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August 2021

# QIAseq<sup>®</sup> Ultralow Input Library Kit Handbook

For preparation of DNA libraries for  
next-generation sequencing (NGS)  
applications that use Illumina<sup>®</sup> instruments

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# Kit Contents

<b>QIaseq Ultralow Input Library Kit (12)</b>	<b>(12)</b>
<b>Catalog no.</b>	<b>180492</b>
<b>No. of reactions</b>	<b>12</b>
End-Polishing Enzyme Mix (violet cap)	1 tube
End-Polishing Buffer, 10x (blue cap)	1 tube
Ultralow Input Ligase (orange cap)	1 tube
Ultralow Input Ligation Buffer, 4x (yellow cap)	1 tube
HiFi PCR Master Mix, 2x (green cap)	1 tube
Primer Mix Illumina Libr. Amp (clear cap)	1 tube
RNase-free Water (1.9 ml) (clear cap)	1 tube
Quick-Start Protocol	1

<b>QIaseq Ultralow Input Lib CDI/UDI Kit</b>	<b>CDI (96)</b>	<b>UDI-A (96)</b>	<b>UDI-B (96)</b>	<b>UDI-C (96)</b>	<b>UDI-D (96)</b>
<b>Catalog no.</b>	<b>180501</b>	<b>180497</b>	<b>180498</b>	<b>180499</b>	<b>180500</b>
<b>No. of reactions</b>	<b>96</b>	<b>96</b>	<b>96</b>	<b>96</b>	<b>96</b>
End-Polishing Enzyme Mix (violet cap)	1 tube	1 tube	1 tube	1 tube	1 tube
End-Polishing Buffer, 10x (blue cap)	1 tube	1 tube	1 tube	1 tube	1 tube
Ultralow Input Ligase (orange cap)	1 tube	1 tube	1 tube	1 tube	1 tube
Ultralow Input Ligation Buffer, 4x (yellow cap)	3 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 Libr. Amp (clear cap)	1 tube	1 tube	1 tube	1 tube	1 tube
RNase-free Water (1.9 ml) (clear cap)	4 tubes	4 tubes	4 tubes	4 tubes	4 tubes
QIaseq CDI Y-Adapter Plate (96)	1	–	–	–	–
QIaseq UDI Y-Adapter Plate A,B,C, or D (96)	–	1	1	1	1
QIaseq Y-Adapter Reference Card	1	1	1	1	1
Quick-Start Protocol	1	1	1	1	1

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

The QIAseq Ultralow Input Library Kits (96) 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 Ultralow Input Lib UDI-A (or B or C or D) Kit (96) 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 28.

For the QIAseq Ultralow Input Library Kit (12), adapters are available in tube format: The GeneRead™ Adapter I Set A/B 12-plex (cat. 180985, 180986) can be ordered separately. The QIAseq Ultralow Input Library Kit (12) is also fully compatible with all plate format QIAseq Y-Adapter Kits (24/96).

## Shipping and Storage

The QIAseq Ultralow Input Library Kit is shipped on dry ice and should be stored immediately upon receipt at  $-30$  to  $-15^{\circ}\text{C}$  in a constant-temperature freezer. QIAseq Ultralow Input Library Kits (96) contain an adapter plate shipped in a separate box. Store the adapter plate at  $-30$  to  $-15^{\circ}\text{C}$  upon arrival. When stored correctly, all reagents are stable for at least 6 months after delivery if not otherwise stated on the label.

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## Intended Use

QIAseq Ultralow Input Library Kits are intended for molecular biology applications. These products are 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](http://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 QIAseq Ultralow Input Library Kit is tested against predetermined specifications to ensure consistent product quality.

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# Introduction

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 a process bottleneck for many labs and a limiting factor in the types of samples that can successfully generate NGS data.

QIAseq Ultralow Input Library Kits have been designed to be the definitive solution for generating high-quality libraries from even very challenging NGS samples. Intended for NGS researchers who seek a single library prep kit compatible with a wide range of ultralow-, low-, and standard-input fragmented DNA (including circulating cell-free DNA, fragmented DNA, ancient DNA, and chromatin immunoprecipitation [ChIP] DNA), QIAseq Ultralow Input Library Kits enable new insights by maximizing performance particularly from limited and damaged DNA sample types. The streamlined 2.5-hour protocol for generating libraries from fragmented DNA using QIAseq Ultralow Input Library Kits also enables straightforward automation on different liquid-handling platforms.

