 5' Gene Expression, and Cell Surface Protein libraries may be pooled for sequencing, taking into account the differences in depth requirements between the pooled libraries. 5' Gene Expression libraries may be sequenced using enriched library parameters, however the cost of sequencing using enriched library parameters is higher.

Refer to Post Library Construction Quantification & Pooling chapter for library pooling on the Chromium Connect instrument.

Library Pooling Examples:

Libraries	Sequencing Depth (read pairs per cell)	Library Pooling Ratio
Example 1		
V(D)J library 5' Gene Expression library Cell Surface Protein library	5,000 20,000 5,000	1 4 1
Example 2		
V(D)J library 5' Gene Expression library Cell Surface Protein library	5,000 50,000 5,000	1 10 1

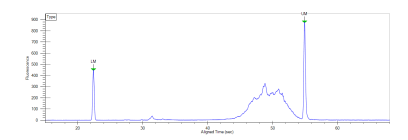
Appendix

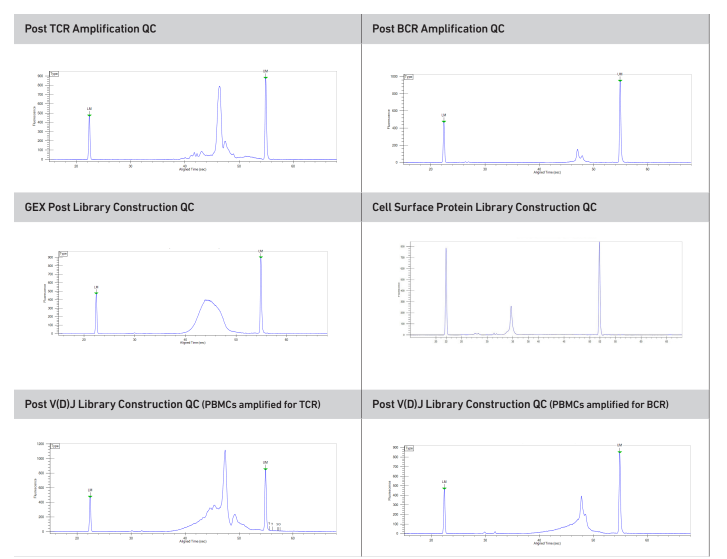
LabChip Traces Oligonucleotide Sequences

LabChip Traces

LabChip Traces DNA High Sensitivity Reagent Kit was used.

