

INSTRUCTIONS FOR USE OF

KAPA mRNA HyperPrep Kit

FOR RESEARCH USE ONLY. NOT FOR USE IN DIAGNOSTIC PROCEDURES.

March 2024, Version 9.0





Contents

The KAPA mRNA HyperPrep Kits contain:

Kit Material Number	Description	Volume
	mRNA Capture Beads	1.3 mL
KK8580	mRNA Bead Binding Buffer	3.9 mL
08098115702	mRNA Bead Wash Buffer	9.6 mL
24 Libraries	Fragment, Prime and Elute Buffer (2X)	300 μL
consists of the following sub kits:	1st Strand Synthesis Buffer	300 μL
KK8542	KAPA Script	25 μL
08105936001	2nd Strand Marking Buffer	780 µL
and	2nd Strand Synthesis & A-Tailing Enzyme Mix	50 μL
KK8543	Ligation Buffer	1 mL
08105880001	DNA Ligase	280 μL
and	PEG/NaCl Solution	1 mL
KK8440	KAPA Pure Beads	3.2 mL
07962231001	Library Amplification Primer Mix (10X)	138 µL
	KAPA HiFi HotStart ReadyMix (2X)	690 μL
	mRNA Capture Beads	5.1 mL
KK8581	mRNA Bead Binding Buffer	15.3 mL
08098123702 96 Libraries	mRNA Bead Wash Buffer	40 mL
	Fragment, Prime and Elute Buffer (2X)	1.4 mL
consists of the following sub kits:	1st Strand Synthesis Buffer	1.4 mL
KK8544	KAPA Script	130 μL
08105952001	2nd Strand Marking Buffer	3.8 mL
and	2nd Strand Synthesis & A-Tailing Enzyme Mix	250 μL
KK8545	Ligation Buffer	5 mL
08105901001	DNA Ligase	1.26 mL
and	PEG/NaCl Solution	5 mL
KK8441	KAPA Pure Beads (3 x 5 mL bottles)	15 mL
07962240001	Library Amplification Primer Mix (10X)	600 μL
	KAPA HiFi HotStart ReadyMix (2X)	3 mL

Note: Accessory kits (KAPA Universal Adapter & KAPA UDI Primer Mixes and KAPA Unique Dual-Indexed Adapters) are sold separately.



Shipping, Storage and Stability

- KAPA Pure Beads and mRNA capture beads and buffers are shipped on dry ice or ice packs, depending on the destination country. Upon receipt, store KAPA Pure Beads and mRNA capture beads and buffers at +2°C to +8°C.
- Enzymes and buffers for cDNA synthesis and library preparation are shipped on dry ice or ice packs, depending on the destination country. These components are temperature sensitive, and appropriate care should be taken during storage.
- Upon receipt, store these reagents at -15°C to -25°C in a constant-temperature freezer.
- The 1st Strand Synthesis Buffer, PEG/NaCl Solution, and KAPA Pure Beads are light sensitive and should be protected from light during storage.
- When stored under these conditions and handled correctly, the kit components will retain full activity until the expiry date indicated on the kit label.

Applications

The KAPA mRNA HyperPrep Kit contains all of the buffers and enzymes required for poly(A) mRNA capture and the rapid construction of stranded mRNA-Seq libraries from 50 ng – 1 µg of intact total RNA. The KAPA mRNA HyperPrep Kit is designed for both manual and automated NGS library construction. The kit enriches for polyadenylated RNAs using magnetic oligo-dT beads. The protocol is applicable to a wide range of RNA-Seq applications, including:

- gene expression
- single nucleotide variation (SNV) discovery
- splice junction and gene fusion identification
- characterization of polyadenylated RNAs.



This kit is not compatible with small RNAs <100 bp in length.

Warnings and Precautions

- Wear the appropriate personal protective equipment, such as gloves and safety glasses, to avoid direct contact while handling the reagents. Wash hands thoroughly after handling samples and reagents.
- Use good laboratory practices to avoid contamination when working with the reagents.
- Do not eat, drink, or smoke in the laboratory area.
- Do not pipette by mouth.
- In the event of a spill, clean up the solution with absorbent pads, allow pads to dry, and dispose of pads.
- Observe all national, regional, and local regulations for waste disposal and management.
- Safety Data Sheets (SDS) are available online or upon request from the local Roche office.

Ordering Information

For a complete overview of Roche Sequencing products, including KAPA mRNA HyperPrep Kits, go to **sequencing.roche.com** products.

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Changes to Previous Versions

Workflow modifications when using KAPA HyperPlex Adapter Kits (KAPA Universal Adapter & KAPA UDI Primer Mixes) added. Branding changes from legacy Kapa Biosystems to Roche Sequencing Solutions.

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Preface

Regulatory Disclaimer

For Research Use Only.

Not for use in diagnostic procedures.

Contact Information

Technical Support

If you have questions, contact your local Roche Technical Support.

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Manufacturing & Distribution

Manufacturer Roche Dia Cape Town,

South Africa

Distribution Roche Diagnostics GmbH

Mannheim, Germany

Distribution in USA Roche Diagnostics Corporation

Indianapolis, IN USA



Conventions Used in This Manual

Symbols

Symbols	Description
\triangle	Important Note: Information critical to the success of the procedure or use of the product. Failure to follow these instructions could result in compromised data.
!	Information Note: Designates a note that provides additional information concerning the current topic or procedure.

Text

Conventions	Description
Numbered listing	Indicates steps in a procedure that must be performed in the order listed.
Italic type, blue	Highlights a resource in a different area of this manual or on a web site.
Italic type	Identifies the external general resources or names.
Bold type	Identifies names of paragraphs, sections or emphasized words.





Chapter 1. Before You Begin

These Instructions for Use describe the process for poly(A) mRNA capture and streamlined RNA library construction of stranded RNA libraries for the Illumina sequencing ecosystem. Specifically, these Instructions for Use provide a protocol for the workflow outlined in *Figure 1* using the KAPA mRNA HyperPrep Kit.

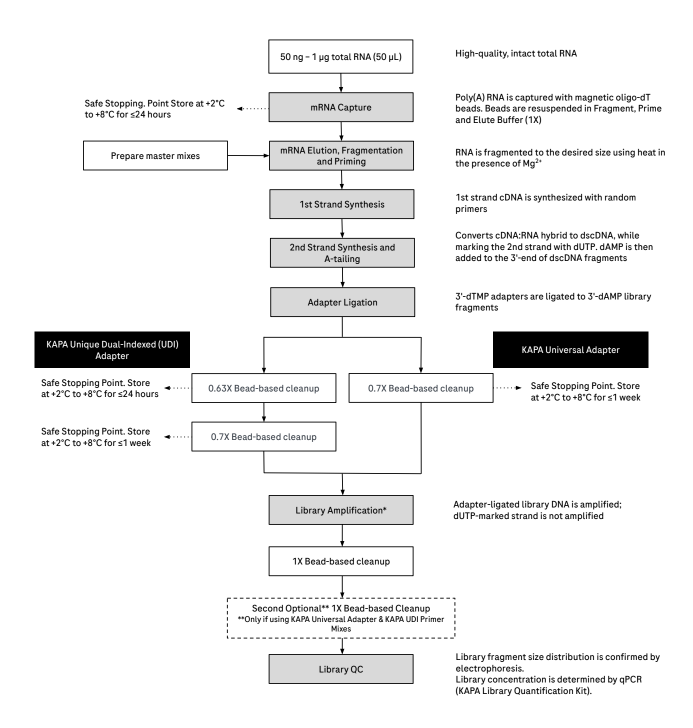


The KAPA mRNA HyperPrep Workflow provides:

- An easy to use, streamlined, and automation-friendly workflow with minimal resource requirements.
- Single vendor service when using the following accessory reagents:
 - KAPA Pure Beads
 - KAPA Unique Dual-Indexed (UDI) Adapters & KAPA Library Amplification Primer Mix or
 - KAPA Universal Adapter & KAPA UDI Primer Mixes 1 384



Figure 1 - KAPA mRNA HyperPrep Quick Guide



^{*}Using KAPA HiFi HS ReadyMix and either KAPA Library Amplification Primer Mix if KAPA UDI Adapters were used or KAPA UDI Primers Mixes if

KAPA Universal Adapter was used.