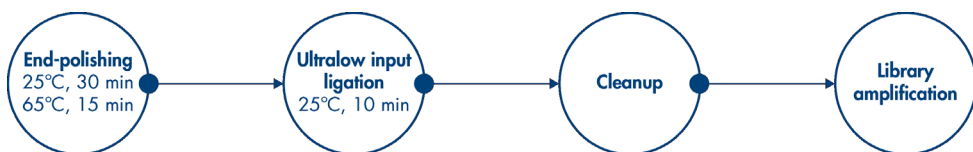
## Principle and procedure

Starting from enzymatically, chemically, mechanically, or naturally fragmented double-stranded DNA, QIAseq Ultralow Input Library Kits use an optimized end-polishing reaction and a new Ultralow Input Ligation formulation along with QIAGEN's proprietary HiFi PCR Master Mix. This combination maximizes the conversion rate of sample DNA into NGS library while efficiently and evenly amplifying even high- and low-GC-content regions of the genome. This protocol enables the highest possible yield sequencing library free of adapter-dimer contamination starting from as little as 10 pg of DNA input. Due to the kit's flexible protocol, the same kit can also be used for higher DNA input amounts of up to 100 ng, including PCR-free library preparation from as little as 10 ng DNA input.

## Important note

Note that the genomic complexity necessary for a given experiment will vary depending on the genome size of the organism and the fraction of that genome included in the target region of interest. For example, 100 pg of DNA represents approximately 30 unique copies of the human haploid genome, but over 5000 copies of a small genome such as *Escherichia coli* (*E. coli*). A conversion rate of 0.3 from sample to library can be applied to these calculations to roughly estimate the maximum genomic complexity of the expected NGS data. For most whole-genome sequencing experiments from human or other organisms with haploid genome sizes over ~1 gigabase, QIAGEN recommends starting the QIAseq Ultralow Input protocol from at least 100 pg of input DNA.

The QIAseq Ultralow Input Library Kit, with its innovative buffer and enzyme formulations, provides an optimized solution to efficiently construct Illumina libraries from as little as 10 pg input DNA. Following adapter ligation and library amplification steps, reaction cleanup and removal of residual adapter-dimers can be achieved by using Agencourt® AMPure® XP beads, which enable easy automation on various high-throughput automation platforms.



**Figure 1. QIAseq Ultralow Input Library Kit Workflow.** Starting from enzymatically, chemically, mechanically or naturally fragmented double-stranded DNA, QIAseq Ultralow Input Library Kits use an optimized end-polishing reaction and a new Ultralow Input Ligation formulation along with QIAGEN's proprietary HiFi PCR Master Mix.

## 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



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each single read to its original source library. This is facilitated by the integration of index sequences into the individual adapter molecules.

QIAseq Ultralow Input Library Kits (96) include a fully compatible indexing solution. We recommend using the QIAseq Dual-Index Y-Adapter Plates delivered with the kit. Each QIAseq Ultralow Input Library Kits (96) includes one of the following:

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

Combining QIAseq Ultralow Input Lib 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 28, and “Ordering Information”, page 58.

Adapter kits compatible with the QIAseq Ultralow Input Library Kit (12) have to be purchased separately:

- GeneRead Adapter I Set A/B 12-plex (single-index adapters)

GeneRead Single-Index adapter sets A and B are available in 12-plex format and can be combined with the QIAseq 1-Step Amplicon Library Kit (12). For more information on GeneRead Adapters and index sequence motives, see Appendix E, page 56, and “Ordering Information”, page 58.

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.

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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.

## Automated library construction

Due to the streamlined protocol and magnetic bead-based purification, QIAseq Ultralow Input library preparation protocols can be easily automated on most commonly used liquid handling platforms.

## Description of protocols

This handbook contains two protocols for generation of DNA libraries that are for use on NGS platforms from Illumina. The first protocol (page 16) describes end-polishing, adapter ligation and cleanup of DNA – to generate libraries that are ready to quantify and use in NGS. The second protocol (page 21) describes an optional, high-fidelity amplification step that can be used to ensure sufficient library for sequencing from as little as 10 pg of starting material.

## Compatible sequencing platforms

- Illumina NovaSeq®
- Illumina MiniSeq®
- Illumina MiSeq®
- Illumina NextSeq®
- Illumina HiSeq®

## Starting materials

- Fragmented genomic DNA
- Fragmented FFPE DNA

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- Circulating cell free DNA (cfDNA)
  - Ancient DNA (naturally fragmented)
  - DNA from CHIP
  - Fragmented double-stranded cDNA
  - Fragmented REPLI-g®-amplified DNA

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# 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.

- Only for 12 reaction kits: GeneRead Adapters (cat. no. 180985 or 180986; can be used with QIAseq Ultralow Input Library Kit [12] for single-indexed libraries)
- Enzymatic, mechanical (e.g., Covaris® instrument) or other method to fragment high molecular weight gDNA samples
- Agencourt® AMPure® XP Beads (Beckman Coulter Inc., 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
- Vortexer
- Microcentrifuge
- Thermocycler
- Magnetic stand (e.g., DynaMag™-2 Magnet or DynaMag™-96 Side Skirted Magnet, Thermo Fisher, cat. no. 12027)
- QIAGEN QIAxcel®, Agilent® Bioanalyzer® or similar method to assess the quality of DNA library
- qPCR instrument and QIAseq Library Quant Assay Kit (cat. no. 333314) or a similar method for qPCR-based library quantitation

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# Important Notes

## DNA preparation and quality control

High-quality DNA is always the best choice for obtaining reliable sequencing results. Sample-handling and DNA-isolation procedures are also critical to the success of the experiment and maximizing the performance of real-world samples. 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.

