

Arima-HiC Kit

User Guide for Nucleated Blood 8 reactions

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U.S. Patent No. US 9,434,985 pertains to the use of this product.

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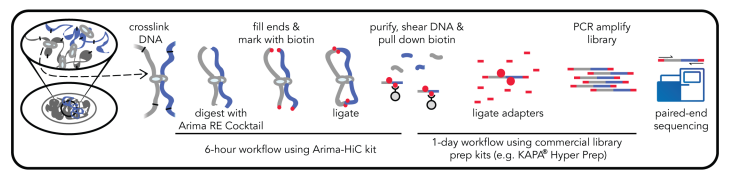
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Introduction



1.1 Arima-HiC Workflow Overview

Arima-HiC is an experimental workflow that captures the sequence and structure (three-dimensional conformation) of genomes. Arima-HiC has been successfully performed on a wide-range of species from the plant and animal kingdoms. As illustrated in the Arima-HiC workflow schematic above, chromatin from a sample source (tissues, cell lines, or blood) is first crosslinked to preserve the genome sequence and structure. The crosslinked chromatin is then digested using a restriction enzyme (RE) cocktail. The 5'-overhangs are then filled in, causing the digested ends to be labeled with a biotinylated nucleotide. Next, spatially proximal digested ends of DNA are ligated, capturing the sequence and structure of the genome. The ligated DNA is then purified, producing pure proximally-ligated DNA. The proximally-ligated DNA is then fragmented, and the biotinylated fragments are enriched. The enriched fragments are then subjected to a <u>custom</u> library preparation protocol utilizing a range of supported commercially available library prep kits. Depending on the choice of library prep kit, a separate Arima-HiC Library Prep user guide is provided that contains a custom protocol for converting proximally-ligated DNA to Arima-HiC libraries.

1.2 Sequencing and Data Analysis

Arima-HiC libraries are sequenced via Illumina® sequencers in "paired-end" mode. The resulting data is referred to as Arima-HiC data. The tools necessary for analyzing Arima-HiC data depend on the application. For example, for studying 3D genome conformation, Arima-HiC data can be processed using publicly available tools such as Juicer (Durand, 2016a) or Hi-C Pro (Servant, 2015), and genome organizational features such as compartments, TADs, and loops can be identified and visualized using tools such as Juicebox (Durand, 2016b). These tools require usage modifications and/or custom input files that are specific to Arima-HiC data, so please contact Technical Support for assistance implementing these tools. Additionally, because paired-end reads of Arima-HiC data can originate from distal sequences along the linear genome, these data capture short- and long-range DNA contiguity information that is valuable for applications such as *de novo* assembly and genome scaffolding. Therefore, Arima-HiC data can be mapped to contigs/unitigs using our mapping pipeline (https://github.com/ArimaGenomics) or Juicer, and then the contigs/unitigs can be scaffolded using tools such as SALSA (Ghurye, 2019) or 3D-DNA (Dudchenko, 2017). Please contact Technical Support for more information.

Arima-HiC Quick Reference Protocol



Kit P/N: A510008 Doc P/N: A160259 Date: Nov 2018

Arima-HiC Quick Reference Protocol



Crosslink Sample



Resuspend cells in 20µL ● Lysis Buffer OR nuclei in 20µL water¹ Incubate 15 min. at 4°C



Add 24µL O Conditioning Solution Incubate 10 min. at 62°C



Add 20µL● **Stop Solution 2** Incubate 15 min. at 37°C



Add $7\mu L \odot$ Buffer A + $1\mu L \odot$ Enzyme A1 + $4\mu L \odot$ Enzyme A2 Incubate 30 min. at 37°C followed by 20 min. at 62°C



Add 12µL ● Buffer B + 4µL ● Enzyme B Incubate 45 min. at 25°C



Add 70µL ● Buffer C + 12µL ● Enzyme C Incubate 15 min. at 25°C



Add $10.5\mu L \odot Buffer D + 25\mu L \odot Enzyme D$ Add $20\mu L \odot Buffer E$ Incubate 30 min. at 55°C followed by 90 min. at 68°C



