KAPA Library Amplification Kit
Illumina® Platforms

KR0408 – v7.17

This Technical Data Sheet provides product information and a detailed protocol for the KAPA Library Amplification Kits.

This document applies to KAPA Library Amplification Kits (07958978001 and 07958986001), KAPA Library Amplification Kits (PCR Mix only) (07958943001, 07958951001 and 07958960001), and KAPA Library Amplification Kits (Primers only) (07958994001).

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Quick Notes

- KAPA HiFi HotStart ReadyMix (2X) is specifically designed to minimize amplification bias, while maintaining industry-leading fidelity.
- Library Amplification Primer Mix (10X) contains primers that target the P5 and P7 regions of Illumina TruSeq® and dual-indexed adapters.
- The primer mix is formulated to limit primer depletion and over-amplification.
- Kits without primers (KK2611, KK2612) are available for library amplification with user-supplied primers.

<table>
<thead>
<tr>
<th>Kapa/Roche Kit Codes and Components</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>KK2620 07958978001 (50 reactions)</td>
<td>1.25 mL 0.25 mL</td>
</tr>
<tr>
<td>KK2621 07958960001 (250 reactions)</td>
<td>6.25 mL 1.25 mL</td>
</tr>
<tr>
<td>KK2610 07958943001 (10 reactions)</td>
<td>0.25 mL</td>
</tr>
<tr>
<td>KK2611 07958951001 (50 reactions)</td>
<td>1.25 mL</td>
</tr>
<tr>
<td>KK2612 07958960001 (250 reactions)</td>
<td>6.25 mL</td>
</tr>
<tr>
<td>KK2623 07958994001 (250 reactions)</td>
<td>1.25 mL</td>
</tr>
</tbody>
</table>
Product Description
KAPA Library Amplification Kits for Illumina platforms are designed for the amplification of next-generation sequencing libraries prepared for Illumina sequencing.

In order to maximize sequence coverage uniformity, it is critical to minimize library amplification bias. KAPA HiFi DNA Polymerase is designed for low-bias, high-fidelity PCR, and is the reagent of choice for NGS library amplification.\(^1\)\(^,\)\(^2\)\(^,\)\(^3\)\(^,\)\(^4\) KAPA Library Amplification Kits include KAPA HiFi HotStart ReadyMix (2X), a ready-to-use PCR mix comprising all the components for library amplification—except primers and template. Kits also include Library Amplification Primer Mix (10X), designed for the high-efficiency amplification of Illumina libraries flanked by adapters containing the P5 and P7 flow cell sequences.


Product Applications
KAPA Library Amplification Kits for Illumina platforms are suited for high-efficiency, high-fidelity, low-bias amplification of libraries prior to Illumina sequencing. This includes libraries prepared for:

- whole-genome shotgun sequencing
- targeted sequencing (pre- and post-capture amplification)
- amplicon sequencing
- ChIP-seq
- RNA-seq

Product Specifications
Shipping and Storage
The enzymes provided in this kit are temperature sensitive, and appropriate care should be taken during shipping and storage. KAPA Library Amplification Kits are shipped on dry ice or ice packs, depending on the destination country. Upon receipt, store the entire kit at -15°C to -25°C in a constant-temperature freezer. When stored under these conditions and handled correctly, all kit components will retain full activity until the expiry date indicated on the kit label.

Handling
Always ensure that components have been fully thawed and thoroughly mixed before use. Keep all reaction components and master mixes on ice whenever possible during handling. KAPA HiFi HotStart ReadyMix (2X) may not freeze completely, even when stored at -15°C to -25°C. Nevertheless, always ensure that the KAPA HiFi HotStart ReadyMix is fully thawed and thoroughly mixed before use.

Quality Control
All kit components are subjected to stringent functional quality control, are free of detectable contaminating exo- and endonuclease activities, and meet strict requirements with respect to DNA contamination. Please contact Technical Support at kapabiosystems.com/support for more information.
Important Parameters

Cycle Number

Excessive library amplification should be avoided to minimize the following adverse effects:

• increased duplicate reads
• uneven coverage depth and sequence dropout
• chimeric library inserts
• nucleotide substitutions
• heteroduplex formation

To minimize over-amplification and associated unwanted artifacts, the number of amplification cycles should be optimized to ensure a sufficient amount of amplified library for the next step in the workflow (capture or sequencing), plus the amount needed for library QC and/or archiving. Depending on the sequencing application and degree of multiplexing, 100 ng – 1.5 µg of amplified library is typically required.

Table 1 indicates the number of cycles typically required to generate 100 ng or 1 µg of amplified library DNA, with or without size-selection. Size selection of libraries at any part in the library construction process results in significant loss of material and as a result, 2 – 4 additional cycles are required for workflows which include a size-selection step prior to library amplification.

These guidelines are for libraries prepared with the KAPA LTP or HTP Library Preparation Kit; the actual optimal number of cycles may be higher, depending the reagents and protocol used for library construction, and the quality of the input DNA. For libraries prepared from FFPE DNA or other challenging samples, or libraries with a broad fragment size distribution, 1 – 3 additional cycles may be required.