**A modified post-amplification cleanup is highly recommended when using KAPA Universal Adapter & KAPA UDI Primer Mixes with KAPA library preparation kits. Especially if libraries will be sequenced on an Illumina NovaSeq, NextSeq 1000/2000 or HiSeq X system (with patterned flow cells). This additional or modified cleanup will serve to remove any indexed primer carryover which may exacerbate index hopping.



To mix samples containing RNA, gently pipette the reaction mixture several times. Vortexing may fragment the RNA, resulting in lower quantity and a reduced library insert size.

Protocol Information & Safety

- Wear gloves and take precautions to avoid sample contamination.
- Perform all centrifugations at room temperature (+15°C to +25°C).
- Unless otherwise specified, all mixing steps are listed as 'mix thoroughly' and indicate that mixing should be performed by vortexing for at least 10 seconds.
- If liquid has collected in a tube's cap after mixing, gently tap or briefly centrifuge the sample to collect the liquid into the tube's bottom, ensuring that the mixture remains homogeneous before progressing to the next step.

References

- Thermocycler Manual
- Qubit Fluorometer Manual
- Qubit dsDNA HS Assay Kit Guide
- Qubit RNA HS Assay Kit Guide



Terminology

Sample Library: The initial shotgun library generated from RNA by fragmentation, 1st and 2nd strand synthesis, A-tailing and adapter ligation.

Amplified Sample Library: The sample library after amplification by PCR.

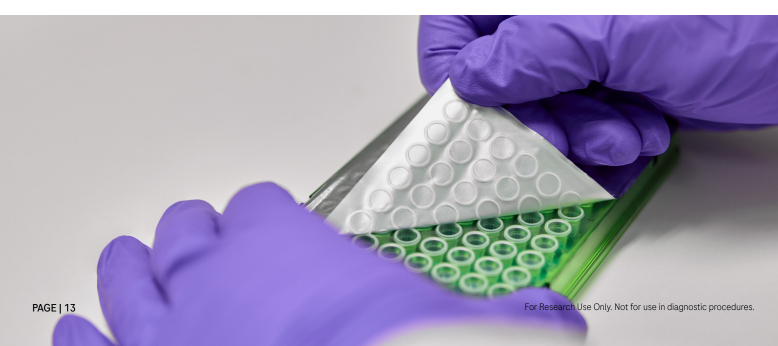
KAPA UDI Adapter: KAPA Unique Dual-Indexed Adapter are full-length, ready-to-use, QC-tested, dual-indexed Illumina adapters for ligation-based library construction.

KAPA UDI Primer Mixes: KAPA Unique Dual-Indexed Primer Mixes to be used in conjunction with KAPA Universal Adapters.

KAPA Universal Adapter: Truncated adapter containing a subset of sequencing motifs. Used in conjunction with KAPA UDI Primer Mixes.

Prepare the Following Equipment and Reagents

- Thermocyclers should be programmed with the following:
 - mRNA Capture program (Chapter 3, Prepare the Sample Library, Step 1)
 - mRNA, Fragmentation and Priming program (Chapter 3, Prepare the Sample Library, Step 2)
 - 1st Strand Synthesis program (Chapter 3, Prepare the Sample Library, Step 3)
 - 2nd Strand Synthesis and A-tailing program (*Chapter 3, Prepare the Sample Library, Step 4*)
 - Adapter Ligation program (Chapter 3, Prepare the Sample Library, Step 5)
 - Library Amplification program (Chapter 4, Amplify the Sample Library, Step 2)
 - It is recommended to use a thermocycler with a programmable heated lid. If not possible, please use the default settings.
- The following steps should be taken before beginning the workflow:
 - Verify you are using the most up-to-date version of these Instructions for Use, go to **sequencing.roche.com/support**





Required Equipment, Labware & Consumables

You assume full responsibility when using the equipment, labware, and consumables described below. This protocol is designed for use with the specified equipment, labware, and consumables.

Laboratory Equipment

Equipment	Supplier
DynaMag-96 Side Magnet	Thermo Fisher
Microcentrifuge (16,000 x g capability)	Multiple Vendors
Qubit Fluorometer	ThermoFisher
Electrophoretic device & associated assays and reagents	Multiple Vendors
Thermocycler	Multiple Vendors
Vortex mixer	Multiple Vendors
Plate Centrifuge (minimum 280 x g capability)	Multiple Vendors

Consumables Available from Roche

Description	Package Size	Material Number
KAPA Library Quantification Kit for Illumina platforms	Various options	Various material numbers
	5 mL	07983271001
KAPA Pure Beads	30 mL	07983280001
	60 mL	07983298001
KAPA PEG/NaCl Solution	20 mL	07961928001
KAPA Unique Dual-Indexed Adapter Kit	96 x 20 μL	08861919702
KAPA Library Amplification Primer Mix	250 reactions (1.25 mL)	07958994001
KAPA Library Amplification Primer Mix	384 reactions (1.92 mL)	09420410001
KAPA Library Amplification Primer Mix 96-well plate	96 x 5 μL	09420479001
WARALL .	96 reactions	09063781001
KAPA Universal Adapter	384 reactions*	09063790001
KAPA UDI Primer Mixes 1 - 96	96 reactions	09134336001
KAPA UDI Primer Mixes 97 - 192	96 reactions	09329838001
KAPA UDI Primer Mixes 193 - 288	96 reactions	09329846001
KAPA UDI Primer Mixes 289 - 384	96 reactions	09329854001

^{*} Virtual kits - consist of 4 x 96 reaction kits



Consumables Purchased from Other Vendors

Component	Supplier
10 mM Tris-HCl, pH 8.0 – 8.5	Multiple Vendors
Ethanol, 200 proof (absolute), for molecular biology	Multiple Vendors
Qubit dsDNA HS Assay Kit & Qubit RNA HS Assay Kit	ThermoFisher
Qubit Assay Tubes	ThermoFisher
Low binding Tubes: 0.2 mL PCR tubes 1.5 mL microcentrifuge tubes (optional)	Multiple Vendors
Nuclease-free, PCR-grade water	Multiple Vendors



Chapter 2.

Store and Prepare the Reagents





Step 1. Store the Reagent Kits

Reagent Kit	Storage Temperature
KAPA RNA HyperPrep Kit	-15°C to -25°C
KAPA mRNA Capture Kit*	+2°C to +8°C
KAPA Pure Beads*	+2°C to +8°C
KAPA UDI Adapter Kit	-15°C to -25°C
KAPA Universal Adapter	-15°C to -25°C
KAPA UDI Primer Mixes or KAPA UDI Primer Mixes (resuspended)	+2°C to +8°C or -15°C to -25°C



^{*}The KAPA mRNA Capture Kit and KAPA Pure Beads must not be frozen.

Step 2. Prepare KAPA Adapters and KAPA UDI Primer Mixes



For multiplexing guidelines, please refer to either the KAPA UDI Adapter or KAPA UDI Primer Mixes Instructions for Use (available online on eLabDoc) as applicable.

Step 2a. Dilute the KAPA Unique Dual-Indexed Adapters (if required)

- 1. Retrieve the KAPA UDI Adapter plate from storage (-15°C to -25°C) and thaw at room temperature.
- 2. Centrifuge the KAPA UDI Adapter plate at room temperature (280 x g for at least 1 min) to ensure the liquid is collected at the bottom of the wells.



Do not vortex the adapter plate as it could result in cross-contamination of the KAPA UDI Adapters. Pipette-mix individual adapters prior to use.

- 3. Before removing the foil cover, please ensure the plate is in the correct orientation. Please refer to the KAPA UDI Adapter Technical Data Sheet (Document number KR1736, available *online* on eLabDoc) for additional handling instructions.
- 4. Upon first use, carefully remove the foil cover of the plate to avoid cross contamination.



Discard the original foil cover. Do not reuse.



Maintain good lab practices when removing the foil seal. E.g., refrain from contact with the underside of the seal. If contact is made, change gloves before handling the adapter or sample plates.