cDNA QC & Quantification

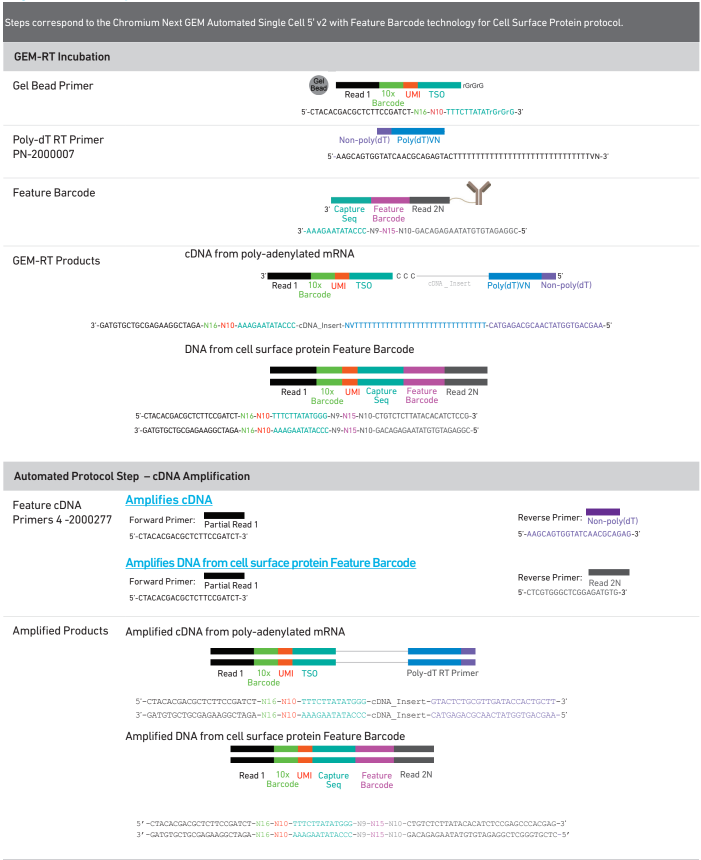




Alternate QC Method: Qubit Fluorometer and Qubit dsDNA HS Assay Kit

Oligonucleotide Sequences Protocol steps correspond to the Chromium Next GEM Automated Single Cell 5' v2 with Feature Barcode technology for Cell Surface Protein protocol. **GEM-RT** Incubation Gel Bead Primer rGrGrG 10x UMI TSO Read 1 Barcode Poly-dT RT Primer Non-poly(dT) Poly(dT)VN PN-2000007 ССС **GEM-RT Products** cDNA Insert 10x UMI TS0 Polv(dT)VN Non-polv(dT) Read 1 Barcode 3'-GATGTGCGCGAGAAGGCTAGA-N16-N10-AAAGAATATACCC-cDNA Insert-NVTTTTTTTTTTTTTTTTTTTTTTTTTTTTCATGAGACGCAACTATGGTGACGAA-5' Automated Protocol Step - cDNA Amplification cDNA Primers Reverse Primer: Non-poly(dT) Forward Primer: Partial Read 1 5'-AAGCAGTGGTATCAACGCAGAG-3' 5'-CTACACGACGCTCTTCCGATCT-3' Amplified Products Amplified cDNA from poly-adenylated mRNA Read 1 10x UMI TSO Barcode Poly-dT RT Primer 5'-CTACACGACGCTCTTCCGATCT-N16-N10-TTTCTTATATGGG-cDNA_Insert-GTACTCTGCGTTGATACCACTGCTT-3' 3'-GATGTGCTGCGAGAAGGCTAGA-N16-N10-AAAGAATATACCC-cDNA_Insert-CATGAGACGCAACTATGGTGACGAA-5' Automated Protocol Step –GEX Adaptor Ligation (for 5' Gene Expression (GEX) Library Construction) Adaptor Read 2 Read 2 5' -GATCGGAAGAGCACACGTCTGAACTCCAGTCAC-3' 3' -TCTAGCCTTCTCG-5' Ligation Product 10x UMI TSO Read 1 Insert Read 2 Barcode 5'-GATCTACACTCTTTCCCTACACGACGCTCTTCCGATCT-N16-N10-TTTCTTATATGGG-cDNA_Insert-AGATCGGAAGAGCACACGTCTGAACTCCAGTCAC-3' 3'-CTAGATGTGAGAAAGGGATGTGCTGCGAGAAGGCTAGA-N16-N10-AAAGAATATACCC-cDNA Insert-TCTAGCCTTCTCG-5' Automated Protocol Step – Sample Index PCR (for 5' Gene Expression (GEX) Library Construction) Dual Indexing Forward Primer: Reverse Primer: P5 Sample Partial Read 1 Partial Read 2 Dual Index TT Set A Index (i5) Index (i7 PN-1000215 5'-AATGATACGGCGACCACCGAGATCT-N10-ACACTCTTTCCCTACACGACGCTC-3' 5'-CAAGCAGAAGACGGCATACGAGAT-N10-GTGACTGGAGTTCAGACGTGT-3' Sample Index PCR Product Sample Read 1 TS0 Read 2 Sample P7 P5 10x UMI Insert Index (i5) Barcode Index (i7) 5'-AATGATAGGGGAGACCACCGAGATCTACAC-NI0-ACACCTCTTCCCCTACCAGGAGGCCCTCTTCCCGATCT-NI6-NI0-TTTCTTATAGGG-CDNA_Insert-AGATCGGAAGAGCACACGTCTGAACTCCAGTCAC-NI0-ATCTCGTTAGCCGTCTTCTCGTT-3 3'-TTACTATGCCGCTGGTGGCTCTAGATGTG-N10-TGTGAGAAAGGGATGTGCTGCGAGAAGGCTAGA-N16-N10-AAAGAATATACCC-cDNA _ Insert-tCTAGCCTCTCGGTGGCAGACTTGAGGTCAGG-N10-