Primer Quality and Concentration

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 – 2 µM each. For libraries constructed from ≥100 ng input DNA, the highest final concentration (2 µM of each primer) is recommended.

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 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 4°C for short-term use, or as single-use aliquots at -20°C.

Primer Depletion and Library Over-amplification

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 primer 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 are typically comprised of library molecules of the desired length, which are separated during denaturation prior to target enrichment (capture) or cluster amplification. 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 quantification methods, such as the KAPA Library Quantification assay, quantify DNA by denaturation and amplification, thereby providing a more accurate measurement of the amount of adapter-ligated molecules—even if the library was over-amplified.

Please refer to the KAPA NGS Library Preparation Technical Guide for a more detailed discussion of factors that can affect the efficiency of library amplification, and the impact of over-amplification on library quantification.

<table>
<thead>
<tr>
<th>Input DNA*</th>
<th>Cycles for 100 ng amplified library DNA</th>
<th>Cycles for 1 µg amplified library DNA</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No SS</td>
<td>With SS</td>
</tr>
<tr>
<td>1 µg</td>
<td>0</td>
<td>2 – 4</td>
</tr>
<tr>
<td>500 ng</td>
<td>0 – 1</td>
<td>3 – 5</td>
</tr>
<tr>
<td>250 ng</td>
<td>1 – 2</td>
<td>4 – 6</td>
</tr>
<tr>
<td>100 ng</td>
<td>2 – 4</td>
<td>6 – 8</td>
</tr>
<tr>
<td>50 ng</td>
<td>5 – 7</td>
<td>8 – 10</td>
</tr>
<tr>
<td>25 ng</td>
<td>6 – 8</td>
<td>9 – 11</td>
</tr>
<tr>
<td>10 ng</td>
<td>7 – 9</td>
<td>11 – 13</td>
</tr>
<tr>
<td>5 ng</td>
<td>8 – 10</td>
<td>12 – 14</td>
</tr>
<tr>
<td>1 ng</td>
<td>11 – 13</td>
<td>14 – 16</td>
</tr>
</tbody>
</table>

*Input into library construction
Optimization of Library Amplification
The quantification of adapter-ligated libraries (prior to library amplification) can greatly facilitate the optimization of library amplification parameters, particularly when a library construction workflow is first established. With the KAPA Library Quantification Kit, the amount of template DNA (adapter-ligated molecules) available for library amplification can be determined accurately. From there, the number of amplification cycles needed to achieve a specific yield of amplified library can be predicted theoretically. Please contact Technical Support at kapabiosystems.com/support regarding a calculator designed to assist with these calculations.

DNA Polymerase
KAPA HiFi HotStart, the enzyme provided in KAPA HiFi HotStart ReadyMix (2X), is an antibody-based hot start formulation of KAPA HiFi DNA Polymerase—a novel B-family DNA polymerase engineered for increased processivity and high fidelity. KAPA HiFi HotStart has 5’→3’ polymerase and 3’→5’ exonuclease (proofreading) activities, but no 5’→3’ exonuclease activity. The strong 3’→5’ exonuclease activity results in superior accuracy during DNA amplification. The error rate of KAPA HiFi HotStart DNA Polymerase is $2.8 \times 10^{-7}$ errors/base, equivalent to 1 error per $3.5 \times 10^6$ nucleotides incorporated.

Reaction Cleanups
- This protocol has been validated for use with either KAPA Pure Beads (KK8000, KK8001, KK8002), or Agencourt® AMPure® XP (Beckman Coulter®). Solutions and conditions for DNA binding and size selection may differ if other beads are used.
- Observe all the storage and handling recommendations for KAPA Pure Beads or AMPure XP. Equilibration to room temperature is essential to achieve specified size distribution and yield of libraries.
- Beads will settle gradually; always ensure that they are fully resuspended before use.
- To ensure optimal DNA recovery, it is critical that the DNA and the KAPA Pure Beads are thoroughly mixed (by vortexing or extensive up-and-down pipetting) before the DNA binding incubation.
- Bead incubation times are guidelines only, and may be modified/optimized according to current protocols, previous experience, specific equipment and samples in order to maximize library construction efficiency and throughput.
- The time required for complete capture of beads varies according to the reaction vessel and magnet used. It is important not to discard or transfer any beads with the removal or transfer of supernatant. Capture times should be optimized accordingly.
- The volumes of 80% ethanol used for bead washes may be adjusted to accommodate smaller reaction vessels and/or limited pipetting capacity, but it is important that the beads are entirely submerged during the wash steps. Always use freshly prepared 80% ethanol.
- It is important to remove all ethanol before proceeding with subsequent reactions. However, over-drying of beads may make them difficult to resuspend, resulting in a dramatic loss of DNA. With optimized aspiration of ethanol, drying of beads for 3 – 5 min at room temperature should be sufficient. Drying of beads at 37°C is not recommended.
- Where appropriate, DNA should be eluted from beads in elution buffer (10 mM Tris-HCl, pH 8.0 – 8.5). Elution of DNA in PCR-grade water is not recommended, as DNA is unstable in unbuffered solutions. However, libraries constructed for target capture must be eluted and stored in PCR-grade water to facilitate drying of DNA prior to probe hybridization. Purified DNA in elution buffer should be stable at 4°C for 1 – 2 weeks, or at -20°C for long-term storage. The long-term stability of library DNA at -20°C depends on a number of factors, including library concentration. Always use low DNA-binding tubes for long-term storage, and avoid excessive freezing and thawing.
- Please refer to the KAPA Pure Beads Technical Data Sheet for additional detailed information and protocols.
Library Amplification Protocol

