

#### **Notices**

#### Limitations of use

For research use only. Not for use in diagnostic procedures. Unless otherwise agreed to in writing, IDT does not intend these products to be used in clinical applications and does not warrant their fitness or suitability for any clinical diagnostic use. Purchaser is solely responsible for all decisions regarding the use of these products and any associated regulatory or legal obligations.

Safety data sheets pertaining to this product are available upon request.

#### **Safety Notices**



**Reminder** symbols call attention to minor details that may be easily overlooked and compromise the procedure resulting in decreased assay performance.



**Caution** symbols denote critical steps in the procedure where risk of protocol failure or damage to the product itself could occur if not carefully observed.



**Stop** symbols indicate where this procedure may be safely suspended and resumed at a later time without risk of compromised assay performance. Make note of these steps and plan your workflow accordingly.



### FUSION*Plex*™ Protocol for Illumina®

### **Contents**

Overview	5
Before getting started	10
Protocol	16

# **Revision history**

Document Number	Date	Description of change
LA135.1	2020	<ul> <li>Updated table formatting.</li> <li>Removal of recommendation to run fragment analysis on final libraries.</li> <li>Updated PCR cycling tables to refer to the product insert for panel-specific cycling recommendations.</li> <li>Updated vendor part numbers for materials required but not supplied.</li> </ul>
PR0028.1	August 2021	<ul> <li>Changed number of document from LA135.1 to PRO028.1</li> <li>No document content changes were made.</li> </ul>
PR0028.2	November 2021	<ul><li>Removal of Quantify, Normalize, and Sequence Section.</li><li>Updated links and email addresses.</li></ul>
PR0028.3	July 2021	<ul> <li>Migrated protocol to new format.</li> <li>Removed liquid accessories.</li> <li>Updated contact information.</li> <li>Updated figures.</li> </ul>
PR0028.4	November 2022	<ul> <li>Updated formatting.</li> <li>Added PreSeq review advice.</li> <li>Added "Data Acquisition" column to PreSeq RNA QC Assay thermal cycler program.</li> <li>Aligned minor inconsistencies across protocols.</li> </ul>
PR0028.5	March 2023	Updated formatting and branding.
RA-DOC-047/REV01	June 2023	<ul> <li>Updated document number from PRO028 to RA-DOC-047.</li> <li>Updated formatting and branding.</li> </ul>



### FUSION*Plex*™ Protocol for Illumina®

RA-DOC-047/REV02	November 2023	<ul> <li>Updated First and Second PCR cycling conditions to include separate anneal and extended steps.</li> <li>Edited reaction mix table under "Step 4: PreSeq RNA QC Assay" for clarity.</li> <li>Clarified instructions under the "Working with lyophilized reagents" section.</li> </ul>
		Updated formatting and branding.
		<ul> <li>Updated links and email addresses.</li> </ul>



#### **Technical support**

Contact us directly at archer-tech@idtdna.com.



#### **Overview**

#### Intended use

The Archer FUSION *Plex* protocol is intended for research applications in conjunction with Archer reagent kits and corresponding target-enrichment panels to produce high-complexity libraries for use with Illumina next- generation sequencing (NGS) platforms.

FUSION*Plex* sequencing data produced by this method should be processed using Archer™ Analysis software—a complete bioinformatics suite that leverages Anchored Multiplex PCR (AMP™) chemistry to identify unique sequence fragments, thus enabling error correction, read deduplication, and ultimately high-confidence alignment and mutation calling. Archer Analysis takes demultiplexed FASTQ files straight from the sequencer as input and produces both high-level and detailed mutation reporting, as well as raw text and BAM outputs for full transparency of the pipeline.

#### **Test principle**

Anchored Multiplex PCR (AMP) is a rapid and scalable method to generate target-enriched libraries for NGS. AMP technology can be used for applications in targeted RNA sequencing, genomic DNA sequencing, and genotyping applications to generate a sequencing library in a matter of hours. Designed for low nucleic acid input, this process delivers robust performance across a variety of sample types.

AMP utilizes unidirectional gene-specific primers (GSPs) that enrich for both known and unknown mutations. Adapters that contain both molecular barcodes and sample indices permit quantitative multiplex data analysis, read deduplication, and accurate mutation calling.

The Archer Analysis software utilizes these molecular barcodes (MBCs) for duplicate read binning, error correction and read deduplication to support quantitative multiplex data analysis and confident mutation detection. Analysis reports both sequencing metrics and number of unique observations supporting called variants.