Oligonucleotide Sequences



uman T Cell Mix 1 v2	Forward Primer:		Reverse Outer Primers:	
V-2000242	PCR Prime	er	5'-TGAAGGCGTTTGCACATGCA-3'	Outer Primer
	5'-GATCTACACTCTTTCCCTACACGACGC-3'		5'-TCAGGCAGTATCTGGAGTCATTGAG-3	
Human B Cell Mix 1 v2 PN-2000254	Forward Primer:	-r	Reverse Outer Primers:	Outer Primer
	5'-GATCTACACTCTTTCCCTACACGACGC-3'		5'-CAGGGCACAGTCACATCCT-3' 5'-TGCTGGACCACGCATTTGTA-3'	outer i filliter
	5-GATCIACACICITITCCCIACACGAC	.00-3	5'-GGTTTTGTTGTCGACCCAGTCT-3'	
			5'-TTGTCCACCTTGGTGTTGCT-3' 5'-CATGACGTCCTTGGAAGGCA-3'	
			5'-TGTGGGACTTCCACTG-3'	
			5'-TTCTCGTAGTCTGCTTTGCTCAG-3'	
ouse T Cell Mix 1 v2	Forward Primer:		Reverse Outer Primers:	
N-2000256		PCR Primer	5'-CTGGTTGCTCCAGGCAATGG-3'	Outer Primer
	5'-GATCTACACTCTTTCCCTACACGAC	:GC-3'	5'-TGTAGGCCTGAGGGTCCGT-3'	
ouse B Cell Mix 1 v2	Forward Primer:	PCR Primer	Reverse Outer Primers:	Outer Primer
N-2000258	F/ 0.1707101070777777777		5'-TCAGCACGGGACAAACTCTTCT-3' 5'-GCAGGAGACAGACTCTTCTCCA-3'	eater i finner
	5'-GATCTACACTCTTTCCCTACACGAC	:GC-3'	5'-AACTGGCTGCTCATGGTGT-3'	
			5'-TGGTGCAAGTGTGGTTGAGGT-3'	
			5'-TGGTCACTTGGCTGGTGGTG-3' 5'-CACTTGGCAGGTGAACTGTTTTCT-3'	
			5'-AACCTTCAAGGATGCTCTTGGGA-3'	
			5'-GGACAGGGATCCAGAGTTCCA-3' 5'-AGGTGACGGTCTGACTTGGC-3'	
			3 ACCIONCOCICICACITOCC 3	
			5'-GCTGGACAGGGCTCCATAGTT-3'	
utomated Protocol S	tep – V(D)J Amplification 2		5'-GCTGGACAGGGCTCCATAGTT-3' 5'-GGCACCTTGTCCAATCATGTTCC-3' 5'-ATGTCGTTCATACTCGTCCTTGGT-3'	
uman T Cell Mix 2 v2	tep – V(D)J Amplification 2 Forward Primer:	PCR Primer	5'-GGCACCTTGTCCAATCATGTTCC-3'	Inner Primer
uman T Cell Mix 2 v2			5'-GGCACCTTGTCCAATCATGTTCC-3' 5'-ATGTCGTTCATACTCGTCCTTGGT-3' Reverse Inner Primers:	Inner Primer
uman T Cell Mix 2 v2 N-2000246	Forward Primer:	:GC-3 [,]	5'-GGCACCTTGTCCAATCATGTTCC-3' 5'-ATGTCGTTCATACTCGTCCTTGGT-3' Reverse Inner Primers: 5'-AGTCTCTCAGCTGGTACACG-3'	_
uman T Cell Mix 2 v2 N-2000246 uman B Cell Mix 2 v2	Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC		5'-GGCACCTTGTCCAATCATGTTCC-3' 5'-ATGTCGTTCATACTCGTCCTTGGT-3' Reverse Inner Primers: 5'-AGTCTCTCAGCTGGTACACG-3' 5'-TCTGATGGCTCAAACACAGC-3' Reverse Inner Primers: 5'-GGGAAGTTTCTGGCGGTCA-3'	Inner Primer Inner Primer
uman T Cell Mix 2 v2 N-2000246 uman B Cell Mix 2 v2	Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC	PCR Primer	5'-GGCACCTTGTCCAATCATGTTCC-3' 5'-ATGTCGTTCATACTCGTCCTTGGT-3' Reverse Inner Primers: 5'-AGTCTCTCAGCTGGTACACG-3' 5'-TCTGATGGCTCCAACACAGC-3' Reverse Inner Primers:	_
uman T Cell Mix 2 v2 N-2000246 uman B Cell Mix 2 v2	Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer:	PCR Primer	5'-GGCACCTTGTCCAATCATGTTCC-3' 5'-ATGTCGTTCATACTCGTCCTTGGT-3' S'-AGTCTCCAGCTGGTACACG-3' 5'-AGTCTCTCAGCTGGTACACG-3' S'-GGCAAGTTTCTGGCGGTCA-3' S'-GGTGGTACCCAGTTATCAAGCAT-3' S'-GGTGCCCAGGTCACCATCAC-3' S'-GTGTCCCAGGTCACCATCAC-3' S'-GTGTCCCAGGTCACCATCAC-3'	_
