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QIAseq[®] FX DNA Library Kit Handbook

For combined DNA fragmentation
and preparation of DNA libraries for
next-generation sequencing (NGS)
applications that use Illumina[®] instruments

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Kit Contents

| QIAseq FX DNA Library UDI Kit | UDI (24) | UDI-A (96) | UDI-B (96) | UDI-C (96) | UDI-D (96) |
|---|-----------------|-------------------|-------------------|-------------------|-------------------|
| Catalog no. | 180477 | 180479 | 180480 | 180481 | 180482 |
| Number of reactions | 24 | 96 | 96 | 96 | 96 |
| FX Enzyme Mix (violet cap) | 1 tube | 1 tube | 1 tube | 1 tube | 1 tube |
| FX Buffer, 10x (blue cap) | 1 tube | 1 tube | 1 tube | 1 tube | 1 tube |
| FX Enhancer (black cap) | 1 tube | 1 tube | 1 tube | 1 tube | 1 tube |
| DNA Ligase (red cap) | 1 tube | 1 tube | 1 tube | 1 tube | 1 tube |
| DNA Ligase Buffer, 5x (yellow cap) | 1 tube | 2 tubes | 2 tubes | 2 tubes | 2 tubes |
| RNase-Free Water (clear cap) | 2 tubes | 3 tubes | 3 tubes | 3 tubes | 3 tubes |
| HiFi PCR Master Mix, 2x (green cap) | 2 tubes | 2 tubes | 2 tubes | 2 tubes | 2 tubes |
| Primer Mix Illumina Library Amp, 10 μ M (clear cap) | 2 tubes | 1 tube | 1 tube | 1 tube | 1 tube |
| QIAseq UDI Y-Adapter Plate A,B,C, or D (96) | N/A | 1 plate | 1 plate | 1 plate | 1 plate |
| QIAseq UDI Y-Adapter Plate (24) | 1 plate | N/A | N/A | N/A | N/A |
| QIAseq Y-Adapter Reference Card | 1 | 1 | 1 | 1 | 1 |
| Quick-Start Protocol | 1 | 1 | 1 | 1 | 1 |

| QIAseq FX DNA Library CDI Kit | CDI (24) | CDI (96) |
|---|-----------------|-----------------|
| Catalog no. | 180483 | 180484 |
| Number of reactions | 24 | 96 |
| FX Enzyme Mix (violet cap) | 1 tube | 1 tube |
| FX Buffer, 10x (blue cap) | 1 tube | 1 tube |
| FX Enhancer (black cap) | 1 tube | 1 tube |
| DNA Ligase (red cap) | 1 tube | 1 tube |
| DNA Ligase Buffer, 5x (yellow cap) | 1 tube | 2 tubes |
| RNase-Free Water (clear cap) | 2 tubes | 3 tubes |
| HiFi PCR Master Mix, 2x (green cap) | 2 tubes | 2 tubes |
| Primer Mix Illumina Library Amp, 10 μ M (clear cap) | 2 tubes | 1 tube |
| QIAseq CDI Y-Adapter Plate (24) | 1 | N/A |
| QIAseq CDI Y-Adapter Plate (96) | N/A | 1 |
| QIAseq Y-Adapter Reference Card | 1 | 1 |
| Quick-Start Protocol | 1 | 1 |

| QIAseq CDI/UDI Y-Adapter Kit | CDI (24) | CDI (96) | UDI (24) | UDI A (96) | UDI B (96) | UDI C (96) | UDI D (96) |
|-------------------------------------|-----------------|-----------------|-----------------|-------------------|-------------------|-------------------|-------------------|
| Catalog no. | 180301 | 180303 | 180310 | 180312 | 180314 | 180316 | 180318 |
| Number of reactions | 24 | 96 | 24 | 96 | 96 | 96 | 96 |
| Adapter plate | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Reference card | 1 | 1 | 1 | 1 | 1 | 1 | 1 |

The QIAseq FX DNA Library Kits contain a QIAseq Y-Adapter plate with either combinatorial dual-index adapters (CDI) or unique dual-index adapters (UDI). To multiplex more than 96 libraries in a single sequencing run, combine kits with different UDI Y-adapter plates. For example, combining the QIAseq FX DNA Library UDI-A (or B or C or D) Kit (96) Kit will allow the generation of 384 libraries with different sample indexes for 384-plex sequencing. For more information on QIAseq Y-Adapter Plates, please refer to Appendix C, page 27.

Shipping and Storage

The QIAseq FX DNA Library Kits are shipped in 2 boxes (Library Core Kit and Adapter Kit). Store both at -30 to -15°C upon receipt. When stored correctly, all reagents are stable for at least 6 months after delivery if not otherwise stated on the label.

Intended Use

QIAseq FX DNA Library Kits are intended for molecular biology applications. This product is not intended for the diagnosis, prevention, or treatment of a disease.

All due care and attention should be exercised in the handling of the products. We recommend all users of QIAGEN[®] products to adhere to the NIH guidelines that have been developed for recombinant DNA experiments, or to other applicable guidelines.

Safety Information

When working with chemicals, always wear a suitable lab coat, disposable gloves, and protective goggles. For more information, please consult the appropriate safety data sheets (SDSs). These are available online in a convenient and compact PDF format at www.qiagen.com/safety where you can find, view, and print the SDS for each QIAGEN kit and kit component.

Quality Control

In accordance with QIAGEN's ISO-certified Quality Management System, each lot of the QIAseq FX DNA Library Kits is tested against predetermined specifications to ensure consistent product quality.

Introduction

Next-generation sequencing (NGS) is a driving force for numerous new and exciting applications, including cancer research, stem cell research, metagenomics, population genetics, and medical research. While NGS technology is continuously improving, library preparation remains one of the biggest bottlenecks in the NGS workflow and includes several time-consuming steps that can result in considerable sample loss and the potential to introduce handling errors. QIAGEN's QIAseq FX technology incorporates enzymatic DNA fragmentation into a streamlined, optimized protocol that does not require sample cleanup between fragmentation and adapter ligation, saving time and preventing errors. Optimized enzyme and buffer compositions ensure high sequencing library yield. Streamlined library construction protocols also enable straightforward automation of library prep on different liquid-handling platforms.

Principle and procedure

The QIAseq FX DNA Library Kit provides a fast, fully enzymatic procedure from DNA fragmentation to NGS library with no cleanup steps until after adapters have been ligated to the sample DNA. The kit allows library preparation from input DNA amounts ranging from as little as 20 pg up to 1 µg.

Samples consisting of longer DNA fragments, such as genomic DNA or amplicons from long-range PCR, are first enzymatically sheared into smaller fragments. The median fragment size is dependent on the applications and sequencing read length, and can be adjusted by varying the QIAseq FX DNA fragmentation reaction conditions. The fragmented DNA is directly end-repaired and an "A" is added to the 3' ends during the FX reaction, making the DNA fragments ready for adapter ligation. Following this step, platform-specific adapters are ligated to both ends of the DNA fragments. These adapters contain sequences essential for binding dual-barcoded libraries to a flow cell for sequencing, allowing for PCR amplification of adapter-ligated fragments, and binding standard Illumina sequencing primers.

To ensure maximum yields from limited amounts of starting material, a high-fidelity amplification step can be performed using the reagents included in the QIAseq FX DNA Library Kit. The proprietary HiFi PCR Master Mix can evenly amplify DNA regions with vastly different GC contents, minimizing sequencing bias caused by PCR.

Following library construction, the reaction cleanup and removal of adapter-dimers can be achieved by using Agencourt® AMPure® XP Beads (Beckman Coulter, cat. no. A63880), which enables easy automation on various high throughput automation platforms.

NGS adapter and index technologies

Sample multiplexing is one of the most important NGS tools for increasing throughput and reducing costs. It works by combining multiple samples to be processed together in a single sequencing run; as a consequence, sequencing reads need to be demultiplexed by reassigning each single read to its original source library. This is facilitated by the integration of index sequences into the individual adapter molecules.

QIAseq FX DNA Library CDI/UDI Kits include a fully compatible indexing solution. We recommend using the QIAseq Dual-Index Y-Adapter Plates delivered with the kit. Each QIAseq FX DNA Library CDI/UDI Kit includes one of the following:

- QIAseq Combinatorial Dual-Index (CDI) Y-Adapter Plate (24)
- QIAseq Combinatorial Dual-Index (CDI) Y-Adapter Plate (96)
- QIAseq Unique Dual-Index (UDI) Y-Adapter Plate (24)
- QIAseq Unique Dual-Index (UDI) Y-Adapter Plate A, B, C, or D (96)

Combining QIAseq FX DNA Library UDI-A/B/C/D (96) Kits enables multiplexing of up to 384 samples per sequencing run. For more information on QIAseq Dual-Index Y-Adapters and index sequence motives, see Appendix C, page 27, and “Ordering Information”, page 55.

CDI adapters use twelve i7 and eight i5 barcode motives that can be combined to form up to 96 combinatory dual indices. In contrast, QIAseq UDI Adapters use a fixed combination of 2 unique barcode motives per adapter molecule. Therefore, each single-index motive is only used once on any UDI adapter plate.

Usage of UDI adapters effectively mitigates the risk of read misassignment due to index hopping. This is enabled by filtering misassigned reads during the demultiplexing of individual samples, thus generating highly accurate output data.

Equipment and Reagents to Be Supplied by User

When working with chemicals, always wear a suitable lab coat, disposable gloves, and protective goggles. For more information, consult the appropriate safety data sheets (SDSs), available from the product supplier.

- Agencourt AMPure XP Beads (cat. no. A63880, A63881) for bead-based library purification
- 100% ethanol (ACS grade)
- Nuclease-free water
- Buffer EB (cat. no. 19086)
- PCR tubes or plates
- Pipette tips and pipettes
- DNA LoBind tubes (from Axygen or Eppendorf)
- Vortexer
- Microcentrifuge
- Thermocycler
- Magnetic racks for magnetic beads separation (e.g., Thermo Fisher Scientific DynaMag™ Magnet)
- Capillary electrophoresis device, e.g., QIAGEN QIAxcel, Agilent® Bioanalyzer® or similar to evaluate the DNA fragmentation profile (optional)

Important Notes

General precautions

- Use good laboratory practices to minimize cross-contamination of nucleic acid products.
- Always use PCR tubes, microcentrifuge tubes, and pipette tips that are certified sterile, DNase-free, and RNase-free.
- Before starting, wipe down work area and pipettes with an RNase- and DNA-cleaning product
- For consistent library construction and amplification, ensure that the thermocycler used in this protocol is in good working order and has been calibrated within the manufacturer's specifications.
- Read the entire protocol before beginning. Take note of stopping points where samples can be frozen at -20°C and plan your workflow accordingly.
- Enzyme-based DNA fragmentation is sensitive to many factors, such as reaction temperature, time, and setup conditions, as well as the quality of the input DNA.

DNA preparation and quality control

High-quality DNA is essential for obtaining reliable sequencing results. The most important prerequisite for any DNA sequence analysis experiment is consistent, high-quality DNA from every experimental sample. Therefore, sample handling and DNA isolation procedures are critical to the success of the experiment. Residual traces of proteins, salts, or other contaminants will degrade the DNA or decrease the efficiency of the enzymatic activities necessary for optimal library preparation.

It is important to remove all cations and chelators from DNA preparations, therefore, make sure DNA is eluted in QIAGEN's Buffer EB or H_2O , not 1x TE buffer containing 1mM EDTA. If the DNA was eluted or dissolved in 1x TE, or if you are not certain about the EDTA

concentration in the input DNA, we strongly recommend purifying the input DNA using Agencourt AMPure XP beads, following the instructions in “Appendix A: Removal of Divalent Cations and EDTA from Input Nucleic Acid”. Alternatively, we recommend setting up the FX reaction using the FX Enhancer as described in “Appendix B: Fragmentation, End-Repair, and A-Addition of DNA in 1x TE”.

Recommended genomic DNA preparation method

To prepare purified DNA, we recommend using an appropriate QIAGEN DNA purification kit that supports DNA elution in Buffer EB or 10mM Tris pH 8.0. For the best FX fragmentation performance, do not elute samples in a buffer containing >0.1 mM EDTA.