Purify DNA

Proceed to library prep using the unique protocol outlined in the User Guide

¹ If nuclei have been isolated prior to lysis step (e.g. from plant tissue, FAC-sorted nuclei)

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Arima-HiC Kit Contents and Storage Info



Kit P/N: A510008 Doc P/N: A160259 Date: Nov 2018

Arima-HiC Box Contents & Storage Info

Box A							
Component	Сар	Storage					
Stop Solution 1	()						
Elution Buffer	()						
Wash Buffer*	()						
Conditioning Solution	0	20 to 25°C					
Stop Solution 2	•						
Buffer D							
Buffer E							

Вох В							
Component	Сар	Storage					
Lysis Buffer							
Buffer A	<u> </u>						
Enzyme A1	0						
Enzyme A2	0						
Buffer B	0	-20°C					
Enzyme B							
Buffer C	0						
Enzyme C							
Enzyme D							

Box C							
Component Cap Storage							
Enrichment Beads*	0	2 to 8°C					
QC Beads**		21080					

^{*} Required for library prep. Depending on choice of library prep kit, the User Guide has a unique library prep protocol to be followed

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^{**} Required for Arima-QC1. QC Beads are not interchangeable with Enrichment Beads

Getting Started

2.1 Handling and Preparation

- Several steps during the *Arima-HiC Protocol* require preparation of a master mix. Sufficient reagent has been included in the kit to make master mixes with 10% excess volume. Use the master mix calculation tables provided.
- When handling reagents, room temperature (RT) is defined as 20 to 25°C.
- If the Arima-HiC Protocol is performed in PCR plates or PCR tubes, ensure to have a total volume capacity of at least 320µL. See Section 2.2 for recommended PCR plates and PCR tubes. Also, ensure that plates and/or tubes are compatible with thermal cyclers and other required equipment. Using seals and caps for PCR plates and tubes is required.
- All kit reagents should be fully thawed and thoroughly mixed before use.
- Stop Solution 1, Conditioning Solution, and Buffer D from Box A may contain precipitates. If present, these precipitates must be dissolved before use. Heating these reagents at 37°C for 5-15 minutes may be necessary to dissolve precipitates.
- During handling and preparation, reagents from **Box A** should be kept at RT.
- During handling and preparation, reagents from Box B should be kept on ice, except for Enzyme D, which should be kept on ice but warmed to room temperature just before use.
- Enzyme solutions from Box B are viscous and require special attention during pipetting.

2.2 User-supplied reagents, consumables, and equipment checklist

Freshly prepared Resuspension Buffer (see Section 2.3 for recipe)

1X PBS, pH 7.4 (e.g. Fisher Scientific® Cat # 50-842-949)

37% Formaldehyde (e.g. Fisher Scientific® Cat # F79-500)

Freshly prepared 80% Ethanol

DNA Purification Beads (e.g. Beckman Coulter Cat # A63880)

Qubit® Fluorometer, dsDNA HS Assay Kit and consumables (e.g. Thermo Fisher Scientific Cat # 32851, 32856)

Liquid Nitrogen or dry ice

15mL conical tubes

1.7mL microcentrifuge tubes, PCR tubes (e.g. SSIbio[®] Cat # 3247-00), or PCR plates (e.g. Bio-Rad[®] Cat # HSS9641) and magnetic rack compatible with tube selection.

Centrifuge

Thermal cycler (if performing Arima-HiC in PCR tubes or PCR plate)

Thermomixer (if performing Arima-HiC in 1.7mL microcentrifuge tubes)

Fetal Bovine Serum (e.g. Fisher Scientific® Cat # A3160501)

2.3 Buffer Recipes

Resuspension Buffer – The **Resuspension Buffer** must be prepared fresh directly before use in the *Crosslinking* protocol. The following recipe is enough for crosslinking 8 samples. This recipe should be scaled accordingly if more or less than 8 samples are processed simultaneously. If using a 16% formaldehyde stock, please contact Technical Support for a different **Resuspension Buffer** formulation. The table below includes a *suggested* vendor and catalog number for each reagent. After the **Resuspension Buffer** is prepared, store at 4°C until use.