uman T Cell Mix 2 v2 N-2000246 uman B Cell Mix 2 v2	Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer:	PCR Primer	5'-GGCACCTTGTCCAATCATGTTCC-3' 5'-ATGTCGTTCATACTCGTCCTTGGT-3' S'-AGTCTCCAGCTGGTACACG-3' 5'-ATGTCGTCCAACAGCGC-3' 5'-GGGAAGTTTCTGGCGGTCA-3' 5'-GGTGGTACCCAGTTATCAAGCAT-3' 5'-GTGTCCCAGGTCACCATCAC-3'	_
uman T Cell Mix 2 v2 N-2000246 uman B Cell Mix 2 v2	Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer:	PCR Primer	5'-GGCACCTTGTCCAATCATGTTCC-3' 5'-ATGTCGTTCATACTCGTCCTTGGT-3' S'-AGTCTCCAGCTGGTACACG-3' 5'-GGCACCTCCAGCTGGTACACG-3' 5'-TCTGATGGCTCAAACACAGC-3' S'-GGGGAACTTTCTGGCGGTCA-3' S'-GGTGGTACCCAGTTATCAAGCAT-3' S'-GTGTCCCAGGTCACCATCAC-3' S'-GTGTCCCAGGTCACCATCAC-3' S'-GTGTCCCAGGTCACCATCAC-3' S'-TCCTGAGGACCGTCAGGACAGC-3' S'-CCTGAGGACTGTAGGACAGC-3'	_
uman T Cell Mix 2 v2 N-2000246 uman B Cell Mix 2 v2 N-2000255	Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer:	PCR Primer	5'-GGCACCTTGTCCAATCATGTTCC-3' 5'-ATGTCGTTCATACTCGTCCTTGGT-3' Reverse Inner Primers: 5'-AGTCTCTCAGCTGGTACAC6-3' 5'-GGGAAGTTTCTGGCGGTACAC6-3' S'-GGGAAGTTTCTGGCGGTCA-3' 5'-GGGGAAGCTTTCTGAGCGGTCA-3' 5'-GGGGACCCAGTTATCAAGCAT-3' 5'-GGTGTCCCAGGTCACC-3' 5'-CCTGAGGACTGTAGGACAGC-3' 5'-CCTGAGGACTGTAGGACAGC-3' 5'-CCGTGTCCCGA-3' 5'-CCGTGTCCCGC-3' 5'-GCGTTATCCACCTTCCACTGT-3' Reverse Inner Primers:	_
uman T Cell Mix 2 v2 N-2000246 uman B Cell Mix 2 v2 N-2000255	Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC	PCR Primer	5'-GGCACCTTGTCCAATCATGTTCC-3' 5'-ATGTCGTTCATACTCGTCCTTGGT-3' Reverse Inner Primers: 5'-AGTCTCTCAGCTGGTACACG-3' 5'-TCTGATGGCTCAAACACAGC-3' Reverse Inner Primers: 5'-GGGAAGTTTCTGGCGGTCA-3' 5'-GGTGGTACCCAGTTATCAAGCAT-3' 5'-GGTGGTCCCAGGTCACCATCAC-3' 5'-TAGCTGCTCGAGGACTGAGGACAGC-3' 5'-CACGCTGCTCGTATCCGA-3' 5'-TAGCTGCTGGCGCCG-3' 5'-GCGTTATCCACCTTCCACTGT-3'	Inner Primer
uman T Cell Mix 2 v2 N-2000246 uman B Cell Mix 2 v2 N-2000255 ouse T Cell Mix 2 v2 N-2000257	Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC	PCR Primer PCR Primer PCR Primer	5'-GGCACCTTGTCCAATCATGTTCC-3' 5'-ATGTCGTTCATACTCGTCCTTGGT-3' Reverse Inner Primers: 5'-AGTCTCTCAGCTGGTACACG-3' 5'-GGCAAGTTTCTGGCGGTCA-3' 5'-GGGAAGTTTCTGGCGGTCA-3' 5'-GGTGGTACCCATTATCAAGCAT-3' 5'-GGTGGTCCCAGTCACAC-3' 5'-GGTGGTCCCAGTCACAC-3' 5'-CCTGAGGGCCGCA3' 5'-CCTGAGGCCGCA3' 5'-CCTGAGGCCGCA3' 5'-CCTGAGGCCGCA3' 5'-CCGGTGTCCCAGTCTACCA3' 5'-CCGGTCACCTGTATCCGA-3' 5'-CGGTTATCCACCTTCCACTGT-3' Reverse Inner Primers: 5'-AGTCAAAGTCGGTGAACAGGCA-3'	Inner Primer
uman T Cell Mix 2 v2 N-2000246 uman B Cell Mix 2 v2 N-2000255 ouse T Cell Mix 2 v2 N-2000257	Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer:	PCR Primer PCR Primer PCR Primer PCR Primer PCR Primer	5'-GGCACCTTGTCCAATCATGTTCC-3' 5'-ATGTCGTTCATACTCGTCCTTGGT-3' S'-ATGTCGTTCATACTCGTCCTTGGT-3' Control Conte	Inner Primer