- QIAamp® DNA Mini Kit (cat. no. 51304) for the preparation of genomic DNA samples from fresh tissues and cells
- GeneRead™ DNA FFPE Kit (cat. no. 180134) for efficient recovery of high-quality gDNA from FFPE tissue
- MagAttract® HMW DNA Kit (cat. no. 67563) for isolation of high-molecular-weight genomic DNA
- QIAamp DNA Microbiome Kit (cat. no. 51704) for isolation of bacterial microbiome DNA from mixed samples

It is critical to accurately determine the input DNA concentration, especially when the input amount is below 100 ng. We recommend using Qubit®, PicoGreen® or another fluorometric method to accurately quantify DNA with a concentration below 1.5 ng/μl.

Protocol: Fragmentation, End-Repair, and A-addition

This protocol describes the FX reaction for single-tube fragmentation, end-repair, and A-addition.

Important points before starting

- Before setting up the reaction, it is critical to accurately determine the amount of the input DNA.
- Ensure input DNA is in water, 10 mM Tris, QIAGEN's Buffer EB or low TE (0.1x TE, 0.1 mM EDTA). If input DNA is in 1x TE, please set up the FX reaction according to the protocol in Appendix B.

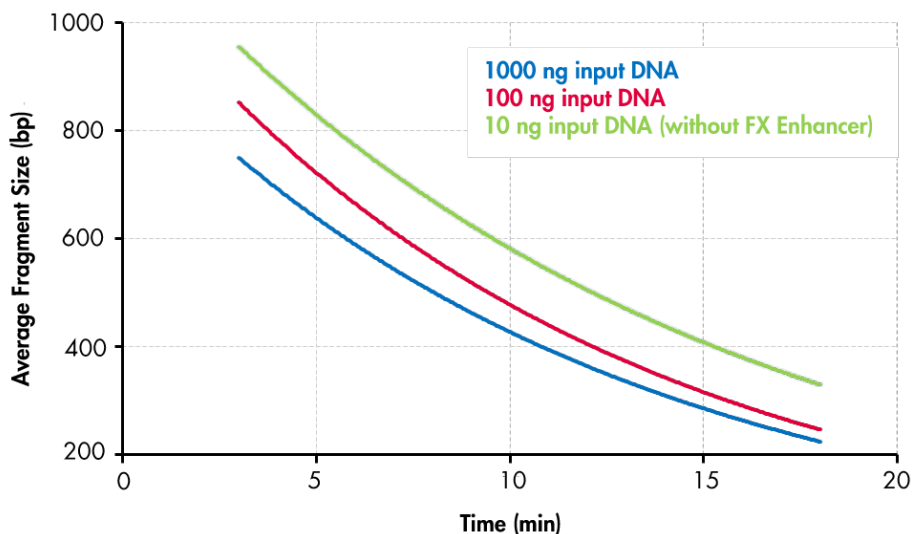


Figure 1. Fragmentation profile of different amounts of input DNA.

Table 1. Guideline for choosing the initial fragmentation time

| Fragment peak size | 250 bp | 350 bp | 450 bp | 550 bp |
|---|--------|--------|--------|--------|
| Fragmentation time (min) at 32°C | | | | |
| 50 pg –1 ng input DNA* | 14 | 4 | 1 | – |
| 10 ng input DNA† | 24 | 16 | 14 | 10 |
| 100 ng input DNA | 16 | 10 | 8 | 6 |
| 1000 ng input DNA | 14 | 8 | 6 | 4 |

Note: The same FX fragmentation time will produce a consistent fragment size within an approximately 5-fold range of input DNA amounts. The exact reaction time may need to be optimized for DNA samples of variable quality.

* For input DNA amounts between 20 and 50 pg, incubate the FX reaction including the FX Enhancer for 25 min to produce a fragment distribution centered around 250 bp.

† For input DNA <10 ng, FX Enhancer is required for optimal performance (Table 4). To produce a fragment distribution centered around 300 bp from 1 ng input, incubate the FX reaction including FX Enhancer for 10 min.

Things to do before starting

- Refer to Figure 1 and Table 1 to determine the time required to fragment input DNA to the desired size. If input DNA is less than 10 ng, add FX Enhancer according to the protocol and use half the reaction time listed for 10 ng input DNA. For example, to produce a fragment distribution centered around 300 bp from 1 ng input, incubate the FX reaction including FX Enhancer for 10 min.
- Make fresh 80% ethanol.
- Thaw reagents on ice. Once reagents are thawed, mix buffers thoroughly by quick vortexing to avoid any localized concentrations. Briefly spin down vortexed reagents before use.
- Program thermocyclers. For increased speed and convenience, all incubation steps of the protocol can be preprogrammed and saved in advance.

Procedure

1. Program a thermocycler according to Table 2 using the predetermined FX fragmentation time for step 2. Be sure to use the instrument's heated lid, and if possible, set the temperature of the heated lid to 70°C.

Table 2. Input DNA (20 pg –1000 ng) free of EDTA, Buffer EB, or in 0.1x TE

| Step | Incubation temperature | Incubation time |
|------|------------------------|-----------------|
| 1 | 4°C | 1 min |
| 2 | 32°C | 1–30 min* |
| 3 | 65°C | 30 min |
| 4 | 4°C | Hold |

* To determine the reaction time for step 2, refer to Figure 1 and Table 1.

2. Start the program. When the thermocycler block reaches 4°C, pause the program.
3. Prepare the FX reaction mix in a PCR plate or tube on ice according to Table 3 for >10 ng input DNA or Table 4 for <10 ng input DNA. Mix well by gently pipetting (do not vortex to mix).

Table 3. FX reaction mix setup (per sample) for >10 ng input DNA

| Component | Volume (µl) |
|------------------------------------|-------------|
| FX Buffer, 10x | 5 |
| Purified DNA | Variable |
| Nuclease-free water | Variable |
| Total without FX Enzyme Mix | 40 |

Table 4. FX reaction mix setup (per sample) for <10 ng input DNA

| Component | Volume (µl) |
|------------------------------------|-------------|
| FX Buffer, 10x | 5 |
| Purified DNA | Variable |
| FX Enhancer | 2.5 |
| Nuclease-free water | Variable |
| Total without FX Enzyme Mix | 40 |

4. Add 10 µl FX Enzyme Mix to each reaction and mix well by pipetting up and down 20 times. It is critical to keep the reactions on ice for the entire time during reaction setup.

-
5. Briefly spin down the PCR plate/tubes and immediately transfer to the prechilled thermocycler (4°C). Resume the cycling program.
 6. When the thermocycler program is complete and the sample block has returned to 4°C, remove samples and place them on ice.
 7. Immediately proceed with adapter ligation as described in the next protocol.

Protocol: Adapter Ligation

This protocol describes adapter ligation.

Things to do before starting

- Equilibrate Agencourt AMPure XP beads to room temperature (15–25°C) for 20–30 min before use.
- Vortex and spin down the thawed adapter plate before use.

Procedure

1. Remove the protective adapter plate lid, pierce the foil seal for each adapter well to be used, and transfer 5 µl from one DNA adapter well to each 50 µl sample from the previous protocol. Track the barcodes from each adapter well used for each sample.

Note: If your DNA input is <10 ng, dilute the adapters according to Table 5.

Table 5. Adapter dilution factors

| Sample DNA amount | Adapter dilution |
|-------------------|------------------|
| 20–99 pg | 1:1000 |
| 100–999 pg | 1:100 |
| 1–9 ng | 1:10 |

2. Replace the adapter plate lid and freeze unused adapters. The adapter plate is stable for a minimum of 10 freeze-thaw cycles.

Important: Only 1 single adapter should be used per ligation reaction. If adapters from another supplier are used, follow the manufacturer’s instructions. Do not reuse adapter wells once the foil seal has been pierced.

3. Prepare the ligation Master Mix (per DNA sample, Table 6) in a separate PCR plate or tube on ice, and mix well by pipetting.

Table 6. Ligation master mix setup (per sample)

| Component | Volume (µl) |
|---------------------|-------------|
| Ligation buffer, 5x | 20 |
| DNA ligase | 10 |
| Nuclease-free water | 15 |
| Total | 45 |

4. Add 45 µl of the ligation Master Mix to each sample, for a total of 100 µl, and mix well by pipetting. Incubate the ligation reaction at 20°C for 15 min.
Important: Do not use a thermocycler with a heated lid.
5. Proceed immediately to adapter ligation cleanup using 0.8x (80 µl) Agencourt AMPure XP beads.
6. Add 80 µl of resuspended Agencourt AMPure XP beads to each ligated sample and mix well by pipetting.
7. Incubate the mixture for 5 min at room temperature. Pellet the beads on a magnetic stand (e.g., DynaMag) for 2 min, then carefully discard the supernatant.
8. Wash the beads by adding 200 µl of 80% ethanol. Pellet the beads on the magnetic stand and discard the supernatant. Repeat the wash once, for a total of 2 ethanol washes. Remove as much excess ethanol as possible.
9. Incubate the beads on the magnetic stand for 5–10 min or until the beads are dry. Overdrying of beads may result in lower DNA recovery. Remove from the magnetic stand.
10. Elute by resuspending in 52.5 µl of Buffer EB or 10 mM Tris-Cl, pH 8.0. Pellet the beads on the magnetic stand. Carefully transfer 50 µl of supernatant into a new plate or tube.
11. Perform a second purification using 1x (50 µl) Agencourt AMPure XP beads following steps 7–9 for DNA binding and washing. Elute DNA by adding 26 µl Buffer EB or 10 mM Tris-Cl, pH 8.0. Pellet the beads and carefully collect 23.5 µl of purified DNA sample in a DNA LoBind tube for library amplification. If not proceeding immediately, the sample can be stored at –30 to –15°C.

Protocol: Amplification of Library DNA

PCR-based library amplification is normally required if the input DNA amount is below 100 ng or if large amounts of libraries are required for downstream hybrid capture. This protocol is for high-fidelity amplification of the DNA library using the amplification reagents provided in the QIAseq FX DNA Library Kit.

Things to do before starting

- Thaw QIAseq HiFi PCR Master Mix and Primer Mix on ice. Once reagents are thawed, mix them thoroughly by quick vortexing to avoid any localized concentrations.
- Always start with the cycling conditions specified in this protocol. The cycling has been optimized for use with QIAseq HiFi PCR Master Mix for even and high-fidelity amplification of sequencing libraries.

Procedure

1. Program a thermocycler with a heated lid according to Table 7.

Table 7. Library amplification cycling conditions

| Time | Temperature | Number of cycles |
|-------|-------------|--|
| 2 min | 98°C | 1 |
| 20 s | 98°C | |
| 30 s | 60°C | 6 (100 ng input DNA) 10 (10 ng input DNA) 12 (1 ng input DNA) 14 (100 pg input DNA) 16 (20 pg input DNA) |
| 30 s | 72°C | |
| 1 min | 72°C | 1 |
| ∞ | 4°C | Hold |

Note: 6–16 amplification cycles are recommended based on the input DNA amount and quality.

2. Prepare a reaction mix on ice according to Table 8. Mix the components in a PCR tube or 96-well PCR plate.

Table 8. Reaction mix for library enrichment

| Component | Volume (µl) |
|------------------------------|-------------|
| HiFi PCR Master Mix, 2x | 25 |
| Primer Mix (10 µM each) | 1.5 |
| Library DNA | 23.5 |
| Total reaction volume | 50 |

3. Transfer the PCR tube or plate to the thermocycler and start the program.
4. Once PCR is complete, add 50 µl of resuspended Agencourt AMPure XP Beads to each reaction (50 µl) and pipet up and down thoroughly to mix.
5. Incubate the mixture for 5 min at room temperature. Pellet the beads on a magnetic stand (e.g., DynaMag) and carefully discard the supernatant.
6. Wash the beads by adding 200 µl of 80% ethanol. Pellet the beads on the magnetic stand and discard the supernatant. Repeat the wash once, for a total of 2 ethanol washes. Remove as much excess ethanol as possible.
7. Incubate on the magnetic stand for 5–10 min or until the beads are dry. Overdrying of beads may result in lower DNA recovery. Remove from the magnetic stand.
8. Elute by resuspending in 25 µl of Buffer EB or 10 mM Tris-Cl, pH 8.0. Pellet the beads on the magnetic stand. Carefully transfer 23 µl of the supernatant into a new tube.
9. Assess the quality of the library using a capillary electrophoresis device such as QIAGEN QIAxcel or Agilent BioAnalyzer. Check for the expected size distribution (see Figure 2) of library fragments and for the absence of an adapters or adapter-dimers peak around 120 bp.