Reagent	Stock Vendor	Stock Cat #	Stock Concentration	Final Concentration	Stock Amount
PBS	Fisher Scientific®	50-842-949			54.45mL
FBS	Fisher Scientific®	A3160501	100%	1%	550µL
				Total	55mL

2.4 Optimal read length, sequencing depth, and number of Arima-HiC reactions per sample

Arima-HiC libraries must be sequenced in paired-end mode, and are compatible with most Illumina® sequencing machines (e.g. MiSeq®, NextSeq®, HiSeq®, NovaSeq™) and a variety of read lengths. We generally recommend 2x150bp read length on the HiSeq® or NovaSeq™ instruments to optimize for sequencing throughput and Arima-HiC data alignment quality, although shorter read lengths (e.g. 2x50bp, 2x100bp) and lower throughput instruments can certainly be used for certain applications of Arima-HiC data such as 3D genome conformation analysis and genome scaffolding.

The optimal sequencing depth for Arima-HiC libraries also depends on the application. For studying 3D genome conformation, the ability to detect certain genome organization features depends on the sequencing depth. For ~3Gb genomes such as mouse and human, we generally recommend obtaining at least 600 million read-pairs *per biological condition* for high-resolution analyses of A/B compartments, TADs, and chromatin loops. One way of obtaining at least 600 million read-pairs is by combining at least 300 million read-pairs from 2 biological replicates. In doing so, you will be able to assess the overall reproducibility of the Arima-HiC data across replicates, and then used the combined replicate Arima-HiC dataset for high-resolution chromatin conformation analyses. Alternatively, one can obtain at least 600 million read-pairs *per biological replicate* and then use the common set of identified genome conformational features across replicates as a "high confidence" set of structural features supported by their observation in both replicates. For lower resolution analyses of A/B compartments and TADs, we generally recommend obtaining at least 300 million read-pairs per biological condition. For help estimating the optimal sequencing depth for different genome sizes or analysis goals, please contact Technical Support.

For applications such as *de novo* assembly and genome scaffolding, the required sequencing depth can vary depending on the quality of contig/unitigs that are being scaffolded using Arima-HiC data.

For a 3Gb genome, we recommend obtaining up to 600M read-pairs, as this is the amount of sequencing that is currently utilized from Arima-HiC libraries for genome scaffolding by the Vertebrate Genome Project (VGP) consortia. The amount of sequencing required scales linearly with the genome size (e.g. up to 200M read-pairs for a 1Gb genome).

Lastly, it is important to note that each Arima-HiC library should pass the Arima-QC2 assay and be evaluated for library complexity prior to deep sequencing. As a general rule, each Arima-HiC library should be complex enough to sequence up to ~600M read-pairs without reaching saturation. If >600M read-pairs of Arima-HiC data are needed, it may be more efficient to sequence a second Arima-HiC library than sequence deeper into the first Arima-HiC library.

2.5 How to cite Arima-HiC in publications

When citing the Arima-HiC protocol or kit, one may write: "Hi-C data was generated using the Arima-HiC kit, according to the manufacturers protocols". Please reference the catalog number found on the kit packaging.

Crosslinking

Input: Whole nucleated blood in ethanol **Output:** Crosslinked nucleated blood cells

Before you begin: The Arima-HiC workflow for nucleated blood begins with washing of ethanol-preserved nucleated blood, followed by crosslinking. The protocol assumes one knows how much whole blood (in μ L) was originally collected and then preserved in ethanol. Less than 25 μ L of whole nucleated blood (i.e. not the volume of the ethanol diluted blood) is typically needed for a single Arima-HiC reaction, but we recommend crosslinking 25 μ L of whole nucleated blood if sufficient blood is available. For example in Step 2 below, if 50 μ L of whole blood was collected in 1 μ L of ethanol, then 500 μ L of the ethanol diluted blood would be used for crosslinking because it contains the equivalent of 25 μ L of whole nucleated blood. Lastly, we recommend performing all centrifugation steps for 5 min at 2000 x G.