#### Modular assay format Archer library preparation reagents include:

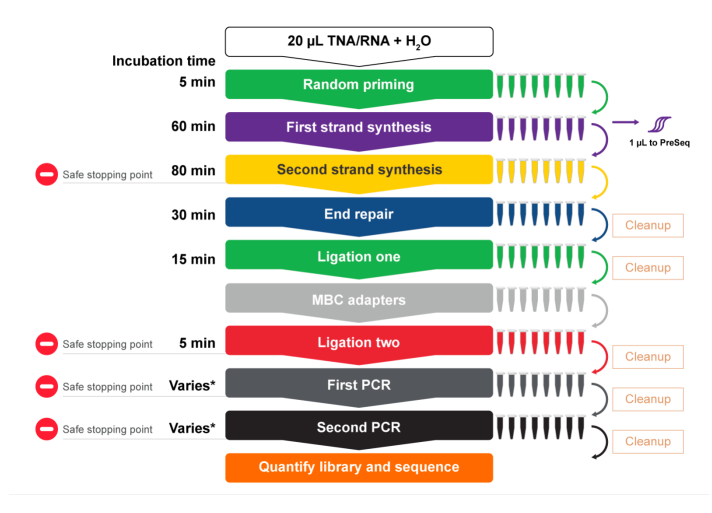
- Archer FUSION*Plex* reagents in lyophilized format for each step of library preparation.
- GSPs that target panel-specific regions of interest during PCR amplification.
- Archer MBC adapters are proprietary adapters that tag each unique molecule with a barcode and common region prior to amplification.
- Archer PreSeq™ RNA QC assay to determine the quality of starting material.

#### **Modular assay format**

Archer kits include library preparation reagents and assay-specific liquid primers, which are used in conjunction with Archer MBC adapters to construct sequencing-ready libraries from total nucleic acid (TNA) or RNA samples. See individual product inserts for panel targets and read depth requirements.



#### Workflow



<sup>\*</sup> First and Second PCR times vary based on the specific panel as well as individual lab cycling conditions. See Product Insert for more information.



#### FUSION Plex Reagents for Illumina (SK0093)

Store at 2°C to 8°C - Allow pouches to reach room temperature before opening.

Materials Supplied					
Description Part Number Quantity					
Random Priming 2.0	SA0194				
First Strand cDNA Synthesis	SA0002				
Second Strand cDNA Synthesis	SA0003				
End Repair	SA0204	1 pouch			
Ligation Step 1	SA0196	(8 reactions/one 8-tube strip)			
Ligation Step 2	SA0197				
First PCR (Illumina-P)	SA0109				
Second PCR Reactions 1 thru 8 (Illumina- P)	SA0110				
Ligation Cleanup Beads	SA0210	1 tube			
Ligation Cleanup Buffer	SA0209	(sufficient for processing 8 samples)			

#### **Additional components**

#### Store at -30°C to -10°C

	Materials Supplied	
Description	Part Number	Quantity
FUSIONPlex Panel GSP1	Refer to product insert	O reactions
FUSIONPlex Panel GSP2	Refer to product insert	8 reactions
10X VCP Primer Mix	SA0126	16 reactions*

<sup>\*</sup>Supplied volume is sufficient to run duplicate qPCR reactions for 16 samples and 4 No Template Controls (NTC).



### FUSION*Plex*™ Protocol for Illumina®



Materials required, but not supplied have been optimized for proper application of the Archer Library preparation. Use of other materials have not been tested by IDT.

Materials Required, but not Supplied				
Description	Supplier	Part Number		
Archer MBC adapters for Illumina	Archer	Varies		
AMPure® XP beads	Beckman Coulter	A63880		
iTaq™ Universal SYBR® Green Supermix	Bio-Rad Laboratories	172-5120		
1 M Tris-HCl, pH 8.0 (molecular biology grade)	Various	-		
Ultrapure water (molecular biology grade)	Various	-		
200 mM Tris-HCl, pH 7.0 (for sequencing)	Various	-		
100% ethanol (ACS grade)	Various	-		
Concentrated NaOH solution (ACS grade)	Various	-		
RNase AWAY™	Thermo Fisher Scientific	7003		
KAPA Universal Library Quantification Kit	KAPA Biosystems	KK4824		
MiSeq® or NextSeq® Reagent Kit (300 cycle minimum)	Illumina	-		
PhiX Control v3	Illumina	FC-110-3001		
Standard PCR thermal cycler	Various	-		
Real-Time PCR thermal cycler	Various	-		
qPCR tubes	Various	-		
0.2 mL PCR tubes	Various	-		
DynaMag™-96 Side Magnet	Thermo Fisher Scientific	12331D		
Microcentrifuge	Various	-		



### FUSION*Plex*™ Protocol for Illumina®

Plate centrifuge	Various	-
Pipettes (P10, P20, P200 and P1000)	Pipetman or equivalent	-
Sterile, nuclease-free aerosol barrier pipette tips	Various	-
Vortex mixer	Various	-
PCR tube cooling block	Various	-
Gloves	Various	-
Qubit® Fluorometer (3.0 or higher)	Thermo Fisher Scientific	Q33216
Qubit RNA HS Assay Kit	Thermo Fisher Scientific	Q32852



## **Before getting started**

#### **Important precautions**

- Read through the entire protocol before starting your library preparation.
- Take note of safe stopping points throughout the protocol where samples can be safely frozen (-30° C to -10° C) to plan your workflow.
- Use good laboratory practices to prevent contamination of samples by PCR products.
- Use nuclease-free PCR tubes, microcentrifuge tubes, and aerosol-barrier pipette tips.
- Wipe down workstation and pipettes with nuclease and nucleic acid cleaning products (e.g., RNase AWAY, Thermo Fisher Scientific).
- Verify that the thermal cycler used for library preparation is in good working order and currently calibrated according to manufacturer specifications. When using a thermal cycler with a 100% ramp rate >6°C/sec, set the ramp rate to go no higher than 6°C/sec for First PCR and Second PCR.
- Reaction cleanup with AMPure XP beads (Beckman Coulter) is performed at room temperature (20°C to 25°C) and is used repeatedly throughout the workflow. Ensure that AMPure XP beads are equilibrated to room temperature and fully resuspended by vortexing until homogenous in both color and appearance prior to drawing out material for **each** use.