Note: The library should show a distribution centered around the size of the fragmented DNA plus 120 bp (see Figure 2). The increase in library length reflects the addition of sequencing adapters to the DNA fragments.

Note: The median fragment size can be used in subsequent qPCR-based quantification methods to calculate library concentration (step 10).

10. Quantify the library using a qPCR-based method such as the QIAseq Library Quant Assay Kit (cat. no. 333314; not provided), or a comparable method.
11. The purified library can be safely stored at -30 to -15°C in a DNA LoBind tube until ready to use for sequencing or other applications.

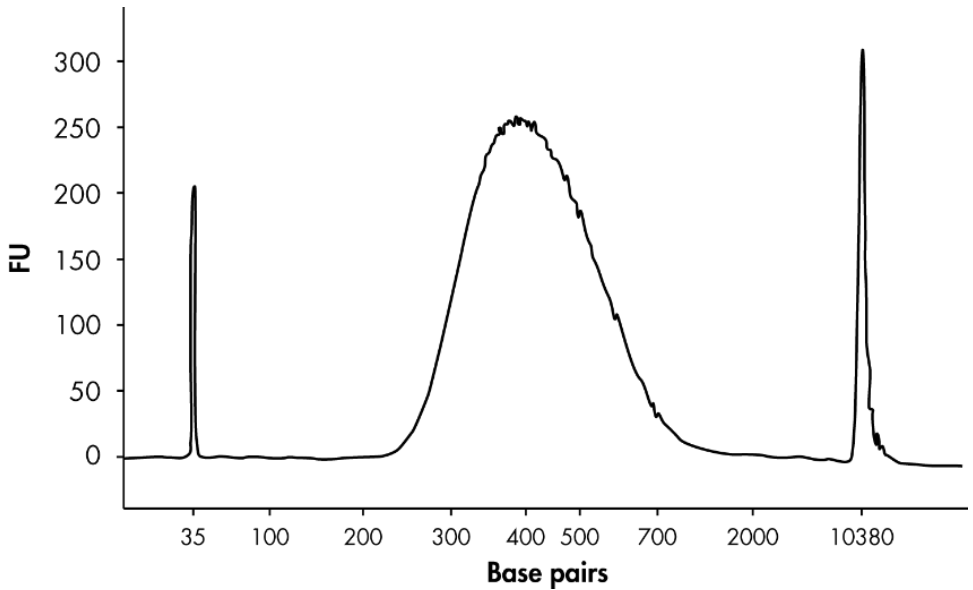


Figure 2. Capillary electrophoresis device trace data.

Troubleshooting Guide

This troubleshooting guide may be helpful in solving any problems that may arise. For more information, see also the Frequently Asked Questions page at our Technical Support Center: www.qiagen.com/FAQ/FAQList.aspx. The scientists in QIAGEN Technical Services are always happy to answer any questions you may have about either the information and/or protocols in this handbook or sample and assay technologies (for contact information, visit www.qiagen.com).

Comments and suggestions

Low library yields

- | | |
|--|---|
| a) Suboptimal reaction conditions due to low DNA quality | Make sure to use high-quality DNA to ensure optimal activity of the library enzymes. |
| b) Insufficient amount of starting DNA for direct sequencing without library amplification | Typically, 100 ng of sheared genomic DNA generates enough Illumina-compatible library to use directly for sequencing without amplification. If the final library yield is not sufficient for the expected number of sequencing runs, a library amplification step can be performed following the adapter ligation step. |
| c) Inaccurate quantification of starting DNA due to RNA contamination. | RNA from the sample material can be co-purified with genomic DNA. This contaminating RNA will affect the accuracy of DNA quantification. To remove RNA during the sample preparation protocol, we recommend to perform RNase A treatment of the DNA. |

Unexpected signal peaks in capillary electrophoresis device traces

- | | |
|--|---|
| a) Presence of shorter peaks between 60 and 120 bp | These peaks represent library adapters and adapter-dimers that occur when there is no, or insufficient, adapter depletion after library preparation. As adapter-dimers can form clusters on the flow cell and will be sequenced, this will reduce the capacity of the flow cell for the library fragments, even though a low ratio of adapter-dimers versus library will not be a problem. Agencourt AMPure XP Beads or GeneRead Size Selection Kit (cat. no. 180514) efficiently remove adapter-dimers, as well as free adapter molecules. |
| b) Presence of larger library fragments after library enrichment | If the fragment population shifts higher than expected after adapter ligation and PCR enrichment (e.g., more than the expected 120 bp shift), this can be a PCR artifact due to over-amplification of the DNA library. Ensure that you use as few amplification cycles as possible to avoid this effect. |

Comments and suggestions

- | | |
|---|--|
| c) Incorrect library fragment size after adapter ligation | During library preparation, adapters of approximately 60 bp are ligated to both ends of the DNA library fragments. This should be reflected on a capillary electrophoresis device by a shift in size of all library fragments of 120 bp. If using library adapters from other suppliers, refer to the size information given in the respective documentation. The absence of a clear size shift may indicate no, or only low, adapter ligation efficiency. Ensure that you use the parameters and incubation times described in the handbook for end-repair, A-addition and ligation, as well as the correct amount of starting DNA. |
| d) Incorrect DNA fragment size prior to adapter ligation | The wrong DNA fragment size prior to adapter ligation can be due to the wrong conditions used for enzymatic DNA fragmentation. The reaction time should be optimized for different amount of input DNA. For input DNA >10 ng, we recommend 12 min as a starting point as it produces fragmentation size centers around 300 to 500 bp. Depending on the size requirement and type of input DNA, either increase or decrease reaction time by 2–4 min incrementally until expected size range is achieved. |

Appendix A: Removal of Divalent Cations and EDTA from Input Nucleic Acid

Refer to manufacturer's protocol for details on methods of purification

1. If DNA is in a volume of less than 50 μ l, adjust the volume to 50 μ l with nuclease-free water.
2. Add 90 μ l of resuspended Agencourt AMPure XP beads to the reaction for a ratio of 1.8x and mix well by pipetting. If DNA is in a volume greater than 50 μ l, scale the volume of Agencourt AMPure XP beads appropriately such that the ratio of beads to DNA is 1.8x.
3. Incubate the mixture for 5 min at room temperature. Pellet the beads on a magnetic stand for 2–4 min and carefully discard the supernatant without disturbing the beads.
4. Wash the beads with 200 μ l of 80% ethanol. Pellet the beads on the magnetic stand and discard the supernatant. Repeat the wash once for a total of 2 ethanol washes. Remove as much excess ethanol as possible.
5. Incubate the beads on the magnetic stand for 5–10 min or until the beads are dry. Remove from the magnetic stand.
6. Elute by resuspending in 45 μ l of QIAGEN's Buffer EB or 10 mM Tris-Cl, pH 8.0. Pellet the beads on the magnetic stand for 2 min. Carefully transfer 42.5 μ l of supernatant into a new tube.
7. Determine the concentration of the purified DNA using Qubit, PicoGreen, or another fluorometric method.

Appendix B: Fragmentation, End-Repair, and A-Addition of DNA in 1x TE

Follow the instructions below for input DNA in 1X TE buffer.

1. Enter the following program into a thermocycler (Table 9). Ensure that you use the instrument's heated lid, and if possible, set the temperature of the heated lid to ~70°C.

Table 9. Input DNA (20 pg – 1000 ng) in 1x TE

| Step | Incubation temperature | Incubation time |
|------|------------------------|-----------------|
| 1 | 4°C | 1 min |
| 2 | 32°C | 1–30 min* |
| 3 | 65°C | 30 min |
| 4 | 4°C | Hold |

* To determine the reaction time for step 2, please refer to Figure 1 and Table 1.

2. Prepare the FX reaction mix in a PCR plate on ice according to Table 10 for >10 ng input DNA or Table 11 for <10 ng input DNA. Mix well by gently pipetting (do not vortex). The reaction can be scaled as needed for the desired number of samples.

Table 10. Input DNA (10–1000 ng) in 1x TE

| Component | Volume (µl) |
|---------------------|-------------|
| FX Buffer, 10x | 5 |
| DNA in 1x TE | Variable |
| FX Enhancer | 2.5 |
| Nuclease-free water | Variable |
| Total | 40 |

Table 11. Input DNA 20 pg – 10 ng in 1x TE

| Component | Volume (µl) |
|---------------------|--------------------|
| FX Buffer, 10x | 5 |
| DNA in 1x TE | Variable |
| FX Enhancer | 5 |
| Nuclease-free water | Variable |
| Total | 40 |

3. Add 10 µl FX Enzyme Mix to reach reaction and mix well by pipetting up and down 20 times. It is critical to keep the PCR tube on ice for the entire time during reaction setup.
4. Briefly spin down the PCR plate/tubes and immediately transfer to the prechilled thermocycler (4°C). Resume the cycling program.
5. When thermocycler program is complete and sample block has returned to 4°C, remove samples and place on ice.
6. Immediately proceed to adapter ligation as described in the adapter ligation protocol.

Appendix C: QIAseq Dual-Index Y-Adapters

Generation of sample sheets for Illumina instruments

Index sequences for QIAseq Unique and Combinatorial Dual-Index Y-Adapters are available for download at www.qiagen.com. Sequencing on the NextSeq, HiSeq X™, or HiSeq 3000/4000 system follows a different dual-indexing workflow than other Illumina systems. If you are manually creating sample sheets for these instruments, enter the reverse complement of the i5 index adapter sequence. If you are using Illumina Experiment Manager, BaseSpace, or Local Run Manager for run planning, the software will automatically reverse complement index sequences when necessary.

Ready-to-use sample sheets containing all QIAseq CDI and UDI Y-Adapter barcode sequences are available for MiSeq, NextSeq, MiniSeq, HiSeq, and NovaSeq instruments. These can be imported and edited using the Illumina Experiment Manager Software, Illumina Local Run Manager, or any text editor. Make sure to download the appropriate sample sheet for NextSeq, HiSeq X, or HiSeq 3000/4000 systems depending on whether you are using Local Run Manager or manually configuring the sequencing run

Unique Dual-Index Y-Adapters

The layout of the 24-plex and 96-plex (A/B/C/D) single-use UDI adapter plate is shown in Figure 3 to Figure 7. The index motives used in the QIAseq Unique Dual-Index Kits are listed in Table 12. To make sequencing preparation more convenient, you can download Illumina-compatible sample sheets for different sequencing instruments at www.qiagen.com.

| | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 |
|---|---------|---------|---------|-------|-------|-------|-------|-------|-------|-------|-------|-------|
| A | UDI 001 | UDI 009 | UDI 017 | empty | empty | empty | empty | empty | empty | empty | empty | empty |
| B | UDI 002 | UDI 010 | UDI 018 | empty | empty | empty | empty | empty | empty | empty | empty | empty |
| C | UDI 003 | UDI 011 | UDI 019 | empty | empty | empty | empty | empty | empty | empty | empty | empty |
| D | UDI 004 | UDI 012 | UDI 020 | empty | empty | empty | empty | empty | empty | empty | empty | empty |
| E | UDI 005 | UDI 013 | UDI 021 | empty | empty | empty | empty | empty | empty | empty | empty | empty |
| F | UDI 006 | UDI 014 | UDI 022 | empty | empty | empty | empty | empty | empty | empty | empty | empty |
| G | UDI 007 | UDI 015 | UDI 023 | empty | empty | empty | empty | empty | empty | empty | empty | empty |
| H | UDI 008 | UDI 016 | UDI 024 | empty | empty | empty | empty | empty | empty | empty | empty | empty |

Figure 3. QIAseq UDI Y-Adapter Plate (24) layout (UDI 1–24).

| | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 |
|---|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|
| A | UDI 001 | UDI 009 | UDI 017 | UDI 025 | UDI 033 | UDI 041 | UDI 049 | UDI 057 | UDI 065 | UDI 073 | UDI 081 | UDI 089 |
| B | UDI 002 | UDI 010 | UDI 018 | UDI 026 | UDI 034 | UDI 042 | UDI 050 | UDI 058 | UDI 066 | UDI 074 | UDI 082 | UDI 090 |
| C | UDI 003 | UDI 011 | UDI 019 | UDI 027 | UDI 035 | UDI 043 | UDI 051 | UDI 059 | UDI 067 | UDI 075 | UDI 083 | UDI 091 |
| D | UDI 004 | UDI 012 | UDI 020 | UDI 028 | UDI 036 | UDI 044 | UDI 052 | UDI 060 | UDI 068 | UDI 076 | UDI 084 | UDI 092 |
| E | UDI 005 | UDI 013 | UDI 021 | UDI 029 | UDI 037 | UDI 045 | UDI 053 | UDI 061 | UDI 069 | UDI 077 | UDI 085 | UDI 093 |
| F | UDI 006 | UDI 014 | UDI 022 | UDI 030 | UDI 038 | UDI 046 | UDI 054 | UDI 062 | UDI 070 | UDI 078 | UDI 086 | UDI 094 |
| G | UDI 007 | UDI 015 | UDI 023 | UDI 031 | UDI 039 | UDI 047 | UDI 055 | UDI 063 | UDI 071 | UDI 079 | UDI 087 | UDI 095 |
| H | UDI 008 | UDI 016 | UDI 024 | UDI 032 | UDI 040 | UDI 048 | UDI 056 | UDI 064 | UDI 072 | UDI 080 | UDI 088 | UDI 096 |

Figure 4. QIAseq UDI Y-Adapter Plate A (96) layout (UDI 1–96).

| | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 |
|----------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| A | UDI 097 | UDI 105 | UDI 113 | UDI 121 | UDI 129 | UDI 137 | UDI 145 | UDI 153 | UDI 161 | UDI 169 | UDI 177 | UDI 185 |
| B | UDI 098 | UDI 106 | UDI 114 | UDI 122 | UDI 130 | UDI 138 | UDI 146 | UDI 154 | UDI 162 | UDI 170 | UDI 178 | UDI 186 |
| C | UDI 099 | UDI 107 | UDI 115 | UDI 123 | UDI 131 | UDI 139 | UDI 147 | UDI 155 | UDI 163 | UDI 171 | UDI 179 | UDI 187 |
| D | UDI 100 | UDI 108 | UDI 116 | UDI 124 | UDI 132 | UDI 140 | UDI 148 | UDI 156 | UDI 164 | UDI 172 | UDI 180 | UDI 188 |
| E | UDI 101 | UDI 109 | UDI 117 | UDI 125 | UDI 133 | UDI 141 | UDI 149 | UDI 157 | UDI 165 | UDI 173 | UDI 181 | UDI 189 |
| F | UDI 102 | UDI 110 | UDI 118 | UDI 126 | UDI 134 | UDI 142 | UDI 150 | UDI 158 | UDI 166 | UDI 174 | UDI 182 | UDI 190 |
| G | UDI 103 | UDI 111 | UDI 119 | UDI 127 | UDI 135 | UDI 143 | UDI 151 | UDI 159 | UDI 167 | UDI 175 | UDI 183 | UDI 191 |
| H | UDI 104 | UDI 112 | UDI 120 | UDI 128 | UDI 136 | UDI 144 | UDI 152 | UDI 160 | UDI 168 | UDI 176 | UDI 184 | UDI 192 |

Figure 5. QIAseq UDI Y-Adapter Plate B (96) layout (UDI 97–192).

| | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 |
|----------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| A | UDI 193 | UDI 201 | UDI 209 | UDI 217 | UDI 225 | UDI 233 | UDI 241 | UDI 249 | UDI 257 | UDI 265 | UDI 273 | UDI 281 |
| B | UDI 194 | UDI 202 | UDI 210 | UDI 218 | UDI 226 | UDI 234 | UDI 242 | UDI 250 | UDI 258 | UDI 266 | UDI 274 | UDI 282 |
| C | UDI 195 | UDI 203 | UDI 211 | UDI 219 | UDI 227 | UDI 235 | UDI 243 | UDI 251 | UDI 259 | UDI 267 | UDI 275 | UDI 283 |
| D | UDI 196 | UDI 204 | UDI 212 | UDI 220 | UDI 228 | UDI 236 | UDI 244 | UDI 252 | UDI 260 | UDI 268 | UDI 276 | UDI 284 |
| E | UDI 197 | UDI 205 | UDI 213 | UDI 221 | UDI 229 | UDI 237 | UDI 245 | UDI 253 | UDI 261 | UDI 269 | UDI 277 | UDI 285 |
| F | UDI 198 | UDI 206 | UDI 214 | UDI 222 | UDI 230 | UDI 238 | UDI 246 | UDI 254 | UDI 262 | UDI 270 | UDI 278 | UDI 286 |
| G | UDI 199 | UDI 207 | UDI 215 | UDI 223 | UDI 231 | UDI 239 | UDI 247 | UDI 255 | UDI 263 | UDI 271 | UDI 279 | UDI 287 |
| H | UDI 200 | UDI 208 | UDI 216 | UDI 224 | UDI 232 | UDI 240 | UDI 248 | UDI 256 | UDI 264 | UDI 272 | UDI 280 | UDI 288 |

Figure 6. QIAseq UDI Y-Adapter Plate C (96) layout (UDI 193–288).

| | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 |
|---|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| A | UDI 289 | UDI 297 | UDI 305 | UDI 313 | UDI 321 | UDI 329 | UDI 337 | UDI 345 | UDI 353 | UDI 361 | UDI 369 | UDI 377 |
| B | UDI 290 | UDI 298 | UDI 306 | UDI 314 | UDI 322 | UDI 330 | UDI 338 | UDI 346 | UDI 354 | UDI 362 | UDI 370 | UDI 378 |
| C | UDI 291 | UDI 299 | UDI 307 | UDI 315 | UDI 323 | UDI 331 | UDI 339 | UDI 347 | UDI 355 | UDI 363 | UDI 371 | UDI 379 |
| D | UDI 292 | UDI 300 | UDI 308 | UDI 316 | UDI 324 | UDI 332 | UDI 340 | UDI 348 | UDI 356 | UDI 364 | UDI 372 | UDI 380 |
| E | UDI 293 | UDI 301 | UDI 309 | UDI 317 | UDI 325 | UDI 333 | UDI 341 | UDI 349 | UDI 357 | UDI 365 | UDI 373 | UDI 381 |
| F | UDI 294 | UDI 302 | UDI 310 | UDI 318 | UDI 326 | UDI 334 | UDI 342 | UDI 350 | UDI 358 | UDI 366 | UDI 374 | UDI 382 |
| G | UDI 295 | UDI 303 | UDI 311 | UDI 319 | UDI 327 | UDI 335 | UDI 343 | UDI 351 | UDI 359 | UDI 367 | UDI 375 | UDI 383 |
| H | UDI 296 | UDI 304 | UDI 312 | UDI 320 | UDI 328 | UDI 336 | UDI 344 | UDI 352 | UDI 360 | UDI 368 | UDI 376 | UDI 384 |

Figure 7. QIAseq UDI Y-Adapter Plate D (96) layout (UDI 289–384).

Table 12. UDI motives used in the QIAseq UDI Y-Adapter Kits (24 and 96 A/B/C/D)

Unique Dual-Index adapters 1–24 are identical on the adapter plates of the QIAseq UDI Y-Adapter Kit (24) and QIAseq UDI Y-Adapter Kit A (96).

Note: Sequencing on the MiniSeq, NextSeq, HiSeqX, and HiSeq 3000/4000 systems follow a different dual-indexing workflow than other Illumina systems, which requires the reverse complement of the i5 index adapter sequence.

| Indices for entry on sample sheet | | | |
|-----------------------------------|---|---|--|
| Unique dual-index number | i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000) | i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000) | i7 bases for entry on sample sheet (all instruments) |
| UDI 001 | ATGGCCGACT | AGTCGGCCAT | TGAACGTTGT |
| UDI 002 | CGATGAGCAC | GTGCTCATCG | ACCAGACTTG |
| UDI 003 | GATAAGTCGA | TCGACTTATC | ACTGGCGAAC |
| UDI 004 | TCACGCCTTG | CAAGGCGTGA | GCGTTAGGCA |
| UDI 005 | AGGAACACAA | TTGTGTTCTT | TTATCGGCCT |
| UDI 006 | CTCAGTAGGC | GCCTACTGAG | GAGGTATAAG |
| UDI 007 | GAAGTGCCTG | CAGGCACCTC | TCAAGGATTC |
| UDI 008 | TCTCTCGCCT | AGGCGAGAGA | CGAACCGAGA |
| UDI 009 | AGGCACCTTC | GAAGGTGCCT | GAGCCAAGTT |
| UDI 010 | CTGTTGGTAA | TTACCAACAG | AAGGCCGTAG |
| UDI 011 | GCTGGTACCT | AGGTACCAGC | TTAGAGAAGC |
| UDI 012 | TAAGGAGCGG | CCGCTCCTTA | TCTAAGACCA |
| UDI 013 | AATCGCTCCA | TGGAGCGATT | TGTAACCACT |
| UDI 014 | CTCCTAATTG | CAATTAGGAG | CCGACACAAG |
| UDI 015 | GCCTCATAAT | ATTATGAGGC | CTCTGATGGC |
| UDI 016 | TGTATTGAGC | GCTCAATACA | CGGCCTGTTA |
| UDI 017 | AGCCATAACA | TGTTATGGCT | TGCATAGCTT |
| UDI 018 | CCACAAGTGG | CCACTTGTGG | AACCTTCTCG |
| UDI 019 | GTTATCACAC | GTGTGATAAC | AAGAGATCAC |
| UDI 020 | TACCGTTCTT | AAGAACGGTA | GCCTGAAGGA |

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| Indices for entry on sample sheet | | | |
|--|--|--|---|
| Unique dual-index number | i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000) | i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000) | i7 bases for entry on sample sheet (all instruments) |
| UDI 021 | AGGCGTTAGG | CCTAACGCCT | ATTGTGCCTT |
| UDI 022 | CCGTAACGTC | GACGTACGG | TCCTCTACCG |
| UDI 023 | GTAATAGCCA | TGGCTATTAC | TACCATGAAC |
| UDI 024 | TAGCGCCGAT | ATCGGCGCTA | CATTGGCAGA |
| UDI 025 | CATTCTTGGGA | TCCAAGAATG | CACTGCTATT |
| UDI 026 | ATGCAAGGTT | AACCTGCAT | AATGGTAGGT |
| UDI 027 | CGCCAGACAA | TTGTCTGGCG | GATACCTATG |
| UDI 028 | GAAGGTTGGC | GCCAACCTTC | CACTAGGTAC |
| UDI 029 | TCGCATCACG | CGTGATGCGA | AGCTCGTCA |
| UDI 030 | CCGTCATGA | TCATGACCGG | TGTCAGTCTT |
| UDI 031 | ATCACAAGC | GCTTGTAAT | GATGAACAGT |
| UDI 032 | CAACCTGTAA | TTACAGGTTG | ACAATCGGCG |
| UDI 033 | GCCAGTCGTT | AACGACTGGC | GATTGAGTTC |
| UDI 034 | TGCCTTGTCG | CGACAAGGCA | GTAATGCCAA |
| UDI 035 | CTATCCGCTG | CAGCGGATAG | TCGTTGCGCT |
| UDI 036 | AATGCCGGAA | TTCCGGCATT | AGGTGAGTAT |
| UDI 037 | CGGTTATCCG | CGGATAACCG | TCGATAATGG |
| UDI 038 | GCGGAAGAGT | ACTCTCCGC | GCGTCTCTTC |
| UDI 039 | TTGGTTAGTC | GACTAACCAA | GTCTCCTGCA |
| UDI 040 | TTCAGTGTGA | TCACACTGAA | GAGCTTCATT |