- 1. Prepare Resuspension Buffer and 1X PBS. Chill buffers on ice until cold.
- 2. If only using a portion of whole blood preserved in ethanol for crosslinking, <u>resuspend</u> the whole blood in ethanol mixture thoroughly by pipetting and transfer a portion to a new 1.5mL tube for crosslinking and proceed to the next step. With the remaining blood and ethanol mixture, <u>freeze</u> on dry ice or liquid nitrogen and <u>store</u> at -80°C.
- 3. Pellet cells by centrifugation and remove supernatant.
- 4. Resuspend pellet in 1mL Resuspension Buffer.
- 5. <u>Pellet</u> cells by centrifugation and <u>remove</u> supernatant.
- 6. Resuspend pellet in 1mL Resuspension Buffer, and transfer to a 15mL conical tube.
- 7. <u>Add</u> 4mL **Resuspension Buffer**, bringing the total volume to 5mL.
- 8. Add 286µL of **37% formaldehyde**, bringing the final formaldehyde concentration to 2%.
- 9. Mix well by inverting 10 times and incubate at RT for 10 min.
- 10. Add 460µL of **Stop Solution 1**, mix well by inverting 10 times and incubate at RT for 5 min.
- 11. <u>Place</u> sample on ice and <u>incubate</u> for 15 min.
- 12. <u>Pellet</u> cells by centrifugation and <u>remove</u> supernatant.
- 13. Resuspend cells in 1mL 1X PBS.
- 14. To prepare for the *Estimating Input Amount* protocol in the following section, <u>mix</u> the sample by inversion and then immediately <u>aliquot</u> sample such that 1 aliquot contains 10% of the sample, while the rest of the aliquots each contain the equivalent of ~20-25% of the sample. <u>Mix</u> sample by inversion between aliquots to ensure all aliquots are equally homogeneous.

- * The 10% aliquot will be used in the *Estimating Input Amount* protocol. The remaining 3 aliquots containing 20-25% are meant to be saved as sample material for the *Arima-HiC Protocol*.
- 15. <u>Pellet</u> cells in all aliquots by centrifugation and <u>remove</u> supernatant leaving only the crosslinked cell pellet and no residual liquid.
- 16. <u>Freeze</u> samples on dry ice or liquid nitrogen, and <u>store</u> at -80°C until ready to proceed to the *Estimating Input Amount* protocol in the following section.

Estimating Input Amount

Input: 10% aliquot of crosslinked nucleated blood cells

Output: Purified genomic DNA

Before you begin: The Estimating Input Amount protocol is required if one does not know how many crosslinked cells will comprise 500ng-5µg of DNA, and if sufficient cells are available to perform this protocol. Arima-HiC reactions are optimally performed on crosslinked cells comprising ~500ng-5µg of DNA. The Estimating Input Amount protocol measures the amount of DNA obtained from 10% of the crosslinked cells, which guides the calculation of the optimal cellular input for an Arima-HiC reaction. The Arima-HiC kit contains enough reagents to perform this protocol on 8 samples. This protocol concludes with a descriptive example of how to estimate the optimal amount of crosslinked cells to use per Arima-HiC reaction.

Note: Step 2 requires addition of several reagents in the same step. These reagents should be combined into master mixes with 10% excess volume before use.

- 1. Thaw the aliquot containing 10% of the crosslinked cells prepared during the *Crosslinking* protocol.
- 2. Add 209.5µL of a master mix containing the following reagents:

Reagent	Volume per reaction	10% extra		# reactions		Final
Elution Buffer	174µL	191.4µL	х	2	=	382.8µL
Buffer D	10.5µL	11.55µL	х	2	=	23.1µL
Enzyme D	25µL	27.5µL	х	2	=	55µL
Total	209.5μL					460.9µL

3. Add 20µL of ■ Buffer E, mix gently by pipetting, and incubate as follows. If using a thermal cycler, set the lid temperature to 85°C.