#### **Working with lyophilized reagents**

- Archer reagents are provided as individually lyophilized reaction pellets in 0.2 mL PCR tube strips.
- Allow pouches to reach room temperature (20°C to 25°C) by removing from 2°C to 8°C storage for at least 15 minutes before opening to reduce moisture condensation on tubes.
- Always centrifuge tubes briefly before opening to pull contents down.
- Detach the required number of reaction tubes and return any unused portion to the pouch with desiccant packet, reseal and store at 2°C to 8°C. It is recommended to use the remaining reactions within 4 weeks after opening.
  - For MBC adapters and Second PCR tubes remember to label prior to returning unused portions to storage.
- Dissolve, mix and spin down:
  - Never touch the lyosphere with the pipette tip.
  - Add sample/reagents to pellets in tubes while on ice.



- Allow at least 5 seconds for pellets to dissolve.
- Pipette up and down 8 times to mix after the lyosphere has dissolved.
- Briefly centrifuge and return to ice before proceeding.

#### Input nucleic acid

- Input nucleic acid (TNA or RNA) in EDTA-free buffer (pH 7-8) or ultrapure water is the optimal starting template for AMP library preparation. Do NOT use EDTA-containing buffers.
  - Note: Some FUSIONPlex panels contain gene-specific primers designed for sample tracking that target non-expressed DNA sequence (intronic or intergenic). When using SEX\_ID and SNP\_ID-based sample tracking, TNA should be used as the starting template.
- Use the maximum allowable input mass (ng) whenever possible. Higher input quantities enable more sensitive variant and fusion detection:
  - 20 250 ng of RNA for FUSION Plex
- If using total nucleic acid (TNA), Do not pretreat with DNase. DNA found in total nucleic acid can act as an internal control, verifying assay performance in the absence of RNA.
- Contact technical support (archer-tech@idtdna.com) for commercially available extraction kit recommendations.

#### Reagents to prepare before starting

- Make at least 10 mL fresh 10 mM Tris-HCl, pH 8.0 from 1M Tris-HCl, pH 8.0 and ultrapure water.
  - o Mix 100 μL 1M Tris-HCl, pH 8.0 with 9,900 μL ultrapure water.
  - o 10 mM Tris-HCl, pH 8.0 can be used for up to one week after mixing.
- Make at least 50 mL fresh 70% ethanol from 100% ethanol and ultrapure water.
  - Add 35 mL 100% ethanol to 15 mL ultrapure water.
  - o Tightly close the cap to minimize evaporation when not in use.
  - o 70% ethanol is appropriate for use for up to one week after mixing.
- Make at least 1 mL fresh 5 mM NaOH from concentrated NaOH and ultrapure water on the day of use.
  - If working from 1M NaOH, add 5 μL of 1M NaOH to 995 μL of ultrapure water to yield 5 mM final NaOH.



#### FUSION*Plex*™ Protocol for Illumina®

o If working from 5M NaOH, add 10 μL of 5M NaOH to 990 μL of ultrapure water to yield 50 mM NaOH. Mix well and briefly spin down. Take 100 μL of 50 mM NaOH and combine with 900 μL of ultrapure water to yield 5 mM NaOH. Mix well and briefly spin down.

### Thermal cycler programs

- Preprogram your thermal cycler with these specific programs.
- Use the appropriate programs for the specific Archer assay.
- Verify programming before initiating runs.

	Step	Temperature (°C)	Time (min)
Dondon Brinsing 2.0	1	65	5
Random Priming 2.0	2	4	Hold

	Step	Temperature (°C)	Time (min)
	1	25	10
First Strand cDNA	2	42	30
Synthesis	3	80	20
	4	4	Hold

	Step	Temperature (°C)	Time (min)
	1	16	60
Second Strand cDNA Synthesis	2	75	20
,	3	4	Hold



### FUSION*Plex*™ Protocol for Illumina®

	Step	Temperature (°C)	Time (sec)	Data Acquisition	Cycles
	Activation	95	20 [20*]	Off	1
PreSeq RNA	Denaturation	95	15 [3*]	Off	
QC Assay (on Real-Time PCR instrument)	Primer annealing & extension	60	60 [30*]	On	35
	Melt-curve gradient	60-95	0.5°C/sec increment	On	1

<sup>\*</sup>Times in [] are for fast-mode cycling. Only use fast-mode conditions if specifically supported by the qPCR instrument.