A new pipette tip must be used for each well to avoid cross contamination. If you are not using the entire contents of the KAPA UDI Adapter plate at this time, apply a new adhesive foil seal provided in the kit. Make sure that the foil is properly aligned and fully covers all 96 wells. Use a roller or other appropriate tool to ensure that the foil is evenly applied.

5. If applicable, dilute adapters to the required concentration, see Table 1 below, using the KAPA Adapter Dilution Buffer.



Step 2b. Dilute the KAPA Universal Adapters (if required)

- 1. Retrieve the KAPA Universal Adapters tube from storage (-15°C to -25°C) and thaw at room temperature or on ice.
- 2. Mix thoroughly before first use.
- 3. Centrifuge the KAPA Universal Adapter at room temperature (280 x g for at least 1 min) to ensure the liquid is collected at the bottom of the tube.
- 4. If applicable, dilute KAPA Universal Adapters to the required concentration, see Table 1 below, using 10 mM Tris-HCl pH 8.0 8.5.

Table 1. Recommended adapter concentrations for libraries constructed from different RNA sample quantities

Quantity of starting material	Adapter stock concentration *
50 - 499 ng	1.5 µM
500 - 1000 ng	7 μM

^{*}Concentration required when using both KAPA UDI Adapter (full length) and KAPA Universal Adapter (truncated). Both KAPA UDI Adapter and KAPA Universal Adapter are provided at 15 µM.

Step 2c. Preparation of the KAPA UDI Primer Mixes

Before use of the KAPA UDI Primer Mixes, undertake the following steps to resuspend the primers:

- 1. Retrieve the KAPA UDI Primer Mixes plate from storage (+2°C to +8°C).
- 2. Centrifuge the KAPA UDI Primer Mixes plate at 280 x g for 1 minute to ensure the contents are at the bottom of the wells.
- 3. Before removing the foil cover, please ensure the plate is in the correct orientation before proceeding. In order to have well position A1 on the top left corner, the notched corner must be facing the user on the bottom left, as shown in *Figure 2*.
- 4. Carefully remove the foil cover on the plate ensuring to avoid cross contamination. Discard the original foil cover.
- Using a multichannel pipette, add 10 µL of Nuclease-free, PCR-grade water directly to the bottom of each well and discard tips after dispensing.



A new pipette tip must be used for each well to avoid cross contamination. Be sure to dispense water slowly to the bottom of each well to avoid liquid splash over to adjacent wells.

6. Ensure every well contains 10 μ L of Nuclease-free, PCR-grade water and cover the plate with one of the adhesive foil seals provided in the kit.



Make sure the foil seal is properly aligned and fully covers all 96 wells. Failure to do so can lead to cross contamination of the KAPA UDI Primer Mixes.

- 7. Use a roller or appropriate tool to ensure the foil seal is evenly applied.
- 8. Centrifuge the plate at 280 x g for 30 seconds to ensure the dispensed 10 μ L are at the bottom of the well.
- 9. Thoroughly vortex the plate ensuring all wells are mixed well.



Ensure wells at the corners of the plate are mixed well by vortexing the corners of the plate. Keep the plate upright.

10. Centrifuge the plate at 280 x g for 1 minute to ensure the contents are collected at the bottom of the wells.



- 11. The KAPA UDI Primer Mixes plate is now ready for use in the *Library Amplification* step.
- 12. Store any unused but already resuspended KAPA UDI Primer Mixes at -15°C to -25°C. To avoid repeated freeze/ thaw cycles you may transfer the resuspended primers to separate tubes or tube strips for storage

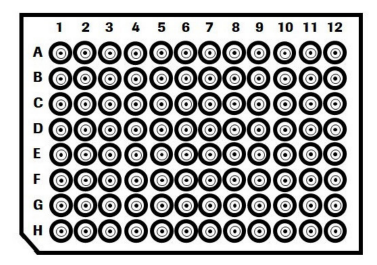


Figure 2: KAPA UDI Primer Mixes plate layout.

Considerations for Adapter Design and Concentration

- Adapter concentration affects ligation efficiency, as well as adapter and adapter-dimer carry-over during the post-ligation cleanup. The optimal adapter concentration for your workflow represents a compromise between the above factors and cost.
- When optimizing workflows for RNA inputs ≤25 ng, two or three adapter concentrations should be evaluated: try the recommended adapter concentration, as well as one or two additional concentrations in a range that is 2 - 10 times higher than the recommended concentration.
- Adapter quality has an impact on the effective concentration of adapter available for ligation. Always source the highest quality adapters from a reliable supplier, dilute and store adapters in a buffered solution with the requisite ionic strength, and avoid excessive freezing and thawing of adapter stock solutions.
- To accommodate different adapter concentrations within a batch of samples processed together, it is best to vary the concentrations of adapter stock solutions, and dispense a fixed volume (5 μL) of each adapter. The alternative (using a single stock solution, and dispensing variable volumes of adapter into ligation reactions) is not recommended.



Chapter 3.

Prepare the Sample Library





Product Description

The KAPA mRNA HyperPrep Kit for Illumina sequencing contains all of the buffers and enzymes required for poly(A) mRNA capture and the rapid construction of stranded mRNA-Seq libraries from 50 ng - 1 µg of intact total RNA via the following steps:

- 1. mRNA capture using magnetic oligo-dT beads;
- 2. fragmentation using heat and magnesium;
- 3. 1st strand cDNA synthesis using random priming;
- 4. combined 2nd strand synthesis and A-tailing, which converts the cDNA:RNA hybrid to double-stranded cDNA (dscDNA), incorporates dUTP into the second cDNA strand for stranded RNA sequencing, and adds dAMP to the 3' ends of the resulting dscDNA;
- 5. adapter ligation, where dsDNA adapters with 3' dTMP overhangs are ligated to library insert fragments; and
- 6. library amplification, to amplify library fragments carrying appropriate adapter sequences at both ends using high-fidelity, low-bias PCR. The strand marked with dUTP is not amplified, allowing strand-specific sequencing.

The kit provides KAPA Pure Beads for reaction cleanups, but does not include RNA or adapters. KAPA Adapters and KAPA UDI Primer Mixes are sold separately.

Reaction buffers are supplied in convenient formats comprising all of the required reaction components. This is designed to minimize the risk of RNase contamination, maximize homogeneous reaction composition and improve uniformity among replicate samples. Similarly, a single enzyme mixture is provided for each step of the library construction process, reducing the number of pipetting steps.

In order to maximize sequence coverage uniformity and to maintain relative transcript abundance, it is critical that library amplification bias be kept to a minimum. KAPA HiFi DNA Polymerase has been designed for low-bias, high-fidelity PCR, and is the polymerase of choice for NGS library amplification 1.2.3.4. The KAPA mRNA HyperPrep Kit includes KAPA HiFi HotStart ReadyMix (2X) and Library Amplification Primer Mix (10X) for library amplification.

- 1. Oyola, S.O., et al., BMC Genomics 13, 1 (2012).
- 2. Quail, M.A., et al., Nature Methods 9, 10 11 (2012).
- 3. Quail, M.A., et al., BMC Genomics 13, 341 (2012).
- 4. Ross, M.G., et al., Genome Biology 14, R51 (2013).

The workflow requires the use of components from the following kits:

- KAPA mRNA HyperPrep Kit
- KAPA Unique Dual-Indexed Adapter Kit & KAPA Library Amplification Primers OR*
- KAPA Universal Adapter** & KAPA UDI Primer Mixes

Ensure that the following are available:

- Nuclease-free, PCR-grade water
- Freshly-prepared 80% ethanol
- 10 mM Tris-HCl, pH 8.0 8.5

*KAPA Unique Dual-Indexed (UDI) Adapters are recommended for use with the KAPA mRNA HyperPrep Kit. However, the kit is also compatible with KAPA Universal Adapter & KAPA UDI Primer Mixes as well as other full-length or truncated adapter designs from 3rd party suppliers.

**The KAPA Universal UMI Adapter (Catalog# 09329862001, and provided at 33 µM), may also be considered for this workflow. The KAPA Universal UMI Adapter with Unique Molecular Identifiers (UMI) allows for molecular barcoding within each sample, enabling proper molecule counting for low-frequency variant detection. However, the KAPA Universal UMI Adapter has only been validated within specific KAPA Target Enrichment workflows and may not confer the same computational benefits for non-target enrichment workflows (such as whole-transcriptome sequencing). Contact your local Roche Technical Support at sequencing.roche.com/support for guidance and important considerations.