Temperature	Time
55°C	30 min.
68°C	90 min.
4°C	8

Note: DNA Purification Beads (e.g. AMPure® XP Beads) should be warmed to RT and thoroughly mixed before use. The DNA Purification Beads are a *user-supplied reagent* and should not be mistaken for the Enrichment Beads or QC Beads provided in the Arima-HiC kit.

4. <u>Add</u> 150μL of **DNA Purification Beads**, <u>mix</u> thoroughly, and <u>incubate</u> at RT for 5 min.

- 5. <u>Place</u> sample against magnet, and <u>incubate</u> until solution is clear.
- 6. <u>Discard</u> supernatant. While sample is still against magnet, <u>add</u> 400µL of 80% ethanol, and incubate at RT for 1 min.
- 7. <u>Discard</u> supernatant. While sample is still against magnet, <u>add</u> 400µL of 80% ethanol, and incubate at RT for 1 min.
- 8. <u>Discard</u> supernatant. While sample is still against magnet, <u>incubate</u> beads at RT for 3 5 min. to air-dry the beads.
- 9. Remove sample from magnet, <u>resuspend</u> beads thoroughly in 20µL of **Elution Buffer**, and <u>incubate</u> at RT for 5 min.
- 10. <u>Place</u> sample against magnet, <u>incubate</u> until solution is clear, and <u>transfer</u> supernatant to a new tube.
- 11. <u>Quantify</u> sample using Qubit[®]. The total DNA yield corresponds to the amount of DNA obtained from 10% of the crosslinked cells.
- 12. <u>Estimate</u> how much crosslinked nucleated blood cells to use per Arima-HiC reaction. See the example description below:

Example: In the following *Arima-HiC Protocol*, it is recommended to use crosslinked cells corresponding to at least 500ng of DNA per Arima-HiC reaction, but no more than 5μg of DNA. If 250ng of DNA was obtained from 10% of the crosslinked cells as calculated in step 11, one can estimate that *at least* 20% of the original crosslinked cells should be used per Arima-HiC reaction (~500ng of DNA). More crosslinked cells should be used if available, as long as the total DNA per reaction is not more than 5μg. If possible, we recommend aiming to use crosslinked cells comprising 3μg of DNA per Arima-HiC reaction. Additionally, please note that the crosslinked cell pellet for one Arima-HiC reaction should occupy no more than 20μL of volume in the sample tube. If the crosslinked cell pellet comprises 500ng-5μg of DNA but occupies greater than 20μL of volume, aliquot the cells into multiple Arima-HiC reactions such that the sum of the DNA input from all reactions is at least 500ng and each cell pellet occupies no more than 20μL of volume, or contact Technical Support for additional guidance.

Recommended HiC Input Amount Explanation: The recommendation to use crosslinked cells comprising at least 500ng of DNA is only a *general* recommendation. If crosslinked cells comprising at least 500ng of DNA cannot be obtained, one should proceed with the Arima-HiC Protocol as described in this user guide and then use our validated low-input library prep protocol.

Arima-HiC Protocol

Input: Crosslinked nucleated blood cells containing ~500ng-5µg of DNA

Output: Proximally-ligated DNA

Before you begin: The cell pellet for one Arima-HiC reaction should occupy no more than 20μL of volume and should be devoid of any residual liquid. If the cell pellet occupies greater than 20μL of volume, aliquot the cells such that the sum of the DNA input from all reactions is between 500ng-5μg and each cell pellet occupies no more than 20μL of volume, or contact Technical Support for additional guidance. Note that steps 2 – 3 require consecutive heated incubations. Make sure your thermal device(s) are set to 62°C and 37°C for these incubations. The safe stopping point in this section is after completing Step 21.

Note: Choose to perform either Step 1a if the input sample type is crosslinked cells, or Step 1b only if the input sample type is crosslinked *nuclei* that have been previously purified from cells.