	Step	Temperature (°C)	Time (min)
Complete End Repair	1	25	30
(Heated lid off)	2	4	Hold

	Step	Temperature (°C)	Time (min)
Ligation Stan 4	1	37	15
Ligation Step 1	2	4	Hold

	Step	Temperature (°C)	Time (min)
Ligation Step 2	1	22	5
Ligation Step 2 (Heated lid off)	2	4	Hold

	Step	Temperature (°C)	Time (min)
Ligation Elution	1	75	10
Ligation Elution	2	4	Hold



#### FUSION*Plex*™ Protocol for Illumina®

	Step	Temperature (°C)	Time	Cycles
	1	95	3 min	1
	2	95	30 sec	
	3		10 sec	Varies-See panel specific product
First PCR	4	Varies-See panel specific product insert)	Varies-See panel specific product insert) 100% ramp rate*	insert)
	5	72	3 min	1
	6	4	Hold	1

<sup>\*</sup>See ramp rate guidelines in the "Before getting started" section.

	Step	Temperature (°C)	Time	Cycles
	1	95	3 min	1
	2	95	30 sec	
	3	Varies-See panel specific product insert)	10 sec	Varies-See panel specific product
Second PCR	4		Varies—See panel specific product insert) 100% ramp rate*	insert)
	5		3 min	1
	6	4	Hold	1

\*See ramp rate guidelines in the "Before getting started" section.

#### Molecular barcoding, sample indexing, and multiplexed sequencing

Molecule-level barcoding (or unique molecule identifier tagging) and sample-level barcoding (also known as index tagging) are both incorporated during Archer MBC ligation. Molecular barcodes are an integral component of the Archer Analysis software suite. Sample barcodes (i.e., index tags) allow pooled libraries to be sequenced simultaneously thereby enabling maximum sequencing throughput and data demultiplexing during downstream bioinformatics analysis.



#### Sample multiplexing

- To efficiently utilize the throughput of the MiSeq (or other Illumina sequencing platform) as well as
  prevent low index diversity within your sequencing run, multiple samples should be sequenced
  simultaneously. Samples can be identified through a combination of two unique nucleotide
  sequences (see below for more details), which are subsequently read during the sequencing
  process. The unique nucleotide sequence is often termed an "index".
- The Archer Library Preparation Reagents for Illumina use a combination of two indices to distinguish between samples. Index 2 is added during Ligation Step 2 and is embedded in the Archer MBC adapters for Illumina (p5/i5 index). Index 1 is added during the Second PCR and is embedded in Illumina Index 1 Primers (p7/i7) within the Second PCR reaction pellets.
- To maintain appropriate coverage depth, it is recommended that users determine the maximum number of samples that can be run on a flow cell. In general, larger panels with more targets will require higher sequencing coverage depth and should be run with fewer samples. Refer to product insert for panel specific read depth recommendations.
- Contact <u>archer-tech@idtdna.com</u> to receive tools that can help define the appropriate number of samples per sequencer and flow cell type.

#### **Barcode diversity**

- Illumina sequencers work best when index diversity within a run is high. For example, if eight
  samples are included in a run, and the user chooses to use only one MBC adapter paired with
  eight different Index 1 Primers, the run may fail due to low barcode diversity. In this example it is
  recommended to use eight different Archer MBC adapters paired with eight different Index 1
  Primers.
- To facilitate a high index diversity, each tube in the 8-strip of MBC adapters for Illumina as well as Second PCR contains a unique index. When preparing one batch of 8 or less samples, this results in 8 samples with unique dual indices and barcode diversity is not a concern. Only when combining very few samples of different batches, it becomes necessary to pay attention to the index diversity, as the i7 indices (added with Second PCR) are the same in each kit.
- If using more than 48 MBCs, contact archer-tech@idtdna.com for adapter compatibility.

#### **Protocol**

Before beginning, review precautions in "Working with Lyophilized Reagents" section for additional information regarding best practices using lyophilized reagents.

### **Step 1: Random Priming 2.0**

- 1. Place an appropriate number of **Random Priming 2.0** (SA0194) reaction tubes on ice.
- 2. Adjust purified RNA/TNA (20 250 ng) samples to a final volume of 20  $\mu$ L and transfer to new PCR tubes. Keep input samples on ice.

Component	Reaction Mix
Ultrapure water	20 - XµI
Purified nucleic acid or RNA	ΧμΙ
Total volume	20µl

- 3. Transfer **20 µL** reaction mix to the Random Priming 2.0 reaction tubes.
  - a. Dissolve, mix and spin down.
  - b. Return tubes to ice.
- 4. Start the following thermal cycler program, and only transfer reactions to the block once temperature reaches 65°C. Pause the program if necessary.
  - a. Use a heated lid (≥100°C).

Random Priming 2.0 Incubation conditions

Step	Temperature (°C)	Time (minutes)
1	65	5
2	4	Hold

b. After the program has reached 4°C, briefly spin down reactions and place tubes on ice for at least 2 minutes.

## **Step 2: First Strand cDNA Synthesis**

1. Place an appropriate number of **First Strand cDNA Synthesis** (SA0002) reaction tubes on ice.





- Spin down the Random Priming 2.0 mixture and transfer 20 μL to the First Strand cDNA Synthesis tubes.
  - a. Dissolve, mix and spin down.
  - b. Return tubes to ice.
- Transfer reactions to a preheated thermal cycler and initiate an incubation using the following program and guidelines:
  - a. Use a heated lid (≥100°C).