Sample Requirements

- The protocol has been validated for library construction from 50 ng − 1 µg of purified, intact total RNA in ≤50 µL of RNase-free water. The quantity of mRNA in a total RNA preparation can vary significantly between samples. An input of 50 ng − 1 µg of total RNA is recommended to ensure that sufficient mRNA is available for downstream library preparation.
- To minimize 3'->5' bias, ensure that RNA is intact and of high quality. The use of fragmented RNA will result in strong bias towards the 3'-end of the mRNA. To determine the quality of RNA, the sample may be analyzed using an Agilent Bioanalyzer RNA kit. RNA with a RNA Integrity Number (RIN) less than 7 is not recommended for this protocol.
- RNA in volumes >50 µL should be concentrated to 50 µL prior to use by either ethanol precipitation, bead purification (e.g., KAPA Pure Beads or RNAClean XP beads, Beckman Coulter), or column-based methods (e.g., RNeasy MinElute Cleanup Kit, QIAGEN). Note that some loss of material is inevitable when using any of the above methods to concentrate RNA.
- When concentrating RNA, elute in 55 μL of RNase-free water to ensure that 50 μL is available for use with this protocol. For guidance on lower input amounts or sample quality, please see contact sequencing.roche.com/support.

RNA Handling

- RNases are ubiquitous and special care should be taken throughout the procedure to avoid RNase contamination.
- To avoid airborne RNase contamination, keep all reagents and RNA samples closed when not in use.
- Use a laminar flow hood if available, or prepare a sterile and RNase-free area. Clean the workspace, pipettes and other equipment with an RNase removal product (e.g., RNaseZAP, Ambion Inc.) according to manufacturer's recommendations.
- To avoid RNase contamination, always wear gloves when handling reagents, and use certified RNase-free plastic consumables. Change gloves after making contact with equipment or surfaces outside of the RNase-free working area.
- To mix samples containing RNA, gently pipette the reaction mixture several times. Vortexing may fragment the RNA, resulting in lower quantity and a reduced library insert size.
- To avoid degradation, minimize the number of freeze-thaw cycles and always store RNA in RNase-free water.

Important Considerations when Handling mRNA Capture Beads

- Beads and bead buffers must be stored at +2°C to +8°C.
- Before use, mRNA Capture Beads must be washed and resuspended in mRNA Bead Binding Buffer.
- When preparing multiple libraries, beads may be washed in batches. A single 1.5 mL microtube can accommodate beads for up to 24 libraries. If more than 24 libraries must be prepared, wash the beads in multiple batches.
- When washing a large volume of beads, allow sufficient time for all the beads to collect on the magnet before removing the supernatant.
- Beads will settle gradually; ensure that they are fully resuspended before use.
- mRNA Capture Beads and mRNA Bead Binding Buffer contain detergent. High speed vortexing and vigorous shaking should be avoided to prevent excessive foaming. Beads may be resuspended by:
 - vortexing at low to medium speed;
 - gentle pipetting, taking care not to aspirate air; or
 - slow mixing on a tube/bottle roller.
- Before adding the Fragment, Prime and Elute Buffer (2X) to the beads, ensure that all of the mRNA Bead Wash Buffer has been removed. Carryover of mRNA Bead Wash Buffer may inhibit 1st strand cDNA synthesis.



Important Considerations for RNA Fragmentation

■ RNA is fragmented by incubating at a high temperature in the presence of magnesium before carrying out 1st strand cDNA synthesis.



After RNA fragmentation, immediately place the heat-treated sample on the magnet and remove the supernatant as soon as the liquid has cleared. Failure to do so may result in rebinding of polyadenylated regions of RNA to the capture beads, resulting in a loss of transcript coverage.

- Fragmentation conditions given in *Chapter 3* should be used as a guideline. It is recommended that a non-precious, representative sample of RNA be evaluated for the optimal fragmentation conditions.
- For fragmentation optimization beyond what is provided in *Chapter 3*, please refer to *Appendix A: Library Size Distribution Optimization*.



Step 1. mRNA Capture

This protocol requires 50 ng – 1 μ g of intact total RNA, in 50 μ L of RNase-free water. Degraded or fragmented total RNA will result in significant 3'-bias.



This protocol has been optimized to isolate mature mRNA from total RNA through two consecutive capture steps using mRNA Capture Beads. Other RNA molecules with homopolymeric adenosine regions may also be isolated.



RNA samples should only be kept on ice where specified in this protocol, since low temperatures may promote non-specific capture, resulting in increased rRNA levels in the captured mRNA.



Before starting, equilibrate mRNA Capture Beads, mRNA Bead Binding Buffer, mRNA Bead Wash Buffer and Fragment, Prime and Elute Buffer (2X) to room temperature.



Before use, beads must be washed with mRNA Bead Binding Buffer (steps 1 – 5).

1. Resuspend the mRNA Capture Beads thoroughly by pipetting up and down gently, or by using a vortex mixer on a low speed setting to prevent foaming.



High-speed vortexing or shaking should be avoided to prevent foaming. Refer to *Important Considerations when handling mRNA Capture Beads* for more information.

- 2. For each library to be prepared, transfer 52.5 μL (50 μL + 5% excess) of the resuspended mRNA Capture Beads into an appropriate tube. When preparing multiple libraries, beads for up to 24 libraries (1260 μL) may be washed in a single tube. When preparing more than 24 libraries, wash beads in multiple batches.
- 3. Place the tube on a magnet and incubate at room temperature until the solution is clear. Carefully remove and discard the supernatant, and replace it with an equal volume of mRNA Bead Binding Buffer (52.5 µL per library).
- 4. Remove the tube from the magnet and resuspend the beads by pipetting up and down, or by low speed vortexing. **Be careful to avoid producing excessive foam.**
- 5. Place the tube on the magnet and incubate at room temperature until the solution is clear. Carefully remove and discard the supernatant, and replace it with an equal volume of mRNA Bead Binding Buffer (52.5 µL per library).
- 6. Remove the tube from the magnet and resuspend the beads by pipetting up and down, or by low speed vortexing. **Be careful to avoid producing excessive foam.**
- 7. For each RNA sample to be captured, transfer $50 \,\mu\text{L}$ of resuspended mRNA Capture Beads into individual tubes or wells of a plate.
- 8. To each well/tube, add 50 μL of the appropriate RNA sample (in RNase-free water).
- 9. Mix thoroughly by gently pipetting up and down several times.
- 10. Place the plate/tube(s) in a thermocycler and perform the 1st mRNA capture as follows:

Step	Temperature	Duration
1st mRNA capture	+65°C	2 min
Cool	+20°C	5 min

- 11. Place the plate/tube(s) containing the mixture of mRNA Capture Beads and RNA, on a magnet and incubate at room temperature until the solution is clear. Remove and discard the supernatant.
- 12. Remove the plate/tube(s) from the magnet and resuspend in 200 µL of mRNA Bead Wash Buffer by pipetting up and down several times.
- 13. Place the plate/tube(s) on the magnet and incubate at room temperature until the solution is clear. Remove and discard the supernatant.
- 14. Resuspend the beads in 50 µL of RNase-free water.
- 15. Place the plate/tube(s) in a thermocycler and perform the 2nd mRNA capture as follows:



Step	Temperature	Duration
2nd mRNA capture	+70°C	2 min
Cool	+20°C	5 min

- 16. Add 50 μ L of Bead Binding Buffer to the mixture of mRNA Capture Beads and RNA, and mix thoroughly by gently pipetting up and down several times.
- 17. Incubate the plate/tube(s) at +20°C for 5 min.
- 18. Place the plate/tube(s) on the magnet and incubate at room temperature until the solution is clear. Remove and discard the supernatant.
- 19. Remove the beads from the magnet and resuspend in 200 μL of mRNA Bead Wash Buffer by pipetting up and down several times.
- 20. Place the plate/tube(s) on the magnet and incubate at room temperature until the solution is clear. Remove and discard the entire volume of supernatant.



Carryover of mRNA Bead Wash Buffer may inhibit 1st strand cDNA synthesis.