uman T Cell Mix 2 v2 N-2000246 uman B Cell Mix 2 v2 N-2000255 ouse T Cell Mix 2 v2 N-2000257	Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC	PCR Primer PCR Primer PCR Primer PCR Primer PCR Primer	5'-GGCACCTTGTCCAATCATGTTCC-3' 5'-ATGTCGTTCATACTCGTCCTTGGT-3' Reverse Inner Primers: 5'-AGTCTCTCAGCTGGTACAC6-3' 5'-GGCAAGTTTCTGGCGGTCA-3' 5'-GGGAAGTTTCTGGCGGTCA-3' 5'-GGGGAAGTTTCTGGCGGTCA-3' 5'-GGTGTCCCAGGTCACCATCAC-3' 5'-GGTGTCCCAGGTCACCATCAC-3' 5'-CCTGAGGCCCCACTATCCAGC-3' 5'-CCTGAGGCCCGC-3' 5'-CCGGTTATCCACCTTCCACTGT-3' Reverse Inner Primers: 5'-AGTCAAAGTCGGTGAACAGGCA-3' 5'-GCCAAGCTCGCGTGAACAGGCA-3' 5'-GCCAAGCCGCGCA3' 5'-GCCAAGCCACCTCCACTGT-3' Reverse Inner Primers: 5'-AGTCAAAGTCGGTGAACAGGCA-3' 5'-GCCAAGCCACCAGGGGTA-3' S'-GCCAAGCCACCACGAGGGTA-3'	Inner Primer
Automated Protocol S uman T Cell Mix 2 v2 N-2000246 uman B Cell Mix 2 v2 N-2000255 louse T Cell Mix 2 v2 N-2000257 Aouse B Cell Mix 2 v2 N-2000259	Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer:	PCR Primer PCR Primer PCR Primer PCR Primer PCR Primer	5'-GGCACCTTGTCCAATCATGTTCC-3' 5'-ATGTCGTTCATACTCGTCCTTGGT-3' S'-ATGTCGTTCATACTCGTCCTTGGT-3' S'-AGTCTCTCAGCTGGTACACG-3' S'-AGTCTCTCAGCTGGTACACG-3' S'-GGCAAGTTTCTGCGCGTACACG-3' S'-GGGAAGTTTCTGCGCGCA-3' S'-GGGAAGTTTCTGCGCGCA-3' S'-GGGGAACCAGTTATCAAGCAT-3' S'-GGTGTCCCAGGTCACCATCAC-3' S'-GGTGTCCCAGGTCACCATCAC-3' S'-TCCTCAGGACCTGTAGGACCAGC-3' S'-AGCTGCTGGCCGC-3' S'-AGCCCAAGTCACCACGAGGCA-3' S'-AGCCAAGTCGGTGAACAGGCA-3' S'-AGCCAAGTCGTGGGCCTT-3' S'-GGCCAAGCCACGTGTGGCCTT-3' S'-CAGGCCACTGTCACCACCT-3' S'-CAGGCCACATCACCAGTGTGGCCTT-3'	Inner Primer
uman T Cell Mix 2 v2 N-2000246 uman B Cell Mix 2 v2 N-2000255 ouse T Cell Mix 2 v2 N-2000257	Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer:	PCR Primer PCR Primer PCR Primer PCR Primer PCR Primer	5'-GGCACCTTGTCCAATCATGTTCC-3' 5'-ATGTCGTTCATACTCGTCCTTGGT-3' S'-ATGTCGTTCATACTCGTCCTTGGT-3' S'-AGTCTCTCAGCTGGTACACG-3' S'-AGTCTCTCAGCTGGTACACG-3' S'-GGCACTTTCTGCGCGTCA-3' S'-GGGGAGTTTCTGCGCGTCA-3' S'-GGTGGTACCCAGTTATCAAGCAT-3' S'-GGTGGTACCCAGTTATCAAGCAT-3' S'-GGTGGTACCCAGTTATCCAGCAC-3' S'-TCCTGAGGACTGTCGCACCATCAC-3' S'-TCCTGAGGACTGTCGGACAGC-3' S'-AGTCAAAGTCGTGACACGC-3' S'-AGTCAAAGTCGGTGAACAGGCA-3' S'-GGCCAAGCCACCGGGGTA-3' S'-AGTCAAAGTCGTGACACGGGCA-3' S'-GGCCAAGCCACGAGGGTA-3' S'-GGCCAAGCACACGAGGGGTA-3' S'-GGCCAAGCCACGAGGGGAA-3' S'-TACACACCAGTGTGGCCTT-3' S'-CAGGCCACTGTCACACCACT-3' S'-CAGGCCACTGTCACACCACT-3'	Inner Primer
uman T Cell Mix 2 v2 N-2000246 uman B Cell Mix 2 v2 N-2000255 ouse T Cell Mix 2 v2 N-2000257	Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer:	PCR Primer PCR Primer PCR Primer PCR Primer PCR Primer	5'-GGCACCTTGTCCAATCATGTTCC-3' 5'-ATGTCGTTCATACTCGTCCTTGGT-3' S'-ATGTCGTTCATACTCGTCCTTGGT-3' S'-AGTCTCTCAGCTGGTACACG-3' 