## Recommended DNA preparation method

We recommend the following QIAGEN kits:

- QIAamp® DNA Mini Kit (cat. no. 51304) for the preparation of genomic DNA samples from fresh tissues and cells
- QIAamp DNA Micro Kit (cat. no. 56304) for purification of genomic and mitochondrial DNA from small samples
- QIAamp Circulating Nucleic Acid Kit (cat. no. 55114) for isolation of free-circulating DNA and RNA from human plasma or serum
- GeneRead DNA FFPE Kit (cat. no. 180134) for the preparation of NGS-ready genomic DNA from FFPE tissue samples
- 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

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## Recommendations for DNA fragmentation

DNA can be fragmented using one the following methods:

- Acoustic shearing (e.g., Covaris Adaptive Focused Acoustics® (AFA) technology)
- Nebulization
- Sonication
- Enzymatic methods

To ensure complete fragmentation of the DNA that is needed for library preparation, follow the recommended parameters given in the manufacturer's instructions. Using too much DNA in a Covaris instrument may, for example, lead to incomplete shearing of the DNA. Check the fragmented DNA for the correct size distribution using an agarose gel or capillary electrophoresis device.

We recommend the QIAxpert® System for convenient quantitation of higher-concentration dsDNA samples. For DNA samples at low concentrations or containing both dsDNA and ssDNA – such as FFPE DNA or cfDNA – we recommend Qubit®, PicoGreen® or another fluorometric method to accurately quantitate input DNA.

## Recommended library quantification method

We recommend final library quantification by qPCR using primers complementary to the platform-specific adapters. This allows measurement of only complete library molecules, which are the only molecules able to perform bridge PCR on Illumina instruments. The QIAseq Library Quant Array Kit (cat. no. 333304) is highly recommended for accurate qPCR quantification of the prepared library. The QIAseq Library Quant Array Kit is compatible with all major NGS platforms and qPCR instruments, and includes predispensed, sequentially diluted DNA standard to eliminate manual titration steps.

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For the most accurate qPCR analysis, the library fragment size distribution should be as narrow as possible, with a known mean value in base pairs. Wide fragment distributions are more difficult to accurately quantitate by qPCR, and the smallest library fragments will be overrepresented in NGS data due to their higher efficiency of amplification during bridge PCR.

### Important points before starting

- This protocol is for constructing sequencing libraries for Illumina NGS platforms.
- QIAseq and GeneRead adapters are dissolved in duplex buffer and are ready to use.
- Adapters are fully compatible with all Illumina instruments, including MiniSeq, MiSeq, NextSeq, HiSeq, and NovaSeq instruments. A PCR step is not required to complete the adapter sequences – they are full length and ready for sequencing following the ligation step.
- The protocol can be used with 10 pg–100 ng double-stranded DNA with suitable fragment sizes for Illumina sequencing. The ideal fragment size will depend on the application and read length. Please refer to the recommendations from the sequencer manufacturer.
- The majority of cfDNA fragments are approximately 170 bp and do not require further fragmentation prior to library preparation.
- Ancient DNA is normally fragmented and does not require further fragmentation prior to library preparation.
- Do not use a heated lid during the adapter ligation step.

# Protocol: End-Polishing and Ultralow Input Ligation

This protocol describes end repair, A-addition, adapter ligation, and library cleanup. It generates libraries that are ready to quantify and use in NGS on instruments from Illumina.

## Things to do before starting

- Fragment high-molecular-weight DNA before starting this protocol, using either an enzymatic method or a physical method (e.g., DNA with specific median fragment length sizes can be prepared using a Covaris instrument, according to the manufacturer's instructions).
- Sample DNA should be dissolved EB/Tris buffer or water before starting.
- Thaw frozen reagents on ice. Once thawed, buffers should be mixed thoroughly by quick vortexing to avoid any localized concentrations. Briefly spin down vortexed reagents before use. Do not vortex enzymes.
- Prepare fresh 80% ethanol

## Procedure

1. Program a thermocycler according to Table 1.

**Table 1. Thermal cycling parameters**

Step	Time	Temperature	Additional comments
End-polishing	30 min	25°C	End repair
	15 min	65°C	Inactivation of end-repair enzymes; A-addition
	∞	4°C	Hold
Ligation	10 min	25°C	Ligation of the adapters to the prepared DNA fragments
	∞	4°C	Hold



2. Prepare the End-Polishing Reaction Mix according to Table 2, adding the components to the PCR tube or plate containing DNA fragments.

**Note:** The reaction mix should be prepared on ice.

3. Mix components by pipetting up and down several times, and then transfer to the thermocycler for end polishing at 25°C, followed by incubation at 65°C to inactivate the End-Polishing Enzyme Mix.

**Table 2. End-Polishing Reaction Mix setup (per sample) for 10 pg – 100 ng input DNA**

Component	Volume/reaction (µl)
10 pg – 100 ng Sample DNA (fragmented)	Variable
End-Polishing Buffer, 10x	5
End-Polishing Enzyme Mix	2
Nuclease-free Water	Variable
<b>Total reaction volume</b>	<b>50</b>

4. During the end-polishing step, prepare and dilute adapters according to the instructions in Appendix D (page 53) or Appendix E (page 56) and Table 3.