First Strand cDNA Synthesis Incubation conditions

Step	Temperature (°C)	Time (minutes)
1	25	10
2	42	30
3	80	20
4	4	Hold

- b. After the program has reached 4°C, briefly spin down reactions and place on ice.
- 4. Make diluted cDNA samples for the PreSeq RNA QC Assay.
  - a. Place a new, empty 8-strip of PCR tubes on ice.
  - b. For each sample, pipette **9 µL** of ultrapure water into the new PCR strip.
  - c. Pipette **1**  $\mu$ L of each First Strand cDNA Synthesis reaction into the water and pipette up and down to mix.
  - d. Keep on ice for use in Step 4: PreSeg RNA QC Assay

### **Step 3: Second Strand cDNA Synthesis**

- Place an appropriate number of Second Strand cDNA Synthesis (SA0003) reaction tubes on ice.
- 2. Add **21 μL** of ultrapure water to each tube containing the **19 μL** of First Strand cDNA Synthesis reaction.
  - a. Pipette up and down to mix.
- 3. Pipette **40 µL** of each diluted First Strand reaction into the new Second Strand cDNA Synthesis tubes.



- a. Dissolve, mix and spin down.
- b. Return tubes to ice.
- 4. Start the following thermal cycler program, and only transfer reactions to the block once temperature reaches 16°C. Pause the program if necessary.
  - a. Use a heated lid (≥100°C).



Secono	Second Strand cDNA Synthesis Incubation conditions		
Step	Temperature (°C)	Time (minutes)	
1	16	60	
2	75	20	
3	4	Hold	

- b. Place samples in the thermal cycler and start the program.
- c. While sample(s) are incubating, proceed to Step 4: PreSeq RNA QC Assay.
- d. After the run has completed, briefly spin down reactions and place on ice.



**Safe stopping point:** It is okay to stop and store the reactions at -30°C to -10°C. It is recommended to review the qPCR results from the PreSeq RNA QC assay at this time to determine predicted sample success.

### Step 4: PreSeq RNA QC Assay

- 1. Thaw the **10X VCP Primer Mix** (SA0126) at room temperature. Once thawed, store 10X VCP Primer Mix on ice.
- 2. Place a fresh qPCR plate or tubes appropriate to your Real-Time PCR instrument on ice.



#### FUSION*Plex*™ Protocol for Illumina®

- 3. Prepare sufficient qPCR reaction mix for:
  - a. Duplicate reactions of each diluted cDNA sample from Step 2: First Strand cDNA Synthesis.

b. One No Template Control (NTC) made using ultrapure water.

Component	Part Number	Reaction Mix (n = 1)
iTaq SYBR Green Supermix	Not Supplied	5 μL
10X VCP Primer Mix	Varies	1 μL
Total reaction mix volume	-	6 μL

- 4. Pipette 6 μL of the reaction mix into each assigned well of a qPCR plate/tube.
- **5.** Pipette **4 μL** of the diluted cDNA samples or NTC (ultrapure water) into the assigned wells/tubes containing reaction mix for a total 0f **10 μL**.
  - a. Mix slowly to avoid introducing bubbles, cap or seal the reactions and spin down.
- 6. Transfer reactions to a Real Time-PCR instrument and initiate a run using the following program:

PreSeg RNA QC Assay Real-Time incubation conditions

Step	Temperature (°C)	Time (sec)	Data Acquisition	Cycles
Activation	95	20 [20*]	Off	1
Denaturation	95	15 [3*]	Off	
Primer annealing & extension	60	60 [30*]	On	35
Melt-curve gradient	60-95	0.5°C/sec increment	On	1

<sup>\*</sup>Times in [] are for fast-mode cycling. Only use fast-mode conditions if specifically supported by the qPCR instrument.

- 7. After completion of the program, review the Ct values of each sample.
  - a. A low Ct value generally correlates with a high probability for good QC values when analyzing the samples in Archer Analysis.
  - b. A high Ct value or N/A result indicates issues with the sample quality or quantity.
  - c. A lab should establish their own Ct-value threshold for when the prospects of success for individual samples are too low to process further in the library prep. It is possible to purchase a set of the first 3 steps of the FUSION Plex library prep individually (SK0107;



PreSeq RNA QC Assay Reagents - 16 reactions), to allow for affordable repetition of difficult samples, without wasting a whole reaction.

### Step 5: End Repair

- 1. Place an appropriate number of **End Repair** (SA0204) reaction tubes on ice.
- 2. Pipette 40 µL of the Second Strand cDNA Synthesis product into the End Repair tubes.
  - a. Dissolve, mix and spin down.
  - b. Return tubes to ice.
- 3. Transfer reactions to a preheated thermal cycler and initiate an incubation using the following program and guidelines:
  - Heated lid off. If the lid is still hot, consider using a different thermal cycler or leaving the lid open.

End Repair Incubation conditions

Step	Temperature (°C)	Time (minutes)
1	25	30
2	4	Hold

- b. Place samples in the thermal cycler and start the program.
- c. When the run has completed, briefly spin down reactions and place on ice.