- 1. <u>Resuspend</u> one reaction of crosslinked cells in 20µL of **Lysis Buffer** in a tube or a well of a PCR plate, and <u>incubate</u> at 4°C for 15 min.
- 2. Add 24μL of O Conditioning Solution, mix gently by pipetting, and incubate at 62°C for 10 min. If using a thermal cycler, set the lid temperature to 85°C.
- 3. Add 20µL of Stop Solution 2, mix gently by pipetting, and incubate at 37°C for 15 min. If using a thermal cycler, set the lid temperature to 85°C.

Note: Steps 4, 6, 8 and 10 require addition of several reagents in the same step. These reagents should be combined into master mixes following the master mix tables.

4. Add 12µL of a master mix containing the following reagents:

Reagent	Volume per reaction	10% extra		# reactions		Final
Buffer A	7µL	7.7µL	X	2	II	15.4µL
Enzyme A1	1µL	1.1µL	Х	2	=	2.2µL
Enzyme A2	4µL	4.4µL	Х	2	=	8.8µL
Total	12µL					26.4µL

5. <u>Mix</u> gently by pipetting, and <u>incubate</u> as follows. If using a thermal cycler, set the lid temperature to 85°C. Note that there are sequential incubations at different temperatures:

Temperature	Time
37°C	60 min.
65°C	20 min.
25°C	10 min.

6. Add 16µL of a master mix containing the following reagents:

Reagent	Volume per reaction	10% extra		# reactions		Final
Buffer B	12µL	13.2µL	Х	2	=	26.4µL
Enzyme B	4µL	4.4µL	х	2	=	8.8µL
Total	16µL					35.2µL

- 7. Mix gently by pipetting, and incubate at room temperature (RT) for 45 min.
- 8. Add 82µL of a master mix containing the following reagents:

Reagent	Volume per reaction	10% extra		# reactions		Final
Buffer C	70μL	77μL	х	2	=	154µL
Enzyme C	12µL	13.2µL	Х	2	=	26.4µL
Total	82µL					180.4µL

9. Mix gently by pipetting, and incubate at RT for 15 min.

Note: Enzyme D should be warmed to RT to prevent precipitation in the below master mix.

10. Add 35.5µL of a master mix containing the following reagents:

Reagent	Volume per reaction	10% extra		# reactions		Final
Buffer D	10.5µL	11.55µL	х	2	=	23.1µL
Enzyme D	25µL	27.5µL	Х	2	=	55µL
Total	35.5µL					78.1µL

11. Add 20µL of ● Buffer E, mix gently by pipetting, and incubate as follows. If using a thermal cycler, set the lid temperature to 85°C.

Temperature	Time		
55°C	30 min.		
68°C	90 min.		
25°C*	10 min.*		

^{*} To provide flexibility, this incubation can also be held overnight at 4°C. *Do not* incubate at 68°C for longer than 90 min. unless doing so using a thermal cycler with a heated lid.

Note: DNA Purification Beads (e.g. AMPure® XP Beads) should be warmed to RT and thoroughly mixed before use. The DNA Purification Beads are a *user-supplied reagent* and should not be mistaken for the Enrichment Beads or QC Beads provided in the Arima-HiC kit.

- 12. Add 100µL of DNA Purification Beads, mix thoroughly, and incubate at RT for 5 min.
- 13. <u>Place</u> sample against magnet, and <u>incubate</u> until solution is clear.
- 14. <u>Discard</u> supernatant. While sample is still against magnet, <u>add</u> 300µL of 80% ethanol, and <u>incubate</u> at RT for 1 min.
- 15. <u>Discard</u> supernatant. While sample is still against magnet, <u>add</u> 300µL of 80% ethanol, and incubate at RT for 1 min.
- 16. <u>Discard</u> supernatant. While sample is still against magnet, <u>incubate</u> beads at RT for 3 5 min. to air-dry the beads.
- 17. <u>Remove</u> sample from magnet, <u>resuspend</u> beads thoroughly in 100µL of **Elution Buffer**, and incubate at RT for 5 min.
- 18. <u>Place</u> sample against magnet, <u>incubate</u> until solution is clear, and <u>transfer</u> supernatant to a new tube.
- 19. Quantify sample using Qubit®.