1. Library Amplification

Library Amplification Primer Mix (10X) (KK2623), sold separately, is designed to eliminate or delay primer depletion during library amplification reactions performed with KAPA HiFi HotStart ReadyMix (2X). 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 previously described. User-supplied primer mixes may be used in combination with incomplete or custom adapters. For guidelines on the formulation of user-supplied library amplification primers, please contact Technical Support at kapabiosystems.com/support.

1.1 Assemble each library amplification reaction as follows:

<table>
<thead>
<tr>
<th>Component</th>
<th>Volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>KAPA HiFi HotStart ReadyMix (2X)</td>
<td>25 μL</td>
</tr>
<tr>
<td>Library Amplification Primer Mix (10X)*</td>
<td>5 μL</td>
</tr>
<tr>
<td>Adapter-ligated library DNA</td>
<td>20 μL</td>
</tr>
<tr>
<td>Total volume:</td>
<td>50 μL</td>
</tr>
</tbody>
</table>

*Or another suitable 10X library amplification primer mix. The recommended final concentration of each primer in the library amplification reaction is 0.5 – 2 μM.

1.2 Mix thoroughly and centrifuge briefly.

1.3 Amplify using the following cycling protocol:

<table>
<thead>
<tr>
<th>Step</th>
<th>Temp.</th>
<th>Duration</th>
<th>Cycles</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial denaturation</td>
<td>98°C</td>
<td>45 sec</td>
<td>1</td>
</tr>
<tr>
<td>Denaturation</td>
<td>98°C</td>
<td>15 sec</td>
<td></td>
</tr>
<tr>
<td>Annealing*</td>
<td>60°C</td>
<td>30 sec</td>
<td>Minimum required for optimal amplification</td>
</tr>
<tr>
<td>Extension</td>
<td>72°C</td>
<td>30 sec</td>
<td></td>
</tr>
<tr>
<td>Final extension</td>
<td>72°C</td>
<td>1 min</td>
<td>1</td>
</tr>
<tr>
<td>HOLD</td>
<td>4°C</td>
<td>∞</td>
<td>1</td>
</tr>
</tbody>
</table>

*Optimization of the annealing temperature may be required for non-standard (i.e., other than Illumina TruSeq®) adapter/primer combinations.

*The optimal cycling number will depend upon the volume and concentration of adapter-ligated, size separated, purified library DNA added to each enrichment PCR reaction.

1.4 Proceed directly to Post-amplification Cleanup (step 2).

2. Post-amplification Cleanup

2.1 In the library amplification plate/tube(s), perform a 1X bead-based cleanup by combining the following:

<table>
<thead>
<tr>
<th>Component</th>
<th>Volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>Library amplification reaction product</td>
<td>50 μL</td>
</tr>
<tr>
<td>KAPA Pure Beads</td>
<td>50 μL</td>
</tr>
<tr>
<td>Total volume:</td>
<td>100 μL</td>
</tr>
</tbody>
</table>

2.2 Mix thoroughly by vortexing and/or pipetting up and down multiple times.

2.3 Incubate the plate/tube(s) at room temperature for 5 – 15 min to bind DNA to the beads.

2.4 Place the plate/tube(s) on a magnet to capture the beads. Incubate until the liquid is clear.

2.5 Carefully remove and discard the supernatant.

2.6 Keeping the plate/tube(s) on the magnet, add 200 μL of 80% ethanol.

2.7 Incubate the plate/tube(s) on the magnet at room temperature for ≥30 sec.

2.8 Carefully remove and discard the ethanol.

2.9 Keeping the plate/tube(s) on the magnet, add 200 μL of 80% ethanol.

2.10 Incubate the plate/tube(s) on the magnet at room temperature for ≥30 sec.

2.11 Carefully remove and discard the ethanol. Try to remove all residual ethanol without disturbing the beads.

2.12 Dry the beads at room temperature for 3 – 5 min, or until all of the ethanol has evaporated. Caution: over-drying the beads may result in reduced yield.

2.13 Remove the plate/tube(s) from the magnet.

2.14 Thoroughly resuspend the beads in an appropriate volume of elution buffer (10 mM Tris-HCl, pH 8.0 – 8.5) or PCR-grade water. Always use PCR-grade water if proceeding to target capture.

2.15 Incubate the plate/tube(s) at room temperature for 2 min to elute DNA off the beads.

2.16 Place the plate/tube(s) on a magnet to capture the beads. Incubate until the liquid is clear.

2.17 Transfer the clear supernatant to a new plate/tube(s). Store purified, amplified libraries at 4°C for 1 – 2 weeks, or at -20°C.
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This product has been manufactured under an ISO 13485:2003 quality management system.