Step 2. mRNA Elution, Fragmentation and Priming

Captured mRNA is resuspended in Fragment, Prime and Elute Buffer (1X) and fragmented off the capture beads to the desired size by incubation at high temperature.

1. Prepare the required volume of Fragment, Prime and Elute Buffer (1X) at room temperature as per the table below:

Component	Volume Per Individual Sample
Fragment, Prime and Elute Buffer (2X)	11 μL
RNase-free water	11 μL
Total volume	22 μL

- 2. Thoroughly resuspend (by gentle pipette-mixing, not vortexing), the mRNA Capture Beads with captured mRNA prepared previously in 22 µL of Fragment, Prime and Elute Buffer (1X).
 - Safe stopping point Resuspended beads with captured mRNA may be stored at +2°C to +8°C for ≤24 hrs. Do not freeze the samples as this will damage the beads. When ready, proceed to Step 3.
- 3. Place the plate/tube(s) in a thermocycler and carry out the fragmentation and priming program given in the table below:



Ensure that the thermocycler lid temperature is set to +105°C (default).

Desired mean library insert size (bp)	Fragmentation
100 - 200	8 min at +94°C
200 - 300	6 min at +94°C
300 - 400	6 min at +85°C

4. Immediately place the plate/tube(s) on a magnet to capture the beads, and incubate until the liquid is clear.



To prevent hybridization of poly(A)-rich RNA to the capture beads, do not allow the sample to cool before placing on the magnet.

- 5. Carefully remove 20 µL of the supernatant containing the eluted, fragmented, and primed RNA into a separate plate or tube.
- 6. Place the plate/tube(s) on ice and proceed immediately to Step 3.1st Strand Synthesis.



Step 3. 1st Strand Synthesis

1. On ice, assemble the 1st strand synthesis master mix as per the table below:

1st Strand Synthesis Master Mix

Component	Volume Per Individual Sample inc. 20% excess
1st Strand Synthesis Buffer eluted from beads	11 μL
KAPA Script	1 μL
Total 1st strand synthesis master mix volume	12 μL

2. Keeping the plate/tube(s) on ice, combine the input RNA (fragmented & primed) and 1st strand synthesis master mix as per the table below:

Component	Volume Per Individual Sample
Fragmented, primed RNA	20 μL
1st strand synthesis master mix	10 μL
Total volume	30 μL

3. Keeping the plate/tube(s) on ice, mix thoroughly by gently pipetting the reaction up and down several times.



Ensure that the thermocycler lid temperature is set to +85°C.

4. Incubate the plate/tube(s) using the following protocol:

Step	Temperature	Duration
Primer extension	+25°C	10 min
1st strand synthesis	+42°C	15 min
Enzyme inactivation	+70°C	15 min
Hold	+4°C	∞

5. Place the plate/tube(s) on ice and proceed immediately to Step 4. 2nd Strand Synthesis & A-tailing.



Step 4. 2nd Strand Synthesis & A-tailing

1. On ice, assemble the 2nd strand synthesis and A-tailing master mix as per table below:

2nd Strand Synthesis & A-tailing Master Mix

Component	Volume Per Individual Sample inc. 10% excess
2nd Strand Marking Buffer	31 μL
2nd Strand Synthesis & A-Tailing Enzyme Mix	2 µL
Total 2nd strand synthesis and A-tailing master mix volume	33 µL

2. Keeping the plate/tube(s) on ice, combine the 1st strand synthesis product and 2nd strand synthesis and A-tailing master mix as per the table below:

Component	Volume Per Individual Sample
1st strand synthesis product	30 μL
2nd strand synthesis & A-tailing master mix	30 μL
Total volume	60 μL

3. Keeping the plate/tube(s) on ice, mix thoroughly by gently pipetting the reaction up and down several times.



Ensure that the thermocycler lid temperature is set to +85°C.

4. Incubate the plate/tube(s) using the following protocol:

Step	Temperature	Duration
2nd strand synthesis	+16°C	30 min
A-tailing	+62°C	10 min
Hold	+4°C	∞

5. Place the plate/tube(s) on ice and proceed immediately to Step 5. Adapter Ligation.



Step 5. Adapter Ligation



KAPA UDI Adapter and KAPA Universal Adapter, unless specified, will hereafter be referred to as KAPA Adapters.



KAPA Adapters must be added to each tube/well individually prior to addition of the Ligation reagents.



If using KAPA Universal Adapter, sample indexes are incorporated during the Library Amplification step (*Chapter 4, Amplify the Sample Library*). Precautions should be taken to avoid sample cross contamination.

1. On ice, assemble each Adapter Ligation reaction as per table below:

Adapter Ligation Master Mix

Component	Volume Per Individual Sample inc. 10% excess
Ligation Buffer	40 μL
DNA Ligase	10 μL
Total Adapter Ligation master mix volume	50 μL

2. Keeping the plate/tube(s) on ice, combine the 2nd strand synthesis product and diluted KAPA Adapter stock, then add the Adapter Ligation master mix as per the table below:

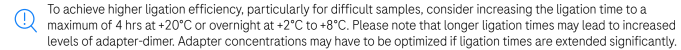
Component	Volume Per Individual Sample
2nd strand synthesis product	60 μL
Adapter ligation master mix	45 μL
Diluted adapter stock (Chapter 2)	5 μL
Total volume	110 μL

3. Keeping the plate/tube(s) on ice, mix thoroughly by pipetting the reaction up and down several times.



Ensure that the thermocycler lid temperature is set to +50°C.

- 4. Incubate the plate/tube(s) at +20°C for 15 min.
- 5. Proceed immediately to the next step.



6. Proceed immediately to Step 6. Purify the Sample Library using KAPA Pure Beads



Step 6. Purify the Sample Library using KAPA Pure Beads

If KAPA UDI Adapters were used for library construction – proceed to Step 6a or,

If KAPA Universal Adapter were used for library construction – proceed to Step 6b

Step 6a. Purify the Sample Library constructed using KAPA UDI Adapter (first cleanup)

 \triangle

Only perform this 0.63X cleanup if full length, KAPA UDI Adapters were used. Proceed to *Step 6b* if the KAPA Universal Adapter was used in order to perform a single 0.7X post-ligation cleanup.

1. To each ligation reaction, add 70 μ L of room temperature KAPA Pure Beads that have been thoroughly resuspended as per the table below:

Component	Volume Per Individual Sample
Adapter-ligated DNA	110 µL
KAPA Pure Beads	70 μL
Total volume	180 μL

2. Once added, mix thoroughly and centrifuge briefly to collect all droplets. Do NOT allow beads to pellet.



It is important at this step to ensure that the solution is thoroughly mixed and appears homogeneous. Insufficient mixing may compromise recovery and result in size selection.

- 3. Incubate the sample at room temperature for 5 min to bind DNA to the beads.
- 4. Place the sample on a magnet to capture the beads. Incubate until the liquid is clear.
- 5. Carefully remove and discard the supernatant.
- 6. Keeping the sample on the magnet, add 200 µL of 80% ethanol.
- 7. Incubate the sample on the magnet at room temperature for ≥30 sec.
- 8. Carefully remove and discard the ethanol.
- 9. Keeping the sample on the magnet, add 200 µL of 80% ethanol.
- 10. Incubate the sample on the magnet at room temperature for ≥30 sec.
- 11. Carefully remove and discard the ethanol. Try to remove all residual ethanol without disturbing the beads.
- 12. Dry the beads at room temperature for 3 5 min, or until all of the ethanol has evaporated.



Over-drying the beads may result in dramatic yield loss. Over-drying is indicated by a dry, cracked appearance of the bead pellet. The bead pellet should have a matte appearance when sufficiently dried.

- 13. Remove the sample from the magnet.
- 14. Thoroughly resuspend the beads in 50 μL of elution buffer (10 mM Tris-HCl, pH 8.0 8.5) to proceed with a 2nd post-ligation cleanup using PEG/NaCl (Step 7. Purify the Sample Library using PEG/NaCl).
- 15. Incubate the sample at room temperature for 2 minutes to allow the sample library to elute off the beads.
- (1)

Safe Stopping Point - If necessary this is a safe stopping point. The solution with resuspended beads can be stored at +2°C to +8°C for \leq 24 hrs. Do not freeze the beads, as this can result in dramatic loss of DNA. When ready, proceed to Step 7. Purify the Sample Library using PEG/NaCl.