#### **Reaction Cleanup after Complete End Repair**

Refer to Important Precautions section for guidance on working with AMPure XP beads.

- 1. Completely resuspend AMPure XP beads by vortexing.
- 2. Remove tubes from ice and add 2.5X volume (100  $\mu$ L) of AMPure XP beads to each Complete End Repair reaction.
- 3. Vortex well or pipette 10 times to mix and visually inspect the color of the sample to ensure a homogenous mixture.
- 4. Incubate for **5 minutes** at room temperature (20°C to 25°C).
- 5. Briefly spin down tubes.





## FUSION Plex™ Protocol for Illumina®

- Place tubes on the magnet for 4 minutes or until beads are fully pelleted against the tube wall.
- 7. Without disturbing the bead pellet, use a pipette to remove and discard the supernatant. If the pellet becomes dislodged from the magnet and a portion is drawn into the pipette tip, return contents to the tube and repeat magnet incubation step.
- 8. Wash beads **two times** with 70% ethanol while still on the magnet. For each wash:
  - a. Add **200 µL** 70% ethanol.
  - b. Incubate for **30 seconds** at room temperature (20°C to 25°C).
  - c. Carefully remove ethanol and discard.
- 9. After the final wash, use a pipette (≤20 µL capacity) to completely remove visible supernatant residue and allow tubes to dry for **3-5 minutes** at room temperature with open lids. **Take care not to over-dry beads** as this will significantly decrease overall recovery (yield) of nucleic acid.
- 10. Elute DNA by resuspending beads in 20 µL 10mM Tris-HCl, pH 8.0.
- 11. Place tubes back on the magnet for 2 minutes.

### **Step 6: Ligation Step 1**

- 1. Place an appropriate number of Ligation Step 1 (SA0196) reaction tubes on ice.
- 2. Transfer **20 μL** of purified cDNA from Reaction Cleanup after End Repair, step 11 into Ligation Step 1 tubes. It is acceptable for a small amount of AMPure XP beads to be transferred.
  - a. Dissolve, mix and spin down.
  - b. Return tubes to ice.
- 3. Transfer reactions to a preheated thermal cycler and initiate an incubation using the following program and guidelines:
  - a. Use a heated lid (≥100°C).

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Ligation Step 1 Incubation conditions

Step	Temperature (°C)	Time (minutes)
1	37	15
2	4	Hold

b. Place samples in the thermal cycler and start the program.





c. After the program has reached 4°C, remove tubes from the temperature block, briefly spin down reactions and place on ice.

#### **Reaction Cleanup after Ligation Step 1**

Refer to Important Precautions section for guidance on working with AMPure XP beads.

- 1. Completely resuspend AMPure XP beads by vortexing.
- 2. Remove tubes from ice and add **2.5X** volume (**50**  $\mu$ L) of AMPure XP beads to each Ligation Step 1 reaction.
- 3. Vortex well or pipette 10 times to mix and visually inspect the color of the sample to ensure even mixing.
- 4. Incubate for **5 minutes** at room temperature (20°C to 25°C).
- 5. Briefly spin down tubes.
- 6. Place tubes on the magnet for 4 minutes or until beads are fully pelleted against the tube wall.
- 7. Without disturbing the bead pellet, use a pipette to remove and discard the supernatant. If the pellet becomes dislodged from the magnet and a portion is drawn into the pipette tip, return contents to the tube and repeat magnet incubation step.
- 8. Wash beads **two times** with 70% ethanol while still on the magnet. For each wash:
  - a. Add **200 µL** 70% ethanol.
  - b. Incubate for **30 seconds** at room temperature (20°C to 25°C).
  - c. Carefully remove ethanol and discard.
- 9. After the final wash, use a pipette (≤20 µL capacity) to completely remove visible supernatant residue and allow tubes to dry for **3-5 minutes** at room temperature with open lids. Take care not to over-dry beads as this will significantly decrease overall recovery (yield) of nucleic acid.
- 10. Elute DNA by resuspending beads in 42 μL 10 mM Tris-HCl, pH 8.0.
- 11. Place tubes back on the magnet for 2 minutes.

### **Step 7: MBC Adapter Incorporation**

1. Label MBC adapter tubes with the sample index tag letter (A, B, or C) and number (1-48) from the MBC adapters pouch label.



#### FUSION*Plex*™ Protocol for Illumina®

a. Use a permanent laboratory marker and orient lid hinges to the back as illustrated:



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Important: As this step incorporates the P5 index tag for sample-level tracking, be sure to record which MBC adapter is being used for each sample. Unused tubes must be labeled before returning to the pouch.

- 2. Place an appropriate number of MBC adapter reaction tubes on ice.
- 3. Add **40 μL** of the purified cDNA sample from Reaction Cleanup after Ligation Step 1, step 11 to a unique MBC adapter reaction tube. **Avoid pipetting AMPure XP beads into this reaction.** 
  - a. If AMPure XP beads are carried over, place the MBC adapter tubes on the magnet for one minute and transfer all liquid to the Ligation Step 2 tubes while the MBC adapter tubes remain on the magnet.
  - b. Dissolve, mix and spin down.
- 4. Immediately proceed to Step 8: Ligation Step 2.