**Table 3. Adapter dilution factors**

Sample DNA Amount	QIaseq or GeneRead Adapters (Plate or Set)
10–99 pg	1:1000 dilution
100–999 pg	1:100 dilution
1–9 ng	1:10 dilution
10– 100 ng	No dilution

**Note:** See Appendix D or Appendix E.

5. After the end-polishing steps are complete, remove the reaction tubes from the thermocycler and place them on ice.

- Proceed immediately to Ultralow Input Adapter Ligation. Add components for the adapter-ligation reaction to end-polished samples according to Table 4.

**Table 4. Adapter ligation setup (per sample) for end-polished DNA**

Component	Volume/reaction (µl)
End-Polished DNA	50
Ultralow Input Ligation Buffer, 4X	25
Ultralow Input Ligase	5
QIAseq or GeneRead Adapter, prepared and diluted adapter; see Table 3 and Appendix D or Appendix E	2
DNase-free Water	18
<b>Total reaction volume</b>	<b>100</b>

- Transfer reactions to the thermocycler to incubate at 25°C for 10 min. Do not use a heated lid.

**Important:** Do not attempt to use diluted adapters more than once, due to the risk of barcode cross-contamination and lower than expected adapter concentration after storage of very dilute oligos.

- Once ligation is complete, place the reactions on ice and proceed with purification using Agencourt AMPure XP beads.
- Add 80 µl resuspended Agencourt AMPure XP beads to each sample and mix well by pipetting.
- Incubate the mixture for 5 min at room temperature (15–25°C).
- Pellet the beads on a magnetic stand (e.g., DynaMag) for 2 min, and then carefully discard the supernatant.

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12. Wash the beads by adding 200  $\mu$ l fresh 80% ethanol to each pellet. Pellet the beads on the magnetic stand and carefully discard the supernatant.
  13. Repeat step 13 for a total of 2 ethanol washes. Remove as much excess ethanol as possible.
  14. Incubate on the magnetic stand for 5–10 min or until the beads are dry. Overdrying of the beads may result in lower DNA recovery. Remove from the magnetic stand.
  15. Elute by resuspending in 52.5  $\mu$ l QIAGEN Buffer EB or 10 mM Tris-Cl, pH 8.0. Pellet beads on the magnetic stand. Carefully transfer 50  $\mu$ l supernatant to a new plate or tube.
  16. Perform a second purification using 1x AMPure beads: Add 50  $\mu$ l resuspended Agencourt AMPure XP beads to each sample and mix well by pipetting.
  17. Repeat steps 11–15. Elute by resuspending in 26  $\mu$ l QIAGEN Buffer EB or 10 mM Tris-Cl, pH 8.0. Pellet the beads on the magnetic stand. Carefully transfer 23.5  $\mu$ l of supernatant into a new PCR plate or tube for QC and storage or amplification.
  18. Assess the quality of the library using a capillary electrophoresis device or comparable method. Check for the correct size distribution of library fragments and for the absence of adapters or adapter-dimers.
  19. If the library will not be amplified, store the DNA at  $-20^{\circ}\text{C}$  in a DNA LoBind tube until ready for sequencing. If amplifying the library, proceed to the protocol “Amplification of Library DNA”).

**Note:** Libraries generated from  $<10$  ng input DNA may not be visible by capillary electrophoresis prior to library amplification.

# Protocol: Amplification of Library DNA

PCR-based library amplification is normally required if the input DNA amount is <100 ng for genomic DNA or <10 ng for cfDNA. This protocol is for optional, high-fidelity amplification of the DNA library using the QIAseq HiFi PCR Master Mix included in the kit.

## Things to do before starting

- Prepare library DNA using the protocol “End-Polishing and Ultralow Input Ligation,” page 16.
- Thaw frozen reagents on ice. Once thawed, buffers should be mixed thoroughly by quick vortexing to avoid any localized concentrations. Briefly spin down vortexed reagents before use. Do not vortex enzymes.

## Procedure

1. Program a thermocycler according to Table 5.

**Note:** Use the minimum number of cycles necessary to generate sufficient library yield. We suggest a starting point of 16 cycles for libraries created from 10 pg DNA, 14 cycles for 100 pg, 10 cycles for 1 ng input, and 8 cycles for 10 ng.