Step 6b. Purify the Sample Library constructed using KAPA Universal Adapter

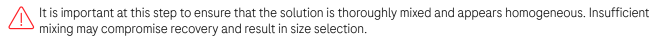


Only perform this 0.7X cleanup if KAPA Universal Adapters were used. Refer to *Step 6a* if the KAPA UDI Adapters were used in order to perform a 0.63X post-ligation cleanup.

1. To each ligation reaction, add 77 μ L of room temperature KAPA Pure Beads that have been thoroughly resuspended as per the table below:

Component	Volume Per Individual Sample
Adapter-ligated DNA	110 µL
KAPA Pure Beads	77 μL
Total volume	187 µL

2. Once added, mix thoroughly and centrifuge briefly to collect all droplets. Do NOT allow beads to pellet.



- 3. Incubate the sample at room temperature for 5 min to bind DNA to the beads.
- 4. Place the sample on a magnet to capture the beads. Incubate until the liquid is clear.
- 5. Carefully remove and discard the supernatant.
- 6. Keeping the sample on the magnet, add 200 µL of 80% ethanol.
- 7. Incubate the sample on the magnet at room temperature for ≥30 sec.
- 8. Carefully remove and discard the ethanol.
- 9. Keeping the sample on the magnet, add 200 µL of 80% ethanol.
- 10. Incubate the sample on the magnet at room temperature for ≥30 sec.
- 11. Carefully remove and discard the ethanol. Try to remove all residual ethanol without disturbing the beads.
- 12. Dry the beads at room temperature for 3-5 min, or until all of the ethanol has evaporated.
- Over-drying the beads may result in dramatic yield loss. Over-drying is indicated by a dry, cracked appearance of the bead pellet. The bead pellet should have a matte appearance when sufficiently dried.
- 13. Remove the sample from the magnet.
- 14. Thoroughly resuspend the beads in 22 µL of 10 mM Tris-HCl, pH 8.0 8.5.
- 15. Incubate the sample at room temperature for 2 minutes to allow the sample library to elute off the beads.
- 16. Place the sample on a magnet to capture the beads. Incubate until the liquid is clear.
- 17. Transfer 20 µL of the clear supernatant to a new plate/tube(s) and proceed to *Library Amplification (Chapter 4, Step 1*).
 - Safe Stopping Point If necessary, this is a safe stopping point. Purified, adapter-ligated library may be stored at +2°C to +8°C for 1 2 weeks or at -15°C to -25°C for ≤1 month before amplification. To avoid degradation, always store DNA in a buffered solution (10 mM Tris-HCl, pH 8.0 8.5) when possible, and minimize the number of freezethaw cycles.



Step 7. Purify the Sample Library using PEG/NaCl (second cleanup)



This is a **mandatory second cleanup** if KAPA UDI Adapters were used during library construction. Only perform this 0.7X cleanup if KAPA UDI Adapters were used. **SKIP** this step if KAPA Universal Adapters were used.

1. To each sample, add $35 \,\mu\text{L}$ of room temperature PEG/NaCl solution that have been thoroughly resuspended as per the table below:

Component	Volume Per Individual Sample
Beads with purified, adapter-ligated DNA (from Step 6a)	50 μL
PEG/NaCl Solution	35 μL
Total volume	85 μL

- 2. Once added, mix thoroughly and centrifuge briefly to collect all droplets. Do NOT allow beads to pellet.
- 3. Incubate the sample at room temperature for 5 min to bind DNA to the beads.
- 4. Place the sample on a magnet to capture the beads. Incubate until the liquid is clear.
- 5. Carefully remove and discard 80 µL of supernatant.
- 6. Keeping the sample on the magnet, add 200 μL of 80% ethanol.
- 7. Incubate the sample on the magnet at room temperature for ≥30 sec.
- 8. Carefully remove and discard the ethanol.
- 9. Keeping the sampleon the magnet, add 200 µL of 80% ethanol.
- 10. Incubate the sample on the magnet at room temperature for ≥30 sec.
- 11. Carefully remove and discard the ethanol. Try to remove all residual ethanol without disturbing the beads.
- 12. Dry the beads at room temperature for 3 5 min, or until all of the ethanol has evaporated.



Over-drying the beads may result in dramatic yield loss. Over-drying is indicated by a dry, cracked appearance of the bead pellet. The bead pellet should have a matte appearance when sufficiently dried.

- 13. Remove the sample from the magnet.
- 14. Thoroughly resuspend the beads in 22 µL of 10 mM Tris-HCl (pH 8.0 8.5).
- 15. Incubate the sample at room temperature for 2 min to allow the sample library to elute off the beads.
- 16. Place the sample on a magnet to capture the beads. Incubate until the liquid is clear.
- 17. Transfer 20 µL of the clear supernatant to a new plate/tube(s) and proceed to *Library Amplification (Chapter 4, Step 1)*.



Safe Stopping Point - If necessary, this is a safe stopping point. Purified, adapter-ligated library may be stored at +2°C to +8°C for 1 - 2 weeks or at -15°C to -25°C for ≤1 month before amplification. To avoid degradation, always store DNA in a buffered solution (10 mM Tris-HCl, pH 8.0 - 8.5) when possible, and minimize the number of freeze-thaw cycles.



Chapter 4.

Amplify the Sample Library





This chapter describes how to amplify the adapter-ligated library, using KAPA HiFi HotStart ReadyMix and appropriate compatible indexes or primer sets.



If KAPA UDI Adapters were used for adapter ligation, ensure KAPA Library Amplification Primer Mix (provided with the kit) is used for library amplification.



If KAPA Universal Adapters were used for adapter ligation, ensure a unique KAPA UDI Primer Mix is added to each sample library.

Ensure that the following is available:

- Freshly-prepared 80% ethanol
- 10 mM Tris-HCl, pH 8.0 8.5
- Nuclease-free, PCR-grade water

Library Amplification Considerations

- KAPA HiFi HotStart, the enzyme provided in the KAPA HiFi HotStart ReadyMix, is an antibody-based hotstart formulation of KAPA HiFi DNA Polymerase—a B-family DNA polymerase engineered for increased processivity and high fidelity. KAPA HiFi HotStart has 5'-->3' polymerase and 3'-->5' exonuclease (proofreading) activity, but no 5'-->3' exonuclease activity. The error rate of KAPA HiFi HotStart is 2.8 x 10⁻⁷ errors/base, equivalent to 1 error in 3.5 x 10⁶ nucleotides incorporated.
- The KAPA Library Amplification Primer Mix (10X) is designed to eliminate or delay primer depletion during library amplification reactions performed with KAPA HiFi HotStart ReadyMix. The primer mix is suitable for the amplification of all Illumina libraries flanked by the P5 and P7 flow cell sequences. Primers are supplied at a 10X concentration of 20 μM each, and have been formulated as described below. User-supplied primer mixes may be used in combination with incomplete or custom adapters. Please contact Technical Support at *sequencing.roche.com/support* for guidelines on the formulation of user-supplied library amplification primers.
- To achieve the highest amplification efficiency and avoid primer depletion, it is critical to use an optimal concentration of high quality primers. Primers should be used at a final concentration of 0.5 4 μM each.
- Library amplification primers should be HPLC-purified and modified to include a phosphorothioate bond at the 3'-terminal of each primer (to prevent degradation by the strong proofreading activity of KAPA HiFi HotStart DNA Polymerase). Always store and dilute primers in a buffered solution (e.g., 10 mM Tris-HCl, pH 8.0 8.5), and limit the number of freeze-thaw cycles. To achieve the latter, store primers at +2°C to +8°C for short-term use, or as single-use aliquots at -15°C to -25°C.
- In library amplification reactions (set up according to the recommended protocol), primers are typically depleted before dNTPs. When DNA synthesis can no longer take place due to substrate depletion, subsequent rounds of DNA denaturation and annealing result in the separation of complementary DNA strands, followed by imperfect annealing to non-complementary partners. This presumably results in the formation of so-called "daisy chains" or "tangled knots", comprising large assemblies of improperly annealed, partially double-stranded, heteroduplex DNA. These species migrate slower and are observed as secondary, higher molecular weight peaks during the electrophoretic analysis of amplified libraries. However, they typically comprise library molecules of the desired length, which are individualized during denaturation prior to cluster amplification or probe hybridization. Since these heteroduplexes contain significant portions of single-stranded DNA, over-amplification leads to the under-quantification of library molecules with assays employing dsDNA-binding dyes. qPCR-based library quantifications methods, such as the KAPA Library Quantification assay, quantify DNA by denaturation and amplification, thereby providing an accurate measure of the amount of adapter-ligated molecules in a library—even if the library was over-amplified. Excessive library amplification can result in other unwanted artifacts such as amplification bias, PCR duplicates, chimeric library inserts and nucleotide substitutions. The extent of library amplification should therefore be limited as much as possible, while ensuring that sufficient material is generated for QC and downstream processing.
- If cycled to completion (not recommended), one 50 µL library amplification can produce 8 − 10 µg of amplified library. To minimize over-amplification and its associated, undesired artifacts, the number of amplification cycles should be tailored to produce the optimal amount of final library required for downstream processes. This is typically in the range of 250 ng − 1.5 µg.