5'-AGTCTCTCAGCTGGTACACG-3' S'-GGGAAGTTTCTGGCGGTCA-3' S'-GGGGAAGTTTCTGGCGGTCA-3' S'-GGTGTCCCAGTGTAGCACAGC-3' S'-GGTGTCCCAGTCACAC-3' S'-TCCTGAGGACTGTAGGACAGC-3' S'-TCCTGAGGACTGTAGGACAGC-3' S'-TCCTGAGGACTGTAGGACAGC-3' S'-TCCTGAGGACTGTCGCAGCA-3' S'-TACCTGCTGCCCGC-3' S'-AGTCAAAGTCGGTGAACAGGCA-3' S'-AGCCAAGCCACTGCGGCTA-3' S'-AGCCAAGTGACCACCACGAGGCA-3' S'-AGCCAAGTGCACCACTGAGGCA-3' S'-AGCCAAGTGCACCACTGGGCTA-3' S'-AGCCAAGTGCACCACT-3' S'-AGGCCACGTCTCACCGTCCC-3' S'-AGGCCACGTCACAGTGACCT-3' S'-CAGGGCAAGTCACAGTGCCC-3' S'-CAGGGAAGTTCACAGTGCC-3' S'-CGCAGGGAAGTTCACAGTGCC-3' S'-CGCAGGGAAGTTCACAGTGCC-3' S'-CGCAGGCAAGTCACAGTGCCT-3' S'-CGCAGGCAAGTCACAGTGCCT-3' S'-CGCAGGCAAGTCACAGTGCCT-3' S'-CGCAGGCAAGTCACAGTGCCT-3' S'-CGCAGGCAAGTCACAGTTGCCT-3' S'-CGCAGGCAAGTCACAGTACTTG-3'	Inner Primer
uman T Cell Mix 2 v2 N-2000246 uman B Cell Mix 2 v2 N-2000255 ouse T Cell Mix 2 v2 N-2000257	Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer:	PCR Primer PCR Primer PCR Primer PCR Primer PCR Primer	5'-GGCACCTTGTCCAATCATGTTCC-3' 5'-ATGTCGTTCATACTCGTCCTTGGT-3' S'-ATGTCGTTCATACTCGTCCTTGGT-3' Reverse Inner Primers: 5'-GGCAAGTTTCTGGCGGTCA-3' 5'-GGGAAGTTTCTGGCGGTCA-3' 5'-GGTGTCCCAGGTCACCA-3' 5'-GGTGTCCCAGGTCACCA-3' 5'-GGTGTCCCAGGTCACCA-3' 5'-GTGTCCCAGGTCACCAC-3' 5'-GCGTACCCAGTATCCAGA-3' 5'-CCTGAGGACTGTAGGACAGC-3' 5'-GCGTTATCCACCTTCCACTGT-3' S'-GCGTTATCCACCTTCCACTGT-3' S'-GCGTTATCCACCTTCCACTGT-3' S'-GCGTATCCACCTGCGCA-3' 5'-GCCAAGCCACGGGGAACTCACGAGGCA-3' 5'-GCCAAGCACAGTGGGCTT-3' S'-GCCAGGCACGTGTCACACACT-3' 5'-CAGGCCACGTGTCACACACT-3' 5'-GCGAGGCAACTTCACGTTCCCACT-3' 5'-GCGAGGCAACTTCACGTTGCCA-3' 5'-GCGGGAACTTCACAGTGCCC-3' 5'-GCGAGGCACGTGACCT-3' 5'-GCGAGGTGGCTACGTACCT-3' 5'-GCGAGGTGGCTACGTTCCCACTCC-3' 5'-GCGAGGTGGCTAGGTACTTG-3' 5'-GCGAGGTGGCTAGGTACTTG-3' 5'-CCTTGACCAGGCAACGTACCTG-3'	Inner Primer
uman T Cell Mix 2 v2 N-2000246 uman B Cell Mix 2 v2 N-2000255 ouse T Cell Mix 2 v2 N-2000257	Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer: 5'-GATCTACACTCTTTCCCTACACGAC Forward Primer:	PCR Primer PCR Primer PCR Primer PCR Primer PCR Primer	5'-GGCACCTTGTCCAATCATGTTCC-3' 5'-ATGTCGTTCATACTCGTCCTTGGT-3' S'-ATGTCGTTCATACTCGTCCTTGGT-3' S'-AGTCTCTCAGCTGGTACACG-3' 5'-AGTCTCTCAGCTGGTACACG-3' S'-GGGAAGTTTCTGGCGGTCA-3' S'-GGGGAAGTTTCTGGCGGTCA-3' S'-GGTGTCCCAGTGTAGCACAGC-3' S'-GGTGTCCCAGTCACAC-3' S'-TCCTGAGGACTGTAGGACAGC-3' S'-TCCTGAGGACTGTAGGACAGC-3' S'-TCCTGAGGACTGTAGGACAGC-3' S'-TCCTGAGGACTGTCGCAGCA-3' S'-TACCTGCTGCCCGC-3' S'-AGTCAAAGTCGGTGAACAGGCA-3' S'-AGCCAAGCCACTGCGGCTA-3' S'-AGCCAAGTGACCACCACGAGGCA-3' S'-AGCCAAGTGCACCACTGAGGCA-3' S'-AGCCAAGTGCACCACTGGGCTA-3' S'-AGCCAAGTGCACCACT-3' S'-AGGCCACGTCTCACCGTCCC-3' S'-AGGCCACGTCACAGTGACCT-3' S'-CAGGGCAAGTCACAGTGCCC-3' S'-CAGGGAAGTTCACAGTGCC-3' S'-CGCAGGGAAGTTCACAGTGCC-3' S'-CGCAGGGAAGTTCACAGTGCC-3' S'-CGCAGGCAAGTCACAGTGCCT-3' S'-CGCAGGCAAGTCACAGTGCCT-3' S'-CGCAGGCAAGTCACAGTGCCT-3' S'-CGCAGGCAAGTCACAGTGCCT-3' S'-CGCAGGCAAGTCACAGTTGCCT-3' S'-CGCAGGCAAGTCACAGTACTTG-3'	Inner Primer