**Table 5. Cycling conditions**

Time	Temperature	Number of cycles
2 min	98°C	1
20 s	98°C	
30 s	60°C	Variable
30 s	72°C	
1 min	72°C	1
∞	4°C	Hold

2. Prepare the library amplification reaction mix on ice according to Table 6.

**Table 6. Reaction mix for library amplification**

Component	Volume/reaction (µl)
HiFi PCR Master Mix, 2x	25
Primer Mix (10 µM each)	1.5
Library DNA (from step 19, page 19)	23.5
<b>Total reaction volume</b>	<b>50</b>

3. Transfer the PCR plate to the thermocycler and start the program.
4. When the program is complete, add 50 µl of resuspended Agencourt AMPure XP beads to each 50 µl PCR sample. Mix well by pipetting up and down several times.
5. Incubate the mixture at 5 min at room temperature.
6. Pellet the beads on a magnetic stand (e.g., DynaMag) for 2 min, then carefully discard the supernatant.
7. Wash the beads by adding 200 µl fresh 80% ethanol to each pellet. Pellet the beads on the magnetic stand and carefully discard the supernatant.
8. Repeat step 27 for a total of 2 ethanol washes. Remove as much excess ethanol as possible.
9. Incubate on the magnetic stand for 5–10 min or until the beads are dry. Overdrying of the beads may result in lower DNA recovery. Remove from the magnetic stand.
10. Elute by resuspending in 25 µl nuclease-free water or QIAGEN Buffer EB. Pellet the beads on a magnetic stand. Carefully transfer 23 µl supernatant into a new LoBind tube.
11. Store libraries at –20°C in a DNA Lo-bind tube until ready for QC, library quantification, and sequencing.

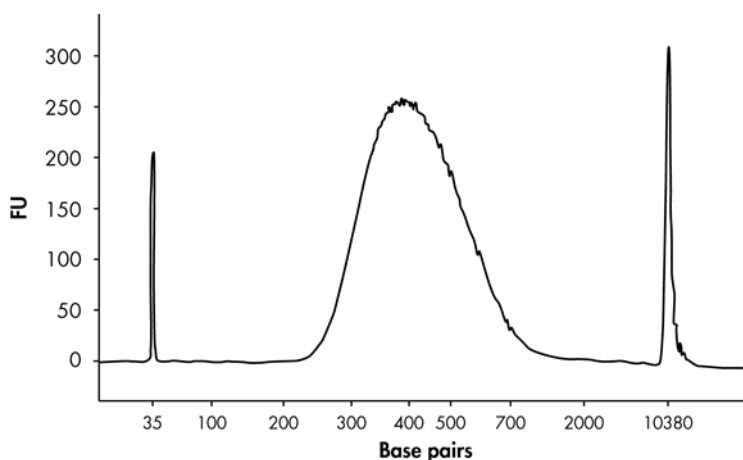
## Library QC and quantification

1. Assess the quality of the library using a capillary electrophoresis device or comparable method. Check for the correct size distribution of library fragments and for the absence of adapters or adapter-dimers.

**Note:** The median size of the DNA fragment should be shifted by the size of the adapters that were ligated to the library fragments (e.g., for QIAGEN adapters, add 120 bp). For PCR-free libraries, be aware of the following:

- The adaptors contain noncomplementary regions prior to PCR amplification. This makes the DNA library migrate slower on the capillary electrophoresis than completely complementary dsDNA.
- In addition to insert molecules with adaptors on both ends, the ligation product may also contain unligated DNA fragments as well as DNA fragments ligated with only one adaptor.

**Note:** The median fragment size can be used for subsequent qPCR-based quantification methods.



**Figure 2.** Capillary electrophoresis device trace data showing the correct size distribution of library fragments and the absence of adapters or adapter-dimers.

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2. Quantify the library using the QIAseq Library Quant Array Kit or a comparable qPCR-based method.
  3. The purified library can be safely stored at  $-20^{\circ}\text{C}$  in DNA LoBind tubes until downstream hybrid capture or sequencing.

# 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](http://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](http://www.qiagen.com)).

## Comments and suggestions

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### Low library yields

- |  |   |
|--|---|
| a) Suboptimal reaction conditions due to low DNA quality                                       | Make sure to use the highest quality sample DNA available to ensure optimal activity of the library enzymes.  |
| b) Insufficient amount of starting DNA for direct sequencing without library amplification     | Typically, 10 ng of cfDNA or 100 ng of fragmented genomic DNA generates enough Illumina-compatible library to use directly for sequencing without amplification. Make sure to use an accurate DNA quantification method (e.g., QIAxpert® or a fluorometric method) to ensure sufficient DNA input. If the final library yield is not sufficient for the expected number of sequencing runs or applications (e.g., hybrid capture), a library amplification step can be performed. |
| c) Insufficient amount of starting DNA due to inaccurate DNA quantification                    | RNA from the sample material can be copurified with genomic DNA. This contaminating RNA will affect the accuracy of DNA quantification. To remove RNA during the sample preparation protocol, it is recommended to perform RNase A treatment of the DNA. For low-concentration DNA samples and FFPE samples, use fluorescence-based methods such as PicoGreen measurement to accurately and sensitively quantify double-stranded DNA.   |
| d) Overdrying of the Ampure beads during cleanup steps   | Overdrying of the Ampure beads can make it difficult to elute the DNA off the beads. Do not dry beads for more than 10 min at room temperature.   |
| e) Wrong reaction volumes or conditions used for end polishing, ligation, or PCR amplification | Make sure to use the exact conditions in the protocol. Make sure not to use heated lid for ligation.  |



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### Comments and suggestions