Step 1. Prepare the Library Amplification Reaction

Make sure the KAPA Pure Beads are removed from storage at least 30 minutes prior to starting the workflow to ensure they are equilibrated to room temperature. For best performance, store the beads protected from light when not in use.

(2X) contains isostabilizers and may not freeze completely, even when stored at -15°C to -25°C. Nevertheless, always ensure that the ReadyMix is **fully thawed** and **thoroughly mixed** before use.

If applicable, retrieve and thaw the KAPA UDI Primer Mixes plate prepared in *Chapter 2, Step 2c.*

Centrifuge the KAPA UDI Primer Mixes plate at 280 x g for 30 seconds to collect the contents to the bottom of the wells and peel off or pierce the foil seal for the appropriate number of wells needed. If only using a subset of the KAPA UDI Primer Mixes from the original plate, remove and discard residual primers from the well and apply a new adhesive foil seal provided in the kit.

If piercing the foil seal, avoid cross contamination by using a new pipette tip for every well.

Proper re-sealing and storage of the KAPA UDI Primer Mixes plate is necessary for unused primer mixes utilization at a later date.

() Keep all components on ice as long as possible during handling.

1. Assemble each library amplification reaction as per table below:

Component	Volume per Individual Library
KAPA HiFi HotStart ReadyMix (2X)	25 μL
KAPA Library Amplification Primer Mix* OR KAPA UDI Primer Mix**	5 μL
Adapter-ligated library	20 μL
Total volume	50 μL

^{*} If KAPA UDI Adapters were used for adapter ligation, ensure KAPA Library Amplification Primer Mix is used for library amplification. The KAPA HiFi HotStart ReadyMix and KAPA Library Amplification Primer Mix should preferably be premixed and added in a single pipetting step.

2. Mix thoroughly and centrifuge briefly. Immediately proceed to the next step.

^{**} If KAPA Universal Adapters were used for adapter ligation ensure a unique KAPA UDI Primer Mix is added to each sample library.



Step 2. Perform the Library Amplification

1. Place the sample in the thermocycler and amplify the Sample library using the following Library Amplification program with the lid temperature set to +105°C:

Step	Temperature	Duration	Cycles
Initial denaturation	+98°C	45 sec	1
Denaturation	+98°C	15 sec	Variable, see Table 2 below
Annealing	+60°C	30 sec	
Extension	+72°C	30 sec	
Final extension	+72°C	1 min	1
Hold	+4°C	∞	1

Table 2: Recommended cycle numbers

Quantity of starting material	Number of cycles*
50 – 100 ng	13 - 16
101 – 250 ng	11 – 14
251 – 500 ng	9 – 12
501 - 1000 ng	7 – 10

Note: The recommended number of cycles should be used as a guide for library amplification. Cycle numbers may require adjustment depending on RNA input quality, library amplification efficiency, presence of adapter-dimer, and the desired yield post amplification. Quantification of material after the second post-ligation cleanup using a qPCR assay, such as the KAPA Library Quantification Kit, can help to determine the number of amplification cycles required for a specific sample type or application.

2. Proceed immediately to the next step.

^{*}The recommended number of cycles is irrespective of which adapter kit was used during library construction (KAPA UDI Adapter or KAPA Universal Adapter).



Step 3. Purify the Amplified Sample Library using KAPA Pure Beads

- If KAPA UDI Adapters and KAPA Library Amplification Primer Mix were used for library construction proceed to Step 3a or,
- If KAPA Universal Adapter and KAPA UDI Primer Mixes were used for library construction proceed to Step 3b.

Step 3a. Purify the Amplified Sample Library constructed using KAPA UDI Adapter & KAPA Library Amplification Primer Mix

- 1. Add 50 µL of room temperature, thoroughly resuspended, KAPA Pure Beads to each amplified sample library.
- 2. Mix the amplified sample library and KAPA Pure Beads thoroughly and centrifuge briefly to collect all droplets. Do **NOT** allow beads to pellet.
- 3. Incubate the sample at room temperature for 5 minutes to allow the amplified sample library to bind to the beads.
- 4. Place the sample on a magnet to capture the beads. Incubate until the liquid is clear.
- 5. Carefully remove and discard the supernatant.
- 6. Keeping the sample on the magnet, add 200 µL of freshly-prepared 80% ethanol.
- 7. Incubate the sample at room temperature for ≥30 seconds.
- 8. Carefully remove and discard the ethanol.
- 9. Keeping the sample on the magnet, add 200 µL of freshly-prepared 80% ethanol.
- 10. Incubate the sample at room temperature for ≥30 seconds.
- 11. Carefully remove and discard the ethanol. Remove residual ethanol without disturbing the beads.
- 12. Allow the beads to dry at room temperature, sufficiently for all of the ethanol to evaporate.
 - bead pellet. The bead pelle

Over-drying the beads may result in dramatic yield loss. Over-drying is indicated by a dry, cracked appearance of the bead pellet. The bead pellet should have a matte appearance when sufficiently dried.

- 13. Remove the sample from the magnet.
- 14. Thoroughly resuspend the beads in 22 µL (or appropriate volume) of 10 mM Tris-HCl, pH 8.0 8.5.
- 15. Incubate the sample at room temperature for 2 minutes to allow the amplified sample library to elute off the beads.
- 16. Place the sample on the magnet to capture the beads. Incubate until the liquid is clear.
- 17. Transfer the clear supernatant to a new tube(s)/well and proceed with library QC (*Chapter 5. Quality Control*), target capture or sequencing, as appropriate.
- 18. Purified, amplified sample libraries can be stored at +2°C to +8°C for 1 2 weeks or at -15 °C to -25°C for up to 3 months.



Step 3b. Purify the Amplified Sample Library constructed using KAPA Universal Adapter & KAPA UDI Primer Mixes



A modified post-amplification cleanup is highly recommended when using KAPA Universal Adapter & KAPA UDI Primer Mixes with KAPA library preparation kits. Especially if libraries will be sequencing on an Illumina NovaSeq or HiSeq X system (with patterned flow cells). This additional or modified cleanup will serve to remove any indexed primer carryover which may exacerbate index hopping.



The KAPA Pure Beads provided with the 24 rxn and 96 rxn pack size, may not be sufficient if processing 24 or 96 samples with the KAPA Universal Adapter & KAPA UDI Primer Mixes. Additional KAPA Pure Beads is sold separately. Alternatively, the unused PEG/NaCl provided with the kit may be repurposed for a "with-bead" protocol. Please contact *Technical Support* for guidance.