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| Indices for entry on sample sheet | | | |
|--|--|--|---|
| Unique dual-index number | i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000) | i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000) | i7 bases for entry on sample sheet (all instruments) |
| UDI 041 | AGAATTCTGG | CCAGAATTCT | AGGCCTACAT |
| UDI 042 | CATTGACTCT | AGAGTCAATG | TGTGGAACCG |
| UDI 043 | GCGGCTTCAA | TTGAAGCCGC | CGTATTAAGC |
| UDI 044 | TTATGGTCTC | GAGACCATAA | CCAGTGGTTA |
| UDI 045 | CGTAACCAGG | CCTGGTACG | GCGTTCGAGT |
| UDI 046 | AGCTCAGATA | TATCTGAGCT | CCTCCGGTT |
| UDI 047 | CCGGTGTTAC | GTAACACCGG | CACAAGACGG |
| UDI 048 | GACCTAACCT | AGGTTAGGTC | GCTTACACAC |
| UDI 049 | TTGTAGAAGG | CCTTCTACAA | AGGATGTCCA |
| UDI 050 | CCTAGCACTA | TAGTGCTAGG | CACCTTATGT |
| UDI 051 | ATCGTGTCT | AGAACACGAT | AAGCGGCTGT |
| UDI 052 | CCAACTTATC | GATAAGTTGG | TTCCTGTGAG |
| UDI 053 | GAAGCCAAGG | CCTGGCCTC | AGTACAGTTC |
| UDI 054 | TGGAGTCAA | TTGAACTCCA | TACAGCCTCA |
| UDI 055 | CTCAATCCT | AGGATTGAAG | GTTCTATTGG |
| UDI 056 | ATCTTGCGTG | CACGCAAGAT | ATATACCGGT |
| UDI 057 | CGTCTAAGGT | ACCTTAGACG | CCTCGGAATG |
| UDI 058 | GAGGTGAACA | TGTTACCTC | GTTCTGGAAC |
| UDI 059 | TCAGAACTAC | GTAGTCTGA | AGATTCACCA |
| UDI 060 | CGGATATTGA | TCAATATCCG | TCGGTCAGAT |

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| Indices for entry on sample sheet | | | |
|--|--|--|---|
| Unique dual-index number | i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000) | i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000) | i7 bases for entry on sample sheet (all instruments) |
| UDI 061 | AGGAGTAGAT | ATCTACTCCT | CACTCTCGCT |
| UDI 062 | CCGCCGAATA | TATTCGGCGG | GTTGGTCCAG |
| UDI 063 | GAGTCTATAC | GTATAGACTC | AGCTCGAAGC |
| UDI 064 | TTATTACCGG | CCGGTAATAA | AGAGGTTCTA |
| UDI 065 | CGCTCGTTAG | CTAACGAGCG | ATGACTCGAA |
| UDI 066 | AACAACGCTG | CAGCGTGTGT | GAACAATCCT |
| UDI 067 | CGCGGTATT | AATAGCCGCG | TGGCAAGGAG |
| UDI 068 | GCTCGACACA | TGTGTCGAGC | GAATATTGGC |
| UDI 069 | TTCTCCAAC | GTTGGAAGAA | CCGGAACCTA |
| UDI 070 | TTGGCGGTTG | CAACCGCCAA | ACTTGTTCGG |
| UDI 071 | AACAGGCAAT | ATTGCCTGTT | CAAGTCCAAT |
| UDI 072 | CAGAATGGCG | CGCCATTCTG | AACCGCAAGG |
| UDI 073 | GTTGAGATTC | GAATCTCAAC | ACGTTGACTC |
| UDI 074 | TGTGTGCGGA | TCCGCACACA | CCACTTAACA |
| UDI 075 | GTTCCGCGAA | TTCGCCGAAC | AGCAGTTCCT |
| UDI 076 | AGCTGTATTG | CAATACAGCT | TCGCCTTCGT |
| UDI 077 | CAGCGGATGA | TCATCCGCTG | TAGGACTGCG |
| UDI 078 | GTCCTGGAT | ATCCAAGGAC | TCCGAGCGAA |
| UDI 079 | TCTAGATGCT | AGCATCTAGA | TTCGGTGTGT |
| UDI 080 | CGAGCCACAT | ATGTGGCTCG | ACAGGAGGAA |

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| Indices for entry on sample sheet | | | |
|--|--|--|---|
| Unique dual-index number | i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000) | i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000) | i7 bases for entry on sample sheet (all instruments) |
| UDI 081 | ATGGAATGGA | TCCATTCCAT | CCTCCATTAA |
| UDI 082 | CATTCTCAC | GTGAGGAATG | AGTCGCGGTT |
| UDI 083 | GCATAGGAAG | CTTCCTATGC | CTCATCCAGG |
| UDI 084 | TGTTCTGTGT | AACACGAACA | TGTGGTTGAA |
| UDI 085 | TAAGACCGTT | AACGGTCTTA | TTATGCGTGG |
| UDI 086 | ATGGTACCAG | CTGGTACCAT | GCGAATGTAT |
| UDI 087 | CCGACAGCTT | AAGCTGTCGG | GTC AAGCTCG |
| UDI 088 | GACGATATGA | TCATATCGTC | TAGAGTTGGA |
| UDI 089 | TTGTA CTCCA | TGGAGTACAA | CTGATGATCT |
| UDI 090 | GTGCACATAA | TTATGTGCAC | ACTAGGTGTT |
| UDI 091 | AGGACAAGTA | TACTTGTCCT | CTGTTAGCGG |
| UDI 092 | CCGATTCGAG | CTCGAATCGG | ATCGCACCAA |
| UDI 093 | GTAGGA ACTT | AAGTTCCTAC | CITACTGGT |
| UDI 094 | TACACTACGA | TCGTAGTGTA | CCTTAATGCG |
| UDI 095 | ATGACCTTGA | TCAAGGTCAT | TCTCGCTAG |
| UDI 096 | CTACGTGACG | CGTCACGTAG | TCTTCAGAGA |
| UDI 097 | AACAATCAGG | CCTGATTGTT | TACCGGTGGT |
| UDI 098 | CTGGTGTGCA | TGCACACCAG | AGGTGTTACG |
| UDI 099 | GCATATCCTT | AAGGATATGC | ACAGACCGAC |
| UDI 100 | TGTCCTGTAC | GTACAGGACA | CGAATACGTA |

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| Indices for entry on sample sheet | | | |
|--|--|--|---|
| Unique dual-index number | i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000) | i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000) | i7 bases for entry on sample sheet (all instruments) |
| UDI 101 | AGAACGTCGC | GCGACGTCT | TAGCATCGAT |
| UDI 102 | CACGGACTAG | CTAGTCCGTG | CCATGAGTCG |
| UDI 103 | GTTGAACACT | AGTGTCAAC | ACTAACATGC |
| UDI 104 | TCGCGTGGTA | TACCACGCGA | ACACTCTCTA |
| UDI 105 | AGCCACTATG | CATAGTGGCT | GCTCTTGCTT |
| UDI 106 | CCACCTACCA | TGGTAGGTGG | AATCTTGAGG |
| UDI 107 | GTTCCGGTGT | ACACCGGAAC | CTAACGGTC |
| UDI 108 | TAGGTCTGAC | GTCAGACCTA | TTGTGACCAA |
| UDI 109 | AGGAAGCATT | AATGCTTCT | TCACACACCT |
| UDI 110 | CCTTAGTTGG | CCAACAAAGG | CTGCAATTAG |
| UDI 111 | GTCCTATTCA | TGAATAGGAC | CTCCTTACTC |
| UDI 112 | TAAGATGGAC | GTCCATCTTA | GCAACGCAGA |
| UDI 113 | AGGCCATGGT | ACCATGGCCT | CCTTACCAAT |
| UDI 114 | CATTGGCCAA | TTGGCCAATG | TTAATCCTCG |
| UDI 115 | GCTATGAATC | GATTCATAGC | TTCCGAGTTC |
| UDI 116 | TTGGTCTCG | CGAGGACCAA | CTCGAGAGGA |
| UDI 117 | AGCGACATAC | GTATGTCGCT | TGTTGGCTGT |
| UDI 118 | CAAGTAGTCT | AGACTACTTG | CGTATCTGCG |
| UDI 119 | GTCAAGAAGA | TCTTCTGAC | CCATAGTATC |
| UDI 120 | TCCTGTTATG | CATAACAGGA | TGGACAGTAA |

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| Indices for entry on sample sheet | | | |
|--|--|--|---|
| Unique dual-index number | i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000) | i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000) | i7 bases for entry on sample sheet (all instruments) |
| UDI 121 | AAGTGC GATA | TATCGCACTT | GTACCTTGTT |
| UDI 122 | AGGCTACACG | CGTGTAGCCT | GAGTGCCTCT |
| UDI 123 | CTATATCGGC | GCCGATATAG | TAAGTAGCGG |
| UDI 124 | GCTAAGGTAA | TTACCTTAGC | CGTGGTGTTT |
| UDI 125 | TAACCTGGTT | AACCAGGTTA | CATTCCTGAA |
| UDI 126 | AGTTGGTCTA | TAGACCAACT | AAGATGCATG |
| UDI 127 | ATGCAGCTGG | CCAGTGCAT | CCTTGGAGCT |
| UDI 128 | CGTTGCCTTC | GAAGGCAACG | ACCGGAACAG |
| UDI 129 | GCGTGGAGAA | TTCTCCACGC | GAATGGAAGC |
| UDI 130 | TACGCCTCCT | AGGAGGCGTA | GTTCTCCATA |
| UDI 131 | AATTCGGTAG | CTACCGAATT | GTCACTATGT |
| UDI 132 | ATTGTGCAAC | GTTGACAAT | TGGTAGAACT |
| UDI 133 | CAACCTTGCG | CGCAAGGTTG | ACGCCTATGG |
| UDI 134 | GCACTGCGTA | TACGCAGTGC | AATCCGTAC |
| UDI 135 | TGCTAGTAGT | ACTACTAGCA | GTTGAGGCTA |
| UDI 136 | AAGTCACGGA | TCCGTGACTT | TATCAACTGG |
| UDI 137 | AGCGATTGAA | TTCAATCGCT | AAGAGGAGAT |
| UDI 138 | CTACCTCTCT | AGAGAGGTAG | GTCTTCTCGG |
| UDI 139 | GACAACGTGC | GACAGTTGTC | GAAGCCACTC |
| UDI 140 | TCCATTGCGG | CCGCAATGGA | GTAGGACACA |

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| Indices for entry on sample sheet | | | |
|--|--|--|---|
| Unique dual-index number | i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000) | i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000) | i7 bases for entry on sample sheet (all instruments) |
| UDI 141 | AGCCTCGCAA | TTGCGAGGCT | CTCCTCGTAT |
| UDI 142 | AATACAGGCT | AGCCTGTATT | CCACATGATT |
| UDI 143 | CGGACCGTTA | TAACGGTCCG | AGACGGTTGG |
| UDI 144 | GCGCTTATGC | GCATAAGCGC | CTAGGTTGAC |
| UDI 145 | TTAACACGAG | CTCGTGTTAA | AAGCGTACCA |
| UDI 146 | CGCCTCTAGA | TCTAGAGGCG | TCATGTTGGT |
| UDI 147 | AATCGACCTT | AAGGTCGATT | TTGGAATGGT |
| UDI 148 | CCGCAATAAC | GTTATTGCGG | GTGTATGTTG |
| UDI 149 | GTTCCAACGA | TCGTTGGAAC | TCCTGTCAAC |
| UDI 150 | TGTTAGACCG | CGGTCTAACA | TAATCAGGCA |
| UDI 151 | AACCTCATAG | CTATGAGGTT | GTAGTGGATT |
| UDI 152 | ATGAATCCAC | GTGGATTCAT | AATTGCGCAT |
| UDI 153 | CGGCTTAATT | AATTAAGCCG | GACAATAACG |
| UDI 154 | GAGTGCAGG | CCTGCAACTC | ACAGTTAAGC |
| UDI 155 | TCCACGAACA | TGTTCTGTTGA | AGCCACACTA |
| UDI 156 | TGACGGAGGA | TCCTCCGTC | CAATCGTCTT |
| UDI 157 | AATGAGTACG | CGTACTCATT | AGGAGCTTGT |
| UDI 158 | CGTCTCCGA | TCGGAAGACG | TTGAGCGGAG |
| UDI 159 | GACAGAGATT | AATCTCTGTC | AGTAGCTCTC |
| UDI 160 | TTACGCTAAC | GTTAGCGTAA | CACGCTGTCA |