Note: If the proximally-ligated DNA yield is less than 275ng, we recommend skipping the Arima-QC1 assay mentioned in Step 20 and described in the following *Arima-QC1 Quality Control* section, and strongly recommend performing the Arima-QC2 assay described in our *Arima-HiC Library Preparation* user guide for low input samples.

- 20. <u>Transfer</u> 75ng of sample into a new tube labelled "Arima-QC1", and <u>add</u> Elution Buffer to Arima-QC1 to bring the volume to 50μL. The "Arima-QC1" sample should now contain 75ng of proximally-ligated DNA in 50μL of Elution Buffer. <u>Store</u> at -20°C until use in the following *Arima-QC1 Quality Control* protocol.
- 21. <u>Store</u> all remaining samples at -20°C until ready to proceed to library preparation following an accompanying *Arima-HiC Library Preparation* user guide.

Arima-QC1 Quality Control

Before you begin: The following protocol quantifies the fraction of proximally-ligated DNA that has been labeled with biotin, and is a quality control metric after completing the *Arima-HiC Protocol* but before proceeding to library preparation. The *Arima-QC1 Quality Control* protocol involves using QC Beads to enrich an aliquot of proximally-ligated DNA, which is then quantified using a Qubit® fluorometer. Unlike standard Qubit® readings which involve quantifying a transparent unobstructed DNA sample, the Arima-QC1 value is obtained by quantifying DNA that is still bound to the QC Beads. This protocol can be performed in either plates or tubes. Set your thermal device (thermal cycler or thermomixer) to hold at 55°C. After completing the *Arima-QC1 Quality Control* protocol, use the provided **Arima-HiC QC** Worksheet to determine the Arima-QC1 values.

- 1. If necessary, <u>thaw</u> the "Arima-QC1" samples prepared during Step 20 of the *Arima-HiC Protocol* in the previous section.
- 2. Add 50µL of QC Beads, mix thoroughly by pipetting, and incubate at RT for 15 min.
- 3. <u>Place</u> sample against magnet, and <u>incubate</u> until solution is clear.
- 4. <u>Discard</u> supernatant, and <u>remove</u> sample from magnet.
- 5. Wash beads by resuspending in 200µL of Wash Buffer, and incubate at 55°C for 2 min.
- 6. <u>Place</u> sample against magnet, and <u>incubate</u> until solution is clear.
- 7. <u>Discard</u> supernatant, and <u>remove</u> sample from magnet.
- 8. Wash beads by resuspending in 200µL of Wash Buffer, and incubate at 55°C for 2 min.
- 9. <u>Place</u> sample against magnet, and <u>incubate</u> until solution is clear.
- 10. <u>Discard</u> supernatant, and <u>remove</u> sample from magnet.
- 11. Wash beads by resuspending in 100µL of Elution Buffer.
- 12. <u>Place</u> sample against magnet, and <u>incubate</u> until solution is clear.
- 13. <u>Discard</u> supernatant, and <u>remove</u> sample from magnet.
- 14. Resuspend beads in $7\mu L$ of Elution Buffer. Proceed to next step with resuspended beads.

Note: The following step involves the quantification of the *bead-bound* DNA using the Qubit® dsDNA HS Assay Kit.

- 15. <u>Quantify</u> the total amount of *bead-bound DNA* using Qubit[®]. Use 2µL of thoroughly mixed bead-bound DNA for the Qubit[®] assay.
- 16. <u>Determine</u> the **Arima-QC1** value by following the **Arima-HiC QC Worksheet**. High quality Arima-QC1 values are expected to be >15%. If the Arima-QC1 value did not obtain a 'PASS' status, please contact Technical Support for troubleshooting assistance.

Warranty and Contact Info

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CONTACT US

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