- 1. Add 50 µL of room temperature, thoroughly resuspended, KAPA Pure Beads to each amplified sample library.
- 2. Mix the amplified sample library and KAPA Pure Beads thoroughly and centrifuge briefly to collect all droplets. Do **NOT** allow beads to pellet.
- 3. Incubate the sample at room temperature for 5 minutes to allow the amplified sample library to bind to the beads.
- 4. Place the sample on a magnet to capture the beads. Incubate until the liquid is clear.
- 5. Carefully remove and discard the supernatant.
- 6. Remove the tubes from the magnet, and resuspend the beads in 50 μ L of nuclease-free PCR-grade water or 10 mM Tris-HCl, pH 8.0 8.5.
- 7. Add 50 µL of KAPA Pure Beads to each sample.
- 8. Mix thoroughly by pipetting or vortexing, and centrifuge briefly.
- 9. Incubate the sample at room temperature for 5 minutes to allow the amplified sample library to bind to the beads.
- 10. Place the sample on a magnet to capture the beads. Incubate until the liquid is clear.
- 11. Carefully remove and discard the supernatant.
- 12. Keeping the sample on the magnet, add 200 μ L of freshly-prepared 80% ethanol.
- 13. Incubate the sample at room temperature for ≥30 seconds.
- 14. Carefully remove and discard the ethanol.
- 15. Keeping the sample on the magnet, add 200 µL of freshly-prepared 80% ethanol.
- 16. Incubate the sample at room temperature for ≥30 seconds.
- 17. Carefully remove and discard the ethanol. Remove residual ethanol without disturbing the beads.
- 18. Allow the beads to dry at room temperature, sufficiently for all of the ethanol to evaporate.



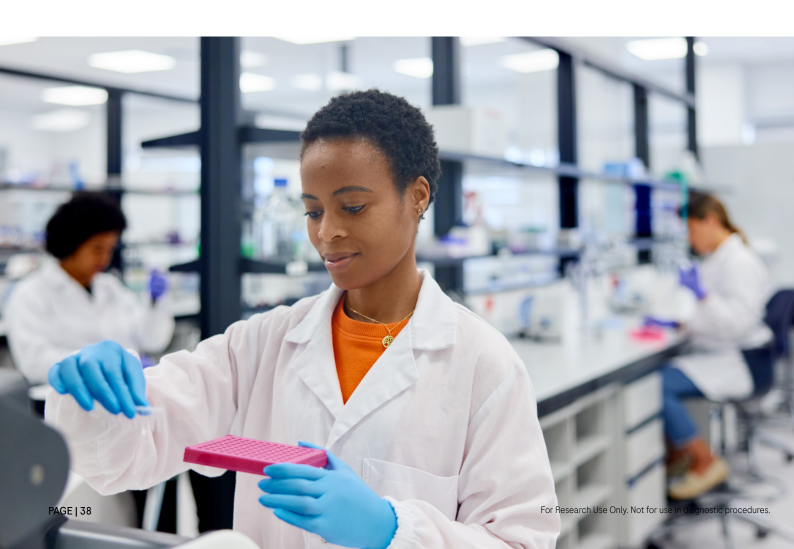
Over-drying the beads may result in dramatic yield loss. Over-drying is indicated by a dry, cracked appearance of the bead pellet. The bead pellet should have a matte appearance when sufficiently dried.

- 19. Remove the sample from the magnet.
- 20. Thoroughly resuspend the beads in 22 μ L (or appropriate volume) of 10 mM Tris-HCl, pH 8.0 8.5.
- 21. Incubate the sample at room temperature for 2 minutes to allow the sample library to elute off the beads.
- 22. Place the sample on the magnet to capture the beads. Incubate until the liquid is clear.
- 23. Transfer the clear supernatant to a new tube(s)/well and proceed with library QC (*Chapter 5. Quality Control*), target capture or sequencing, as appropriate.
- 24. Purified, amplified sample libraries can be stored at +2°C to +8°C for 1 2 weeks or at -15°C to -25°C for up to 3 months.



Chapter 5.

Quality Control





This chapter describes how to determine the concentration and size distribution of the sample library. Your specific library construction workflow should be tailored and optimized to yield a sufficient amount of adapter-ligated molecules of the desired size distribution for the next step in the process (e.g., target capture or sequencing), as well as for library QC and archiving purposes.

Quantification

KAPA Library Quantification Kits for Illumina platforms are recommended for qPCR-based quantification of libraries generated using the KAPA mRNA HyperPrep workflow. Libraries may also be quantified using a Qubit Fluorometer and the Qubit dsDNA HS Assay Kit.

Sizing

The size distribution of the final libraries should be confirmed with an electrophoretic method. A LabChip GX, GXII, or GX Touch (PerkinElmer), Bioanalyzer, TapeStation (Agilent Technologies), Fragment Analyzer (Advanced Analytical) system or similar instruments are recommended over conventional gels. KAPA mRNA HyperPrep Kit libraries may require dilution prior to electrophoretic assessment. Refer to the respective instrument and assay user manuals for guidance on how to dilute your sample library so as not to exceed the detection limit of the specific assay.



Appendices



Appendix A. Library Size Distribution Optimization

The KAPA mRNA HyperPrep Kit offers a tunable RNA fragmentation module in which RNA is fragmented at a high temperature in the presence of magnesium. Final library size distributions can be optimized for specific sample types and applications by varying both incubation time and temperature. Generally:

- higher temperatures and/or longer incubation times result in shorter, narrower distributions; and
- lower temperatures and/or shorter incubation times result in longer, broader distributions.

When using high-quality, intact total RNA, the following recommendations can be used as a starting point for the optimization of final library distributions beyond what is provided within *Chapter 3*. It is recommended that a non-precious, representative RNA sample be used for this optimization. For the following figures and tables, final libraries were generated using 100 ng of high-quality Universal Human Reference (UHR) RNA.

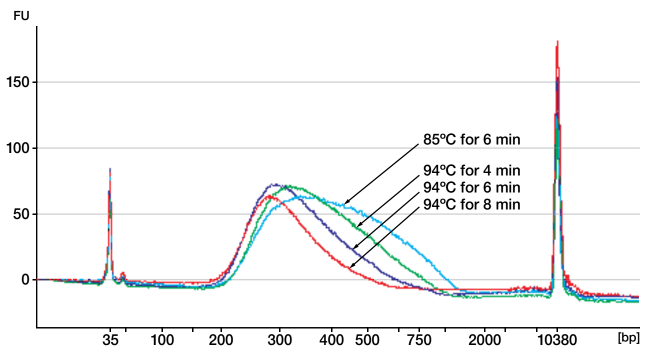


Figure 2. Final library distributions demonstrating fragmentation tunability. Higher temperatures and longer incubation times resulted in shorter, narrower final library distributions. Libraries were constructed using 100 ng of high-quality UHR RNA and various fragmentation conditions. Note that results may differ with other sample sources. Electropherograms were generated with an Agilent High Sensitivity DNA Kit.



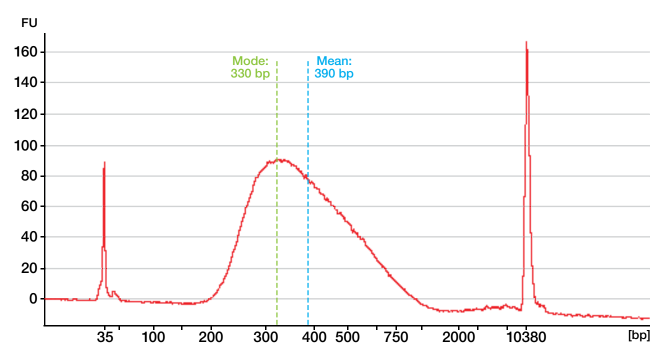


Figure 3. Visual depiction of the mean and mode distribution metrics. For a final library generated using 100 ng UHR fragmented at 94°C for 4 minutes, the mode is the highest peak in the library (~330 bp), while the mean is the numerical average across all molecular lengths in the library (~390 bp). In this example, the mean of the library is calculated across the range of 190 to 1600 bp. The higher molecular weight shoulder of the distribution results in the mean being larger than the mode.

Table 3. Approximate mean and mode final library sizes (bp) for each fragmentation condition assessed

For any substitut	Final library size (bp)		
Fragmentation	Mean	Mode	
+94°C for 8 min	~320	~290	
+94°C for 6 min	~350	~300	
+94°C for 4 min	~390	~330	
+94°C for 2 min	~470	~380	
+85°C for 6 min	~430	~3300	
+85°C for 4 min	~490	~450	



Appendix B. Limited Warranty

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