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|----|--|--|
| f) | Lower than expected library yield with extensive adapter-dimer | Excessive adapter-dimers can compete with the amplification of the real libraries during library amplification PCR step and lead to low yield of specific libraries. Make sure the right dilution of the adapter is used. Make sure the correct AMPure bead purification protocol is used. |
|----|--|--|

### 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 Illumina flow cell and will be sequenced, this will reduce the capacity of the flow cell for the library fragments. A low ratio of adapter-dimers versus library will not be a problem. Make sure the correct dilution of the adapters and the correct volume of the AMPure beads are used for the cleanup steps. |
| b) | Presence of larger library fragments after library enrichment    | If the fragment population shifts higher than expected after adapter ligation and library amplification (e.g., more than the expected 120 bp shift), this can be a PCR artifact due to overamplification of the DNA library. Use as few amplification cycles as possible to avoid this effect.  |
| c) | Presence of larger library fragments after library enrichment    | If the fragment population shifts higher than expected, this can also be due to the carryover of the AMPure beads. Make sure not to aspirate beads while taking supernatant during the cleanup steps.   |
| d) | Presence of larger library fragments prior to library enrichment | 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, please refer to the size information given in the respective documentation.  |

# Appendix A: Adapter Indices for the GeneRead Adapter I Set A 12-Plex

The index sequences used in the GeneRead Adapter I set 12-plex are listed in Table 7. Indexes 1–12 correspond to the respective Illumina adapter indexes.

**Table 7. Adapter indices**

<b>Adapter name</b>	<b>Indices</b>
Adapter Bc1 Illumina	ATCACG
Adapter Bc2 Illumina	CGATGT
Adapter Bc3 Illumina	TTAGGC
Adapter Bc4 Illumina	TGACCA
Adapter Bc5 Illumina	ACAGTG
Adapter Bc6 Illumina	GCCAAT
Adapter Bc7 Illumina	CAGATC
Adapter Bc8 Illumina	ACTTGA
Adapter Bc9 Illumina	GATCAG
Adapter Bc10 Illumina	TAGCTT
Adapter Bc11 Illumina	GGCTAC
Adapter Bc12 Illumina	CTTGTA

## Appendix B: Adapter Indices for the GeneRead Adapter I Set B 12-Plex

The index sequences used in the GeneRead Adapter I set 12-plex B are listed in Table 8. Indices 13–16, 18–23, 25, and 27 correspond to the respective Illumina adapter indices.

**Table 8. Adapter indices**

<b>Adapter name</b>	<b>Indices</b>
Adapter Bc13 Illumina	AGTCAA
Adapter Bc14 Illumina	AGTCC
Adapter Bc15 Illumina	ATGTCA
Adapter Bc16 Illumina	CCGTCC
Adapter Bc18 Illumina	GTCCGC
Adapter Bc19 Illumina	GTGAAA
Adapter Bc20 Illumina	GTGGCC
Adapter Bc21 Illumina	GTTTCG
Adapter Bc22 Illumina	CGTACG
Adapter Bc23 Illumina	GAGTGG
Adapter Bc25 Illumina	ACTGAT
Adapter Bc27 Illumina	ATTCT

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# 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](http://www.qiagen.com). Sequencing on the NextSeq, HiSeq X<sup>®</sup>, 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 1 to Figure 5. The index motives used in the QIAseq Unique Dual-Index Kits are listed in Table 9. To make sequencing preparation more convenient, you can download Illumina-compatible sample sheets for different sequencing instruments at [www.qiagen.com](http://www.qiagen.com).

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

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

	1	2	3	4	5	6	7	8	9	10	11	12
<b>A</b>	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>B</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
<b>C</b>	UDI 003	UDI 011	UDI 019	UDI 027	UDI 035	UDI 043	UDI 051	UDI 059	UDI 067	UDI 075	UDI 083	UDI 091
<b>D</b>	UDI 004	UDI 012	UDI 020	UDI 028	UDI 036	UDI 044	UDI 052	UDI 060	UDI 068	UDI 076	UDI 084	UDI 092
<b>E</b>	UDI 005	UDI 013	UDI 021	UDI 029	UDI 037	UDI 045	UDI 053	UDI 061	UDI 069	UDI 077	UDI 085	UDI 093
<b>F</b>	UDI 006	UDI 014	UDI 022	UDI 030	UDI 038	UDI 046	UDI 054	UDI 062	UDI 070	UDI 078	UDI 086	UDI 094
<b>G</b>	UDI 007	UDI 015	UDI 023	UDI 031	UDI 039	UDI 047	UDI 055	UDI 063	UDI 071	UDI 079	UDI 087	UDI 095
<b>H</b>	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 2. QIAseq UDI Y-Adapter Plate A (96) layout (UDI 1–96).