#### **Step 8: Ligation Step 2**

- 1. Place an appropriate number of **Ligation Step 2** (SA0197) reaction tubes on ice.
- 2. Transfer the **entire volume** of each purified cDNA sample from Step 7: MBC Adapter Incorporation to Ligation Step 2 tubes.
  - a. Dissolve, mix and spin down.
  - b. Return tubes to ice.
- 3. Transfer reactions to a thermal cycler and initiate an incubation using the following program and guidelines:



a. Heated lid off.

Ligation Step 2 Incubation conditions

Step	Temperature (°C)	Time (minutes)
1	22	5
2	4	Hold

b. After the program has reached 4°C, briefly spin down reactions and place on ice.





**Safe stopping point:** It is okay to stop and store the reactions at -30°C to -10°C.

#### **Reaction Cleanup after Ligation Step 2**



**Caution:** This step uses Ligation Cleanup Beads (SA0210) and Ligation Cleanup Buffer (SA0209) instead of AMPure XP beads and 70% ethanol. Make sure to use Ligation Cleanup Beads and Ligation Cleanup Buffer that come with the kit for this step.

#### **Prepare Ligation Cleanup Beads:**

- 1. Completely resuspend Ligation Cleanup Beads by vortexing.
- 2. For each reaction, pipette **50 μL** of Ligation Cleanup Beads into new 0.2 mL 8-strip tubes.
- 3. Place tube(s) on the magnet for 1 minute or until the beads are pelleted.
- 4. Without disturbing the bead pellet, use a pipette to remove and discard the supernatant. If the pellet becomes dislodged from the magnet and a portion is drawn into the pipette tip, return contents to the tube and repeat magnetic pelleting step.
- 5. Pipette **50 µL of Ligation Cleanup Buffer** into each tube to resuspend beads.

#### **Ligation Cleanup Procedure:**



Caution: When vortexing PCR tubes in the subsequent steps, maintain firm pressure on all lids as the contained detergent may allow lids to open.

- 1. Pipette the entire volume of Ligation Step 2 reaction into the tubes with Ligation Cleanup Beads and Buffer.
- Mix samples by vortexing.
- 3. Incubate reactions at room temperature for 5 minutes.
- 4. Mix samples by vortexing.
- 5. Incubate reactions at room temperature for **5 minutes**.
- 6. Briefly spin down tubes.
- 7. Place tubes on the magnet for 1 minute or until beads are fully pelleted against the tube wall.
- 8. Carefully pipette off and discard supernatant (100 μL) without disturbing the beads.
- 9. Wash beads two times with Ligation Cleanup Buffer.



#### FUSION*Plex*™ Protocol for Illumina®

- a. Resuspend beads in **200 µL** Ligation Cleanup Buffer by vortexing, briefly spinning down, and placing back on magnet for **1 minute**.
- b. Once slurry has cleared, discard supernatant.
- 10. Wash beads **once** with **ultrapure water**:
  - a. Resuspend beads in **200 µL** of ultrapure water by vortexing, briefly spinning down and placing back on magnet.
  - b. Once slurry has cleared, discard supernatant.
  - c. Take care to ensure that all supernatant has been removed from tubes.
- 11. Elute DNA from Ligation Cleanup Beads:
  - a. Resuspend ligation cleanup beads in 18 μL of 5 mM NaOH.
  - b. Transfer reactions to a thermal cycler and initiate an incubation using the following program and guidelines:
  - c. Use a heated lid (≥100°C).

Ligation Elution incubation conditions

Step	Temperature (°C)	Time (minutes)
1	75	10
2	4	Hold

d. After sample has reached 4°C, briefly spin down and transfer to the magnet.

### Step 9: First PCR

- 1. Place an appropriate number of First PCR (SA0109) reaction tubes on ice.
  - a. Label tubes by sample number.
- 2. Pipette 2 μL of GSP1 (red cap) into each First PCR tube.
  - a. Spin down and return tubes to ice.
- 3. Pipette **18 µL** of supernatant from Reaction Cleanup after Ligation Step 2, step 11 into appropriately labeled First PCR tubes.
  - a. Dissolve, mix and spin down.
  - b. Return tubes to ice.





- 4. Transfer reactions to a preheated thermal cycler and immediately initiate the program specified in the panel specific **Product Insert (First PCR Reaction).** 
  - a. Use a heated lid (≥100°C).
  - b. After the program has reached 4°C, briefly spin down reactions and place on ice. It is also acceptable to leave tubes in the thermal cycler at 4°C overnight.

#### **Reaction Cleanup after First PCR**

Refer to Important Precautions section for guidance on working with AMPure XP beads.