	CTACACGACGCTCTTCCGATCT-N16-N10-TT 3GATGTGCTGCGAGAAGGCTAGA-N16-N10-A		ner-3'	
Automated Proto	ocol Step – Adaptor Ligation (for V(D)J Library Construct	ion)	
Adaptor (Read 2)	Read 5'-GATCGGAAGAGCACACGTCTGA 3'-TCTAGCCTTCTCG-5'	2 ACTCCAGTCAC-3'		
	CCTACACGACGCTCTTCCGATCT-N16 <mark>-N10-</mark> GGATGTGCTGCGAGAAGGCTAGA-N16- <mark>N10</mark> -/			C-3'
Automated Prote	ocol Step – Sample Index PCR	R (for V(D)J Library Construc	ction)	
Dual Indexing Dual Index Kit TT Set A PN-1000215		Sample Partial Read 1 Index (i5) N10-ACACTCTTTCCCTACACGACGCTC-3'	Reverse Primer: 5'-CAAGCAGAAGACGGCAT	P7 Sample Partial Read 2 Index (i7) ACGAGAT-N10-GTGACTGGAGTTCAGACGTGT-3'
Sample Index PC	P5	Sample Read 1 10x UMI TSO Index (i5) Barcode	V D J C Read 2 Sampl Index	