	1	2	3	4	5	6	7	8	9	10	11	12
<b>A</b>	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>B</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
<b>C</b>	UDI 099	UDI 107	UDI 115	UDI 123	UDI 131	UDI 139	UDI 147	UDI 155	UDI 163	UDI 171	UDI 179	UDI 187
<b>D</b>	UDI 100	UDI 108	UDI 116	UDI 124	UDI 132	UDI 140	UDI 148	UDI 156	UDI 164	UDI 172	UDI 180	UDI 188
<b>E</b>	UDI 101	UDI 109	UDI 117	UDI 125	UDI 133	UDI 141	UDI 149	UDI 157	UDI 165	UDI 173	UDI 181	UDI 189
<b>F</b>	UDI 102	UDI 110	UDI 118	UDI 126	UDI 134	UDI 142	UDI 150	UDI 158	UDI 166	UDI 174	UDI 182	UDI 190
<b>G</b>	UDI 103	UDI 111	UDI 119	UDI 127	UDI 135	UDI 143	UDI 151	UDI 159	UDI 167	UDI 175	UDI 183	UDI 191
<b>H</b>	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 3. QIAseq UDI Y-Adapter Plate B (96) layout (UDI 97–192).

	1	2	3	4	5	6	7	8	9	10	11	12
<b>A</b>	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>B</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
<b>C</b>	UDI 195	UDI 203	UDI 211	UDI 219	UDI 227	UDI 235	UDI 243	UDI 251	UDI 259	UDI 267	UDI 275	UDI 283
<b>D</b>	UDI 196	UDI 204	UDI 212	UDI 220	UDI 228	UDI 236	UDI 244	UDI 252	UDI 260	UDI 268	UDI 276	UDI 284
<b>E</b>	UDI 197	UDI 205	UDI 213	UDI 221	UDI 229	UDI 237	UDI 245	UDI 253	UDI 261	UDI 269	UDI 277	UDI 285
<b>F</b>	UDI 198	UDI 206	UDI 214	UDI 222	UDI 230	UDI 238	UDI 246	UDI 254	UDI 262	UDI 270	UDI 278	UDI 286
<b>G</b>	UDI 199	UDI 207	UDI 215	UDI 223	UDI 231	UDI 239	UDI 247	UDI 255	UDI 263	UDI 271	UDI 279	UDI 287
<b>H</b>	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 4. QIAseq UDI Y-Adapter Plate C (96) layout (UDI 193–288).

	1	2	3	4	5	6	7	8	9	10	11	12
<b>A</b>	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>B</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
<b>C</b>	UDI 291	UDI 299	UDI 307	UDI 315	UDI 323	UDI 331	UDI 339	UDI 347	UDI 355	UDI 363	UDI 371	UDI 379
<b>D</b>	UDI 292	UDI 300	UDI 308	UDI 316	UDI 324	UDI 332	UDI 340	UDI 348	UDI 356	UDI 364	UDI 372	UDI 380
<b>E</b>	UDI 293	UDI 301	UDI 309	UDI 317	UDI 325	UDI 333	UDI 341	UDI 349	UDI 357	UDI 365	UDI 373	UDI 381
<b>F</b>	UDI 294	UDI 302	UDI 310	UDI 318	UDI 326	UDI 334	UDI 342	UDI 350	UDI 358	UDI 366	UDI 374	UDI 382
<b>G</b>	UDI 295	UDI 303	UDI 311	UDI 319	UDI 327	UDI 335	UDI 343	UDI 351	UDI 359	UDI 367	UDI 375	UDI 383
<b>H</b>	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 5. QIAseq UDI Y-Adapter Plate D (96) layout (UDI 289–384).

**Table 9. 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	TTGTGTTCT	TTATCGGCCT
UDI 006	CTCAGTAGGC	GCCTACTGAG	GAGGTATAAG
UDI 007	GAAGTGCCTG	CAGGCACTTC	TCAAGGATTC
UDI 008	TCTCTCGCCT	AGGCGAGAGA	CGAACCGAGA
UDI 009	AGGCACCTTC	GAAGGTGCCT	GAGCCAAGTT
UDI 010	CTGTTGGTAA	TTACCAACAG	AAGCCGTAG
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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<b>Indices for entry on sample sheet</b>			
<b>Unique dual-index number</b>	<b>i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000)</b>	<b>i5 bases for entry on sample sheet (iSeq 100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000)</b>	<b>i7 bases for entry on sample sheet (all instruments)</b>
UDI 021	AGGCCGTTAGG	CCTAACGCCT	ATTGTGCCTT
UDI 022	CCGTAACGTC	GACGTTACGG	TCCTCTACCG
UDI 023	GTAATAGCCA	TGGCTATTAC	TACCATGAAC
UDI 024	TAGCGCCGAT	ATCGGCGCTA	CATTGGCAGA
UDI 025	CATTCTTGA	TCCAAGAATG	CACTGCTATT
UDI 026	ATGCAAGGT	AACCTGCAT	AATGGTAGGT
UDI 027	CGCCAGACAA	TTGTCTGGCG	GATACCTATG
UDI 028	GAAGGTTGGC	GCCAACCTTC	CACTAGGTAC
UDI 029	TCGCATCACG	CGTGATGCGA	AGCTCGTTCA
UDI 030	CCGTCATGA	TCATGACCGG	TGTCAGTCTT
UDI 031	ATTCACAAGC	GCTTGTAAT	GATGAACAGT
UDI 032	CAACCTGTAA	TTACAGGTTG	ACAATCGGCG
UDI 033	GCCAGTCGT	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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<b>Indices for entry on sample sheet</b>			
<b>Unique dual-index number</b>	<b>i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000)</b>	<b>i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000)</b>	<b>i7 bases for entry on sample sheet (all instruments)</b>
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	CCTGGCTTC	AGTACAGTTC
UDI 054	TGGAGTCAA	TTGAACTCCA	TACAGCCTCA
UDI 055	CTTCAATCCT	AGGATTGAAG	GTTCTATTGG
UDI 056	ATCTTGCGTG	CACGCAAGAT	ATATACCGGT
UDI 057	CGCTAAGGT	ACCTTAGACG	CCTCGGAATG
UDI 058	GAGGTGAACA	TGTTACCTC	GTTCTGGAAC
UDI 059	TCAGAACTAC	GTAGTTCTGA	AGATTCACCA
UDI 060	CGGATATTGA	TCAATATCCG	TCGGTCAGAT