- 1. Completely resuspend AMPure XP beads by vortexing.
- 2. Add 1.2X volume (24 µL) of AMPure XP beads to each reaction.
- 3. Vortex well or pipette 10 times to mix and visually inspect the color of the sample to ensure a homogenous mixture.
- 4. Incubate for **5 minutes** at room temperature (20°C to 25°C).
- 5. Briefly spin down tubes.
- 6. Place tubes on the magnet for 4 minutes or until beads are fully pelleted against the tube wall.
- 7. Without disturbing the bead pellet, use a pipette to remove and discard the supernatant. If the pellet becomes dislodged from the magnet and a portion is drawn into the pipette tip, return contents to the tube and repeat magnet incubation step.
- 8. Wash beads **two times** with 70% ethanol while still on the magnet. For each wash:
  - a. Add **200 μL** 70% ethanol
  - b. Incubate for **30 seconds** at room temperature (20°C to 25°C)
  - c. Carefully remove ethanol and discard
- 9. After the final wash, use a pipette (≤20 µL capacity) to completely remove visible supernatant residue and allow tubes to dry for **3-5 minutes** at room temperature with open lids. **Take care not to over-dry beads** as this will significantly decrease overall recovery (yield) of nucleic acid.
- **10.** Elute DNA by resuspending beads in **20 μL** of 10 mM Tris-HCl, pH 8.0.
- 11. Place tubes back on the magnet for **2 minutes**.
- 12. Transfer **20 μL** of purified eluate to a new 0.2mL PCR tube and store reactions as indicated below or proceed directly to Step 10: Second PCR.



### FUSION*Plex*™ Protocol for Illumina®



**Safe stopping point:** It is okay to stop and store the reactions at -30°C to -10°C (or leave in thermal cycler on 4°C hold for no more than 24 hours).

### Step 10: Second PCR

1. Place an appropriate number of **Second PCR** (SA0110) reaction tubes on ice.

#### Important: The Index 1 (P7) index tag is incorporated during this step.

 Use a permanent marker to label the tubes 1 to 8 from left to right as shown below. (Review the Molecular Barcoding, Sample Indexing & Multiplexed Sequencing section above).



- b. Unused tubes must be labeled before returning to the pouch.
- 2. Pipette 2 µL of GSP2 (black cap) into each Second PCR tube.
  - a. Record which P7 index sequence is used with which sample.
- 3. Pipette 18 µL of purified First PCR reaction into each Second PCR tube.
  - a. Dissolve, mix and spin down.
  - b. Return tubes to ice.
- 4. Transfer reactions to a preheated thermal cycler and immediately initiate the program specified in the panel specific **Product Insert (Second PCR Reaction).** 
  - a. Use a heated lid (≥100°C).
  - b. After the program has reached 4°C, briefly spin down reactions and place on ice. It is also acceptable to leave tubes in the thermal cycler at 4°C overnight.

#### Reaction Cleanup after Second PCR

Refer to Important Precautions section for guidance on working with AMPure XP beads.

- 1. Completely resuspend AMPure XP beads by vortexing.
- 2. Add 1.2X volume (24 μL) of AMPure XP beads to each Second PCR reaction.
- 3. Vortex well or pipette 10 times to mix and visually inspect the color of the sample to ensure a homogenous mixture.





- 4. Incubate for **5 minutes** at room temperature (20°C to 25°C).
- 5. Briefly spin down tubes.
- 6. Place tubes on the magnet for 4 minutes or until beads are fully pelleted against the tube wall.
- 7. Without disturbing the bead pellet, use a pipette to remove and discard the supernatant. If the pellet becomes dislodged from the magnet and a portion is drawn into the pipette tip, return contents to the tube and repeat magnet incubation step.
- 8. Wash beads **two times** with 70% ethanol while still on the magnet. For each wash:
  - a. Add **200 µL** 70% ethanol.
  - b. Incubate for **30 seconds** at room temperature (20°C to 25°C).
  - c. Carefully remove ethanol and discard.
- 9. After the final wash, use a pipette (≤20 µL capacity) to completely remove visible supernatant residue and allow tubes to dry for **3-5 minutes** at room temperature with open lids. **Take care not to over-dry beads** as this will significantly decrease overall recovery (yield) of nucleic acid.
- 10. Elute DNA by resuspending beads in **20 μL** 10 mM Tris-HCl, pH 8.0.
- 11. Place tubes back on the magnet for **2 minutes**.
- 12. Transfer **18 μL** of the purified solution to a new 0.2mL PCR tube. Be sure to avoid transferring beads to the fresh tube.
- 13. Stop or proceed directly to Quantify, Normalize and Sequence.



Safe stopping point: It is okay to stop and store the reactions at -30°C to -10°C.



### **Quantify, Normalize, and Sequence**

Refer to the Quantify, Normalize, and sequence Protocol for Illumina (RA-DOC-054) for information on quantifying, normalizing and sequencing Archer AMP libraries.

#### **Data analysis**

Archer AMP libraries should be demultiplexed to FASTQs prior to analysis. Analyze data with Archer Analysis using either a local software installation or Archer Unlimited. Visit our website or contact archertech@idtdna.com for more information.

FUSION*Plex* assays require a one-time upload of a GTF file (a text file, in GTF format, which directs the software on how to analyze data from the panel). Additionally, if the SNV/indel pipeline is chosen, there is an option to select a targeted mutation file (a text file, in VCF format, which lists specific variants of interest). This file also requires a one-time upload. Both files can be obtained by contacting archertech@idtdna.com.

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