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| Indices for entry on sample sheet | | | |
|--|--|--|---|
| Unique dual-index number | i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000) | i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000) | i7 bases for entry on sample sheet (all instruments) |
| UDI 161 | CTCCTCGAAG | CTTCGAGGAG | AAGACCTCTT |
| UDI 162 | ATACCGCAGA | TCTGCGGTAT | GACCTCTTCT |
| UDI 163 | CCTATCTGAT | ATCAGATAGG | TACTTCCTTG |
| UDI 164 | GATCGGTAC | GTAACCGATC | TGCGATACGC |
| UDI 165 | TGGTGAGGTG | CACCTACCA | GCAGGCTTAA |
| UDI 166 | AACCGGCGTA | TACGCCGGTT | TAAGCTTGTG |
| UDI 167 | AATACCGATC | GATCGGTATT | ATGGTCCGCT |
| UDI 168 | CGATACTCAA | TTGAGTATCG | ATGTCAGAAG |
| UDI 169 | GTAAGGCGGT | ACCGCCTTAC | GACGAAGGTC |
| UDI 170 | TTC AAGTCTG | CGACCTTGAA | ATCACCGTGA |
| UDI 171 | TATCCGAGTA | TACTCGGATA | GCTACAGTGT |
| UDI 172 | AGCGCGCTTA | TAAGCGCGCT | CGTCAATAT |
| UDI 173 | CCGGAGACAT | ATGTCTCCGG | CAACCATCGG |
| UDI 174 | GAGATAACTG | CAGTTATCTC | CGGTCCATC |
| UDI 175 | TTGTAAGCGC | GCGCTTACAA | AGAAGAGCCA |
| UDI 176 | CAAGAGGAGG | CCTCTCTTG | CTATGCAATG |
| UDI 177 | AACCTTAGGA | TCCTAAGGTT | CACTGAACCG |
| UDI 178 | CTGGCAACTC | GAGTTGCCAG | TACTGTGTGA |
| UDI 179 | GAACCTGTTG | CAACAAGTTC | GCATTCTGTT |
| UDI 180 | TGTGCAAGAT | ATCTTGACACA | CTCCGCTAAG |

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| Indices for entry on sample sheet | | | |
|--|--|--|---|
| Unique dual-index number | i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000) | i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000) | i7 bases for entry on sample sheet (all instruments) |
| UDI 181 | AATCGAGAGA | TCTCTCGATT | TCGCTTGAGA |
| UDI 182 | AGCGTGTACAG | CTGACACGCT | AACTAGCCTT |
| UDI 183 | CTTGGTGATT | AATCACCAAG | TTGCTCAGG |
| UDI 184 | GAAGCAGCAA | TTGCTGCTTC | CTCTACAACA |
| UDI 185 | TTCCGTGAC | GTCGACGGAA | TGAGTGTGTT |
| UDI 186 | CGAGATGCCA | TGGCATCTCG | TAGTTAGTCG |
| UDI 187 | AAGTTCGTGC | GCACGAACTT | GCCTGATCCT |
| UDI 188 | CGTCCATAAG | CTTATGGACG | CGAGTACAGG |
| UDI 189 | TTGTGGCATA | TATGCCACAA | GCCTAGATTA |
| UDI 190 | AGATCGGAAT | ATTCCGATCT | TCGGCACTGT |
| UDI 191 | CATTCTACTG | CAGTAGAATG | CCGTGCAAGA |
| UDI 192 | ATCGCCGTAG | CTACGGCGAT | CTGGCTGGTT |
| UDI 193 | ATCCTTACAC | GTGTAAGGAT | CGTTAGGATT |
| UDI 194 | CGCAAGGACT | AGTCCTTGCG | TTCCATTACG |
| UDI 195 | GCTGGCGTTA | TAACGCCAGC | TAGCGGTAAC |
| UDI 196 | TACTTAGAGG | CCTCTAAGTA | GTAGCCAGGA |
| UDI 197 | ATGGCGATGC | GCATCGCCAT | AGGATACTCT |
| UDI 198 | CATTGGTGCG | CGCACCAATG | TATCTCCAG |
| UDI 199 | GCGAGATATA | TATATCTCGC | TAAGTCGTTT |
| UDI 200 | TGACTGCTAT | ATAGCAGTCA | TCCGGATTGA |

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| Indices for entry on sample sheet | | | |
|--|--|--|---|
| Unique dual-index number | i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000) | i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000) | i7 bases for entry on sample sheet (all instruments) |
| UDI 201 | AACGTCCGCT | AGCGGACGTT | ACGTCTTGTT |
| UDI 202 | CGCACATGTC | GACATGTGCG | ATGAAGTGCG |
| UDI 203 | GCACACCTGA | TCAGGTGTGC | CGATCACTGC |
| UDI 204 | TTGTCCAGAG | CTCTGGACAA | CCTATCGGAA |
| UDI 205 | AGCCTTCTCG | CAGGAAGGCT | CAGAGAGCTT |
| UDI 206 | CCTTAGCCCA | TGGCGTAAGG | GCAACTTGCG |
| UDI 207 | GAATACGTAC | GTACGTATTC | TATGGAGGAC |
| UDI 208 | TTGGCACCGT | ACGGTGCCAA | TGAGATCAGA |
| UDI 209 | ATTAGGTGGC | GCCACCTAAT | TCAGCCTATT |
| UDI 210 | CGATCAAGAA | TTCTTGATCG | GTTGTGAGCG |
| UDI 211 | GCTGTCTTCT | AGAAGACAGC | TCAGTAACAC |
| UDI 212 | TACATGTCTG | CAGACATGTA | AAGGCTCAGA |
| UDI 213 | AACCAGTTGA | TCAACTGGTT | GTGTGGTGGT |
| UDI 214 | CCGTAAGCT | AGCTTACCGG | CCGAGCTTAG |
| UDI 215 | GTTCAATAG | CTATTGCAAC | ATCACGCTTC |
| UDI 216 | TGTCAGGCTC | GAGCCTGACA | TAGCTATGCA |
| UDI 217 | CAACAGTGTT | AACACTGTTG | TGTTCTCAT |
| UDI 218 | AAGAGAGGAA | TTCTTCTTT | CATACCTTCT |
| UDI 219 | CGGTGTAGC | GCTACAACCG | GCCTTCAATG |
| UDI 220 | GCCTGAAGTG | CAC TTCAGGC | CTTGACCAGC |

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| Indices for entry on sample sheet | | | |
|--|--|--|---|
| Unique dual-index number | i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000) | i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000) | i7 bases for entry on sample sheet (all instruments) |
| UDI 221 | TTACGACACT | AGTGTCTGATA | CTACACACAA |
| UDI 222 | CGCCTAGATC | GATCTAGGCG | TAGGCTGAAT |
| UDI 223 | AATCTGGATG | CATCCAGATT | TCGGAGTCTT |
| UDI 224 | CGACGGTACA | TGTACCGTCG | AACATCGCGG |
| UDI 225 | GTAGTATTGC | GCAATACTAC | GTTGTCTTAC |
| UDI 226 | TCCAGCGGAT | ATCCGCTGGA | GTGGCAACTA |
| UDI 227 | CAACCACCTC | GAGGTGGTTG | GAGCAGGCAT |
| UDI 228 | AGCTTAGGCG | CGCCTAAGCT | AACGGCACCT |
| UDI 229 | CCGTTCTCTT | AAGGAACCGG | AGTAACCTTG |
| UDI 230 | GACATTGAAC | GTTCAATGTC | TCTCATAAGC |
| UDI 231 | TTAGAGGCGA | TCGCCTCTAA | TGCTTGCCAA |
| UDI 232 | CAAGCCGAAC | GTTGCGCTTG | CGGTTCTCTG |
| UDI 233 | AGGAGAACGG | CCGTTCTCTT | CCAAGTAGAT |
| UDI 234 | CCTGTTAGAC | GTCTAACAGG | AAGGTTGGCG |
| UDI 235 | GTTCTACGTT | AACGTAGAAC | TGCTCTGGTC |
| UDI 236 | TAAGTCCACA | TGTGGACTTA | ACTGTAACGA |
| UDI 237 | CAAGAACCAT | ATGGTTCTTG | GATTCCAGGT |
| UDI 238 | AGTTGATGAC | GTCATCAACT | TTCACCAGAT |
| UDI 239 | CCTACTCTTG | CAAGAGTAGG | ACTTCCAAGG |
| UDI 240 | GAACAATCCA | TGGATTGTTC | CCGAATATTC |

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| Indices for entry on sample sheet | | | |
|--|--|--|---|
| Unique dual-index number | i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000) | i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000) | i7 bases for entry on sample sheet (all instruments) |
| UDI 241 | TTCTGTTGGT | ACCAACAGAA | CTCTATCCA |
| UDI 242 | CATCGTCAGG | CCTGACGATG | TCACAGCGGT |
| UDI 243 | ATGCATGAAG | CTTCATGCAT | CCTCTGTCGT |
| UDI 244 | CGTGAATCGC | GCGATTCACG | TCTGTTCTCG |
| UDI 245 | GAGCAGCCTT | AAGGCTGCTC | GATACTTCAC |
| UDI 246 | TCGATTACCA | TGGAATCGA | AGTGCTGATA |
| UDI 247 | CAGTCCAATT | AATTGGACTG | ATCCTTCGGT |
| UDI 248 | AGAGGCTTGG | CCAAGCCTCT | GACAACGATT |
| UDI 249 | CAGGCTCTCA | TGAGAGCCTG | GAACCGGTAG |
| UDI 250 | GTTGCCTCTC | GAGAGCGAAC | AGCAATGAGC |
| UDI 251 | TCGGACTAAT | ATTAGTCCGA | CAAGACTCCA |
| UDI 252 | CGAGATCTTC | GAAGATCTCG | ACCGTGTAGG |
| UDI 253 | ATAACCGGAC | GTCCGGTTAT | AGGCACAGGT |
| UDI 254 | CGGTAGTTA | TAACTACACG | CGACAGATCG |
| UDI 255 | GAACATAGGT | ACCTATGTC | ACGCGACAAC |
| UDI 256 | TCTAACATCG | CGATGTTAGA | ACTTGCCTTA |
| UDI 257 | AACGGTGGCA | TGCCACCGTT | CACCACTCAT |
| UDI 258 | AGGACGGTGT | ACACCGTCTT | CTTCGTAACT |
| UDI 259 | CTGTGACCTG | CAGGTCACAG | CAGTATTCGG |
| UDI 260 | GCTGTAACAA | TTGTTACAGC | CAGTCTGGAC |

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| Indices for entry on sample sheet | | | |
|--|--|--|---|
| Unique dual-index number | i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000) | i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000) | i7 bases for entry on sample sheet (all instruments) |
| UDI 261 | TACGGACGTC | GACGTCCGTA | TACCGTCTCA |
| UDI 262 | CCTAAGGAGC | GCTCCTTAGG | GTGTCCACAG |
| UDI 263 | ATAAGGCCAG | CTGGCCTTAT | TTACGACTGT |
| UDI 264 | CTCATCTGTA | TACAGATGAG | GACGCGAATG |
| UDI 265 | GAAGGCATCT | AGATGCCTTC | CAACGTACGC |
| UDI 266 | TCTCTACTGC | GCAGTAGAGA | AGCTCAGGAA |
| UDI 267 | AACCGAACAA | TTGTTCCGTT | GATAGGCCGT |
| UDI 268 | ATCTCGCCAC | GTGGCGAGAT | AGTAGGAAGT |
| UDI 269 | CCATGCAACG | CGTTGCATGG | CATGTTGTAG |
| UDI 270 | GAATGGTGTA | TACACCATTC | CACATTCTTC |
| UDI 271 | TATATGCCGT | ACGGCATATA | GCAGCTCGTA |
| UDI 272 | CTCGATAGAT | ATCTATCGAG | GTTCAGACGG |
| UDI 273 | AACACAAGAG | CTCTTGTT | TCCTGGAAGT |
| UDI 274 | CGCAATCGGT | ACCGATTGCG | GCATTGTTAG |
| UDI 275 | GTTGCGTAGA | TCTACGCAAC | GACCTACAGC |
| UDI 276 | TAGAGTGATC | GATCACTCTA | CACCGACGTA |
| UDI 277 | AAGACGCAGC | GCTGCGTCTT | CTCTCACCTT |
| UDI 278 | AACTTCTCGA | TCGAGAAGTT | CTCGTTCATT |
| UDI 279 | CGCAACTGAG | CTCAGTTGCG | TGGTGGCAAG |
| UDI 280 | GCTCCGCAAT | ATTGCGGAGC | GATTGCTTGA |