3'-TTACTATGCCGCTGGTGGCTCTAGATGTG-N10-TGTGAGAAAGGGATGTGCTGCGAGAAGGCTAGA-N16-N10-AAAGAATATACCC-Insert-TCTAGCCTTCTCGTGTGCAGACTTGAGGTCAGTG-N10-TAGAGCATACGGCAGAAGACGAAC-5'

Automated Proto	ocol Step – GEX Adaptor Ligation	n (for 5' Gene Expressio	on (GEX) L	ibrary Construc	tion)
Adaptor Read 2	Read 2			5′ –gatcggaaga 3′ –tctagccttc:	gCacacgtctgaactccagtcac-3′ rcg-5′
Ligation Product		ead 1 10x UMI TSO Barcode	Insert	Read 2	
	GATCT-N16-N10-TTTCTTATATGGG-cDNA _ Ins CCTAGA-N16-N10-AAAGAATATACCC-cDNA _ Ins		GAACTCCAGTCA	LC-3'	
Automated Proto	ocol Step – Sample Index PCR (fo	or 5' Gene Expression (GEX) Libr	ary Constructior	1)
Dual Indexing F	Forward Primer:			Reverse Primer:	
Dual Index Kit TT Set A PN-1000215 5	"-AATGATACGGCGACCACCGAGATCT-N10-ACA(CTCTTTCCCTACACGACGCTC-3		5'-CAAGCAGAAGACGGG	CATACGAGAT-N10-GTGACTGGAGTTCAGACGTGT-3'
Sample Index PCR Product	P5 Sample Index (is	e Read 1 10x UMI TSO Barcode	Insert	Read 2 Sample Index (i7)	27
	ACGGCGACCACCGAGATCTACAC-N10-ACACTCTTTCCCTACACG		_		
Automated Proto	ocol Step - Sample Index (for Ce	ll Surface Protein/Imm	nune Rece	ptor Mapping Lil	orary Construction)
Dual Index Kit	Forward Primer: P5 Sample F Index (i5)	Partial Read 1		Reverse Primer:	P7 Sample Partial Read 2N Index (i7)
TN Set A PN-1000250 5	5'-AATGATACGGCGACCACCGAGATCTACAC-N10	-ACACTCTTTCCCTACACGACGCT	C-3'	5'-CAAGCAGAAGACGG	CATACGAGAT-N10-GTCTCGTGGGCTCGG-3'
Sample Index PCR Product	P5 Sample Index (i5)	Read 1 10x UMI Capture Barcode Seq	Feature Re Barcode	ad 2N Sample P7 Index (i7)	-