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| Indices for entry on sample sheet | | | |
|--|--|--|---|
| Unique dual-index number | i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000) | i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000) | i7 bases for entry on sample sheet (all instruments) |
| UDI 281 | GTAACCTCCG | CGGAAGTTAC | CCGTTAAGGT |
| UDI 282 | CTCACGACTA | TAGTCGTGAG | TGCTGAGAGG |
| UDI 283 | AACCAACGGC | GCCGTGGTT | TTGTCACTTG |
| UDI 284 | CCTGCCTGTA | TACAGGCAGG | GCTGTTATGT |
| UDI 285 | TACGCTGCAG | CTGCAGCGTA | GCAGCAGTTG |
| UDI 286 | AATGTTGCGA | TCGCAACATT | GCAGATCAAT |
| UDI 287 | CGACGTTCTG | CAGAACGTCG | TGGTTCACGG |
| UDI 288 | AATAGGACAC | GTGTCCTATT | TCGACCGCAT |
| UDI 289 | ATGTGCCTCA | TGAGGCACAT | TAACCTAGGT |
| UDI 290 | CGACTCCGTT | AACGGAGTCG | AACTCATGCG |
| UDI 291 | GCTGTTGTGG | CCACAACAGC | CCGGATGAAC |
| UDI 292 | TACCAATCAC | GTGATTGGTA | CGTTGCCGTA |
| UDI 293 | ATGCTTACG | CGTAAGACAT | GCTCTACGGT |
| UDI 294 | CGCAACAATA | TATTGTTGCG | TGCATTGGCG |
| UDI 295 | GAACGAAGAC | GTCTTCGTTT | CGATTGTGAC |
| UDI 296 | TCGAGGACGT | ACGTCCTCGA | GACTGCACTA |
| UDI 297 | ATTATGAGCG | CGCTCATAAT | GTTAACTGCT |
| UDI 298 | CGCGTTATAA | TTATAACGCG | TCGGACCTTG |
| UDI 299 | GCGTGTCATGT | ACATGCACGC | TGCAGCAAGC |
| UDI 300 | TAAGCGGCTC | GAGCCGCTTA | CACATGCGAA |

Table continues on next page

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| Indices for entry on sample sheet | | | |
|--|--|--|---|
| Unique dual-index number | i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000) | i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000) | i7 bases for entry on sample sheet (all instruments) |
| UDI 301 | AACATGGAGA | TCTCCATGTT | CAGACGTAAT |
| UDI 302 | CCGAGTCTCT | AGAGACTCGG | ATTCGGTACG |
| UDI 303 | GTA CTCTAC | GTAGAAGTAC | TTAGCACGGC |
| UDI 304 | TGTTACATG | CATGTGAACA | GAGGATAGTA |
| UDI 305 | AAGTAACGC | GCGTTACCTT | AACTGTGGTT |
| UDI 306 | CCGCCTACT | AGTAAGGCGG | ATTACCTCGG |
| UDI 307 | GTTGAGGCAG | CTGCCTCAAC | CGTGTATAC |
| UDI 308 | TGCGGACCTA | TAGGTCGCCA | CTTGCTCACA |
| UDI 309 | AGAAGCGACA | TGTCGCTTCT | CAACACCTGT |
| UDI 310 | CAGGATAATC | GATTATCCTG | CAATTGCTCG |
| UDI 311 | GCTCCTACAG | CTGTAGGAGC | CATAGACAAC |
| UDI 312 | TCAACAGGT | ACCTGTTGAA | TTGGTGTCTA |
| UDI 313 | CCTCGTCCAT | ATGGACGAGG | TATGTCCTGT |
| UDI 314 | AGCGTTGGTT | AACCAACGCT | GCCAAATCGT |
| UDI 315 | CATTGAACA | TGTTCAATG | TAGGCGATCG |
| UDI 316 | GCTTACCGAC | GTCGGTAAGC | ATGAGTGTAC |
| UDI 317 | TTAGCTTAGG | CCTAAGCTAA | CCGAAGGATA |
| UDI 318 | CCGACACACA | TGTGTGTCGG | AGTCCACTGT |
| UDI 319 | ATTCGCTGAT | ATCAGCGAAT | GCGGCTAATT |
| UDI 320 | CCAAGAGGCA | TGCCTCTGG | TCTAACTCAG |

Table continues on next page

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| Indices for entry on sample sheet | | | |
|--|--|--|---|
| Unique dual-index number | i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000) | i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000) | i7 bases for entry on sample sheet (all instruments) |
| UDI 321 | GACGCAGTTC | GAAC TGCGTC | CAAGCTGAGC |
| UDI 322 | TGGAACTCGG | CCGAGTCCA | CCAGGCACA |
| UDI 323 | CCACACCAAT | ATTGGTGTGG | TGTACAAGGT |
| UDI 324 | AGTTCTCGGC | GCCGAGAACT | TAGAATGCCT |
| UDI 325 | CTTGACGACG | CGTCGTCAAG | TGCTTACTG |
| UDI 326 | GAGGTCGCTA | TAGCGACCTC | ATGACTAAGC |
| UDI 327 | TCAGTAGCAT | ATGCTACTGA | ATGTAGGCAA |
| UDI 328 | CTAACGTGGA | TCCACGTTAG | GCGAAGAGGT |
| UDI 329 | ATGCCAACCG | CGGTTGGCAT | CGGTGGTCT |
| UDI 330 | CGGTCGATTC | GAATCGACCG | CTGTCGTGG |
| UDI 331 | GAAGTACAGT | ACTGTACTTC | TGATCGACAC |
| UDI 332 | TCTGCAGTAA | TTACTGCAGA | CCACCAGCTA |
| UDI 333 | CTATCCTAGC | GCTAGGATAG | CACGGTTCGT |
| UDI 334 | AACACTCCTT | AAGGAGTGTT | AGTGAGAGCT |
| UDI 335 | CCGAACCTAA | TTAGGTTCCGG | TTGCATGCGG |
| UDI 336 | GTCTAGTCGC | GCGACTAGAC | TATACGTGTC |
| UDI 337 | TGGATGTACG | CGTACATCCA | TGACGCGTTA |
| UDI 338 | CTACCAGCGT | ACGCTGGTAG | TACAGAACGT |
| UDI 339 | AAGGATTACAG | CTGAATCCTT | CTTGTCAGGT |
| UDI 340 | CGAGGTGTGT | ACACACCTCG | ATCCACAGCG |

Table continues on next page

Table continued from previous page

| Indices for entry on sample sheet | | | |
|--|--|--|---|
| Unique dual-index number | i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000) | i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000) | i7 bases for entry on sample sheet (all instruments) |
| UDI 341 | GTAGACGCTC | GAGCGTCTAC | CCTATCCATC |
| UDI 342 | TCGTCCGTCA | TGACGGACGA | ACCGCGAGTA |
| UDI 343 | CCGTGATAGG | CCTATCACGG | AAGTTCTGGT |
| UDI 344 | AGGATGACCT | AGGTCATCCT | ACAGGTATCG |
| UDI 345 | CCTCGAGTAC | GTACTIONGAGG | ATGACGGATT |
| UDI 346 | GTCACTGAGG | CCTCAGTGAC | GTCTGAGTAG |
| UDI 347 | TACGGTTAGA | TCTAACCGTA | TGCCAGATGT |
| UDI 348 | CAACGAGAAT | ATTCTCGTTG | GCTAAGCATT |
| UDI 349 | AATACACCGG | CCGGTGTATT | ACAGCATGGT |
| UDI 350 | CCGATCCATC | GATGGATCGG | ATAGAGACCG |
| UDI 351 | GAATCTCGCT | AGCGAGATTC | ATATCGCGTA |
| UDI 352 | TGACCGGCAA | TTGCCGGTCA | TTAAGGAGGT |
| UDI 353 | CATGATAGCA | TGCTATCATG | CTGTGCGACT |
| UDI 354 | AACAGCTTCG | CGAAGCTGTT | TCCGTATGCT |
| UDI 355 | CTAGTGCTTA | TAAGCACTAG | CCATCGATGT |
| UDI 356 | TGTGATACGT | ACGTATCACA | GTGAGCCGTT |
| UDI 357 | ATGAGCGTAT | ATACGCTCAT | TGCCGTTAAT |
| UDI 358 | CTAGATATGG | CCATATCTAG | CGGATGTGGT |
| UDI 359 | CGCTATGCTG | CAGCATAGCG | TCGCGTGTG |
| UDI 360 | TACTACGTGA | TCACGTAGTA | CCGCGATCAT |

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Table continued from previous page

| Indices for entry on sample sheet | | | |
|--|--|--|---|
| Unique dual-index number | i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000) | i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000) | i7 bases for entry on sample sheet (all instruments) |
| UDI 361 | ATGTGGAGGT | ACCTCCACAT | CGCGTTATCG |
| UDI 362 | CCATGGCTCA | TGAGCCATGG | GTAGCCTCCT |
| UDI 363 | CCAATCACGC | GCGTGATTGG | ACTAGACACT |
| UDI 364 | TTAGATCCAG | CTGGATCTAA | CGATTCTGTG |
| UDI 365 | AGGAATATCG | CGATATTCTT | GAAGAGATGT |
| UDI 366 | CCTCCTATGT | ACATAGGAGG | AGATCCGACG |
| UDI 367 | TAGAGACACG | CGTGTCTCTA | CCAGGACATT |
| UDI 368 | CCAGCTCAGT | ACTGAGCTGG | ACGTGGCATT |
| UDI 369 | ATGGCTCATA | TATGAGCCAT | AAGCAGGACG |
| UDI 370 | CGGAGTGAAG | CTTCACTCCG | ACGAGTCGGT |
| UDI 371 | TACCTATGGT | ACCATAGGTA | AGTGTACGCG |
| UDI 372 | ATGAGACAGT | ACTGTCTCAT | ACCGACCATT |
| UDI 373 | CTAAGAGTTG | CAACTCTTAG | TTGCTAACGT |
| UDI 374 | TAACCGTATG | CATACGGTTA | CTTGATACTG |
| UDI 375 | AGAGTCCATG | CATGGACTCT | CTGATAAAGT |
| UDI 376 | CTAGACCGCA | TGCGGTCTAG | ATAGCTTACG |
| UDI 377 | TATGGCTTGT | ACAAGCCATA | GTCCATGAGT |
| UDI 378 | CGTTGTCTCT | AGGAACAACG | ACTCCAGTCG |
| UDI 379 | CCGACATTAG | CTAATGTCGG | TCTCAGCACG |
| UDI 380 | TGTGAAGGCA | TGCCTTCACA | ATCGTGATGT |
| UDI 381 | AGCATCGTCT | AGACGATGCT | ACGCAATCCG |
| UDI 382 | CCGACTAGGA | TCCTAGTCGG | GAGATCGGCT |
| UDI 383 | AACATTACCG | CGGTAATGTT | CTACGTCTCG |
| UDI 384 | CCTAATTCGT | ACGAATTAGG | CTCAGGCTGT |

Combinatorial Dual-Index Y-Adapters

The layout of the 24-plex and 96-plex single-use CDI adapter plate is shown in Figure 8 and Figure 9. The index motives used in the QIAseq Combinatorial Dual-Index Kits are listed in Table 13. To make sequencing preparation more convenient, you can download Illumina-compatible sample sheets for different sequencing instruments at www.qiagen.com.

| | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 |
|---|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|
| A | 501/ 701 | 501/ 702 | 501/ 703 | 501/ 704 | 501/ 705 | 501/ 706 | 501/ 707 | 501/ 708 | 501/ 709 | 501/ 710 | 501/ 711 | 501/ 712 |
| B | 502/ 701 | 502/ 702 | 502/ 703 | 502/ 704 | 502/ 705 | 502/ 706 | 502/ 707 | 502/ 708 | 502/ 709 | 502/ 710 | 502/ 711 | 502/ 712 |
| C | 503/ 701 | 503/ 702 | 503/ 703 | 503/ 704 | 503/ 705 | 503/ 706 | 503/ 707 | 503/ 708 | 503/ 709 | 503/ 710 | 503/ 711 | 503/ 712 |
| D | 504/ 701 | 504/ 702 | 504/ 703 | 504/ 704 | 504/ 705 | 504/ 706 | 504/ 707 | 504/ 708 | 504/ 709 | 504/ 710 | 504/ 711 | 504/ 712 |
| E | 505/ 701 | 505/ 702 | 505/ 703 | 505/ 704 | 505/ 705 | 505/ 706 | 505/ 707 | 505/ 708 | 505/ 709 | 505/ 710 | 505/ 711 | 505/ 712 |
| F | 506/ 701 | 506/ 702 | 506/ 703 | 506/ 704 | 506/ 705 | 506/ 706 | 506/ 707 | 506/ 708 | 506/ 709 | 506/ 710 | 506/ 711 | 506/ 712 |
| G | 507/ 701 | 507/ 702 | 507/ 703 | 507/ 704 | 507/ 705 | 507/ 706 | 507/ 707 | 507/ 708 | 507/ 709 | 507/ 710 | 507/ 711 | 507/ 712 |
| H | 508/ 701 | 508/ 702 | 508/ 703 | 508/ 704 | 508/ 705 | 508/ 706 | 508/ 707 | 508/ 708 | 508/ 709 | 508/ 710 | 508/ 711 | 508/ 712 |

Figure 8. QIAseq CDI Y-Adapter Plate (96) layout (CDI 1–96).

| | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 |
|---|---------|---------|---------|-------|-------|-------|-------|-------|-------|-------|-------|-------|
| A | 501/701 | 501/702 | 501/703 | empty | empty | empty | empty | empty | empty | empty | empty | empty |
| B | 502/701 | 502/702 | 502/703 | empty | empty | empty | empty | empty | empty | empty | empty | empty |
| C | 503/701 | 503/702 | 503/703 | empty | empty | empty | empty | empty | empty | empty | empty | empty |
| D | 504/701 | 504/702 | 504/703 | empty | empty | empty | empty | empty | empty | empty | empty | empty |
| E | 505/701 | 505/702 | 505/703 | empty | empty | empty | empty | empty | empty | empty | empty | empty |
| F | 506/701 | 506/702 | 506/703 | empty | empty | empty | empty | empty | empty | empty | empty | empty |
| G | 507/701 | 507/702 | 507/703 | empty | empty | empty | empty | empty | empty | empty | empty | empty |
| H | 508/701 | 508/702 | 508/703 | empty | empty | empty | empty | empty | empty | empty | empty | empty |

Figure 9. QIAseq CDI Y-Adapter Plate (24) layout (CDI 1–24).

Table 13. CDI motives used in the QIAseq CDI Y-Adapter Kits (24 and 96)

Note: Sequencing on the MiniSeq, NextSeq, HiSeqX, and HiSeq 3000/4000 systems follow a different dual-indexing workflow than other Illumina systems, which requires the reverse complement of the i5 index adapter sequence.

| Indices for entry on sample sheet | | | | |
|-----------------------------------|---|---|-------------------|------------------------------------|
| D50X barcode name | i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000) | i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000) | D50X barcode name | i7 bases for entry on sample sheet |
| D501 | TATAGCCT | AGGCTATA | D701 | ATTACTCG |
| D502 | ATAGAGGC | GCCTCTAT | D702 | TCCGGAGA |
| D503 | CCTATCCT | AGGATAGG | D703 | CGCTCATT |
| D504 | GGCTCTGA | TCAGAGCC | D704 | GAGATTCC |
| D505 | AGGCGAAG | CTTCGCCT | D705 | ATTCAGAA |
| D506 | TAATCTTA | TAAGATTA | D706 | GAATTCGT |
| D507 | CAGGACGT | ACGTCCTG | D707 | CTGAAGCT |
| D508 | GTACTGAC | GTCAGTAC | D708 | TAATGCGC |
| | | | D709 | CGGCTATG |
| | | | D710 | TCCGCGAA |
| | | | D711 | TCTCGCGC |
| | | | D712 | AGCGATAG |

Appendix D: Column-based Reaction Cleanup with the GeneRead Size Selection Kit

This protocol is optimized for the removal of primers and adapter-dimers from DNA libraries prepared from at least 100 ng of DNA using the QIAseq FX DNA Library Kits. This protocol may not remove all adapter dimer from libraries prepared from less than 100 ng of DNA.

Notes before starting

- All centrifugation steps should be performed at full speed (maximum 20,000 $\times g$) in a conventional, table-top centrifuge.
- Wash steps should be performed using 80% ethanol prepared from 96–100% ethanol.

Procedure

1. Add 4 volumes of Buffer SB1 to 1 volume of adapter-ligated DNA library prepared using the QIAseq FX DNA Library Kit, and mix. For example, add 360 μ l Buffer SB1 to a 90 μ l sample.
2. To bind DNA, apply the sample to the MinElute[®] spin column, and then centrifuge for 1 min. Discard the flow-through. Place the MinElute spin column back into the same tube.
3. To wash, add 700 μ l of 80% ethanol to the MinElute spin column and centrifuge for 1 min. Discard the flow-through. Place the MinElute spin column back into the same tube.
4. Repeat step 3.
5. Centrifuge the MinElute spin column for an additional 1 min at maximum speed.
6. Place the MinElute spin column into a clean 1.5 ml microcentrifuge tube (provided).

7. Add 90 μ l Buffer TE to the center of the membrane, let the column stand for 1 min, and then centrifuge for 1 min.

Important: Ensure that the buffer is dispensed directly onto the center of the membrane.

Important: Keep the spin column and the flow-through.

8. Place the MinElute spin column from step 7 into a new 2 ml collection tube (provided). Add 4 volumes of Buffer SB1 (~360 μ l) to 1 volume of the flow-through, and mix.

9. Reapply the mixture to the MinElute spin column and centrifuge for 1 min. Discard the flow-through.

10. To wash, add 700 μ l of 80% ethanol to the MinElute spin column and centrifuge for 1 min. Discard the flow-through. Place the MinElute spin column back into the same tube.

11. Repeat step 10.

12. Centrifuge the MinElute spin column for an additional 1 min at maximum speed.

13. Place the MinElute spin column in a clean 1.5 ml microcentrifuge tube (provided).

14. For elution, add 17 μ l Buffer EB to the center of the membrane, let the MinElute spin column stand for 1 min, and then centrifuge for 1 min.

Important: Ensure that the buffer is dispensed directly onto the center of the membrane, for complete elution of the bound DNA.

Appendix E: Library Quantification and Quality Control

Quality control for the library construction process can be performed using QIAGEN's QIAseq Library Quant Assay Kit (cat. no. 333314). With this assay, the correct dilution of the library can also be determined for sequencing. Refer to the corresponding handbook for library quantification and quality control.

Ordering Information

| Product | Contents | Cat. no. |
|--------------------------------------|---|-----------------|
| QIAseq FX DNA Library UDI-A Kit (96) | For 96 reactions: Buffers and reagents for DNA fragmentation, end-repair, A-addition, ligation, and library amplification; for use with Illumina instruments; includes a plate containing 96 adapters with different barcodes (pierceable foil seal allowing usage of defined parts of plate) | 180479 |
| QIAseq FX DNA Library UDI-B Kit (96) | For 96 reactions: Buffers and reagents for DNA fragmentation, end-repair, A-addition, ligation, and library amplification; for use with Illumina instruments; includes a plate containing 96 adapters with different barcodes (pierceable foil seal allowing usage of defined parts of plate) | 180480 |
| QIAseq FX DNA Library UDI-C Kit (96) | For 96 reactions: Buffers and reagents for DNA fragmentation, end-repair, A-addition, ligation, and library amplification; for use with Illumina instruments; includes a plate containing 96 adapters with different barcodes (pierceable foil seal allowing usage of defined parts of plate) | 180481 |
| QIAseq FX DNA Library UDI-D Kit (96) | For 96 reactions: Buffers and reagents for DNA fragmentation, end-repair, A-addition, ligation, and library amplification; for use with Illumina instruments; includes a plate containing 96 adapters with different barcodes (pierceable foil seal allowing usage of defined parts of plate) | 180482 |
| QIAseq FX DNA Library CDI Kit (96) | For 96 reactions: Buffers and reagents for DNA fragmentation, end-repair, A-addition, ligation, and library amplification; for use with Illumina instruments; includes a plate containing 96 adapters with different barcodes (pierceable foil seal allowing usage of defined parts of plate) | 180484 |

| Product | Contents | Cat. no. |
|---|---|----------|
| QIAseq FX DNA Library UDI Kit (24) | For 24 reactions: Buffers and reagents for DNA fragmentation, end-repair, A-addition, ligation, and library amplification; for use with Illumina instruments; includes a plate containing 24 adapters with different barcodes (pierceable foil seal allowing usage of defined parts of plate) | 180477 |
| Related products | | |
| QIAseq Y-Adapter Kits for Illumina | | |
| QIAseq CDI Y-Adapter Kit (24) | Combinatorial Dual-Index Adapters for Illumina | 180301 |
| QIAseq CDI Y-Adapter Kit (96) | Combinatorial Dual-Index Adapters for Illumina | 180303 |
| QIAseq UDI Y-Adapter Kit (24) | Unique Dual-Index Adapters for Illumina (1–24) | 180310 |
| QIAseq UDI Y-Adapter Kit A (96) | Unique Dual-Index Adapters for Illumina (1–96) | 180312 |
| QIAseq UDI Y-Adapter Kit B (96) | Unique Dual-Index Adapters for Illumina (97–192) | 180314 |
| QIAseq UDI Y-Adapter Kit C (96) | Unique Dual-Index Adapters for Illumina (193–288) | 180316 |
| QIAseq UDI Y-Adapter Kit D (96) | Unique Dual-Index Adapters for Illumina (289–384) | 180318 |
| QIAseq Library Quantification Kits for use with Illumina instruments | | |
| QIAseq Library Quant Assay Kit | Laboratory-verified forward and reverse primers for 500 x 25 µl reactions (500 µl); DNA Standard (100 µl); Dilution Buffer (30 ml); (1.35 ml x 5) GeneRead qPCR SYBR® Green Mastermix | 333314 |
| QIAamp Kits – for genomic DNA purification | | |
| QIAamp DNA Mini Kit | For 50 DNA preps: 50 QIAamp Mini Spin Columns, QIAGEN Proteinase K, Reagents, Buffers, Collection Tubes (2 ml) | 51304 |

| Product | Contents | Cat. no. |
|---|---|-----------------|
| QIAamp DNA Microbiome Kit | For 50 DNA preps: 50 QIAamp UCP Mini Columns, 50 Pathogen Lysis Tubes L, buffers, reagents, Collection Tubes (2 ml) | 51704 |
| MagAttract Kits – for high-molecular-weight genomic DNA purification | | |
| MagAttract HMW DNA Kit (48) | For 48 DNA preps: MagAttract Suspension G, Buffer ATL, Buffer AL, Buffer MB, Buffer MW1, Buffer PE, Proteinase K, RNase A, Buffer AE, Nuclease-Free Water | 67563 |
| MagAttract Magnetic Rack | Magnetic rack for convenient processing of up to 12 samples | 19606 |
| REPLI-g® Kits – for MDA-based whole genome amplification | | |
| REPLI-g Mini Kit (100)* | DNA Polymerase, Buffers, and Reagents for 100 x 50 µl whole genome amplification reactions (typical yield 10 µg per reaction) | 150025 |
| REPLI-g Single Cell Kit (96)* | REPLI-g sc Polymerase, Buffers, and Reagents for 96 whole genome amplification reactions (yields up to 40 µg/reaction) | 150345 |
| REPLI-g Mitochondrial DNA Kit (25) | DNA Polymerase, Buffers, and Reagents for 25 x 50 µl mitochondrial DNA specific whole genome amplification reactions | 151023 |

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* Other kit sizes available; see www.qiagen.com.

Document Revision History

| Date | Changes |
|---------|--|
| 01/2020 | <p>Deleted Adapter Plate Illumina from "Kit Contents". Added information about QIAseq Y-adapter kits and NGS adapter and index technologies. Removed information exclusive to formerly recommended adapters. Updated information related to Illumina instruments.</p> <p>Updated Table 1 to add another fragment insert size. Updated headings on Table 2 and Table 9 from 1 ng to 20 pg input and incubation time from 3–20 min to 1–30 min. Revision of measurement quantity symbols in this document. Added new table (Table 5) for adapter dilution factors. Updated the "Number of cycles" column and number of recommended amplification cycles in Table 7. Updated the "Volume" column in tables 10 and 11.</p> |
| 01/2021 | <p>Fixed a typo in Figure 9. Changed precise storage temperatures to range temperatures. Added a statement in the Principle and Procedure section to clarify input range. Corrected the DNA input range in for adapter dilution 1:1000 in Table 5. Increased the number of times to pipette up and down from "6–8" to "20" times in step 4 of Protocol: Fragmentation, End-Repair, and A addition and step 3 of Appendix B.</p> |

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