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<b>Indices for entry on sample sheet</b>			
<b>Unique dual-index number</b>	<b>i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000)</b>	<b>i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000)</b>	<b>i7 bases for entry on sample sheet (all instruments)</b>
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	CAGCGTTGTT	GAACAATCCT
UDI 067	CGCGCTATT	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	GTTGCGCGAA	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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<b>Indices for entry on sample sheet</b>			
<b>Unique dual-index number</b>	<b>i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000)</b>	<b>i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000)</b>	<b>i7 bases for entry on sample sheet (all instruments)</b>
UDI 081	ATGGAATGGA	TCCATTCCAT	CCTCCATTAA
UDI 082	CATTCTCAC	GTGAGGAATG	AGTCGCGGTT
UDI 083	GCATAGGAAG	CTTCCTATGC	CTCATCCAGG
UDI 084	TGTTCTGTT	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	TTGTACTCCA	TGGAGTACAA	CTGATGATCT
UDI 090	GTGCACATAA	TTATGTGCAC	ACTAGGTGTT
UDI 091	AGGACAAGTA	TACTTGCCT	CTGTTAGCGG
UDI 092	CCGATTCGAG	CTCGAATCGG	ATCGCACCAA
UDI 093	GTAGGAACTT	AAGTTCCTAC	CTTACTGGT
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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<b>Indices for entry on sample sheet</b>			
<b>Unique dual-index number</b>	<b>i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000)</b>	<b>i5 bases for entry on sample sheet (iSeq 100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000)</b>	<b>i7 bases for entry on sample sheet (all instruments)</b>
UDI 101	AGAACGTCGC	GCGACGTTCT	TAGCATCGAT
UDI 102	CACGGACTAG	CTAGTCCGTG	CCATGAGTCG
UDI 103	GTTGAACACT	AGTGTCAAC	ACTAACATGC
UDI 104	TCGCGTGGTA	TACCACGCGA	ACACTCTCTA
UDI 105	AGCCACTATG	CATAGTGGCT	GCTCTGCTT
UDI 106	CCACCTACCA	TGGTAGGTGG	AATCTTGAGG
UDI 107	GTCCGGTGT	ACACCGAAC	CCTAACGGTC
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	GTATGTGCT	TGTTGGCTGT
UDI 118	CAAGTAGTCT	AGACTACTTG	CGTATCTGCG
UDI 119	GTCAAGAAGA	TCTTCTGAC	CCATAGTATC
UDI 120	TCCTGTTATG	CATAACAGGA	TGGACAGTAA

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<b>Indices for entry on sample sheet</b>			
<b>Unique dual-index number</b>	<b>i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000)</b>	<b>i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000)</b>	<b>i7 bases for entry on sample sheet (all instruments)</b>
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	CATTCTGAA
UDI 126	AGTTGGTCTA	TAGACCAACT	AAGATGCATG
UDI 127	ATGCAGCTGG	CCAGTGCAT	CCTGGAGCT
UDI 128	CGTTGCCTTC	GAAGGCAACG	ACCGGAACAG
UDI 129	GCGTGGAGAA	TTCTCCACGC	GAATGGAAGC
UDI 130	TACGCCTCCT	AGGAGGCGTA	GTTCTCCATA
UDI 131	AATTCGGTAG	CTACCGAATT	GTCACTATGT
UDI 132	ATTGTGGAAC	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	GTCCTCTCGG
UDI 139	GACAAC TGTC	GACAGTTGTC	GAAGCCACTC
UDI 140	TCCATTGCGG	CCGCAATGGA	GTAGGACACA

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<b>Indices for entry on sample sheet</b>			
<b>Unique dual-index number</b>	<b>i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000)</b>	<b>i5 bases for entry on sample sheet (iSeq 100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000)</b>	<b>i7 bases for entry on sample sheet (all instruments)</b>
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	TGTTTCGTGGA	AGCCACACTA
UDI 156	TGACGGAGGA	TCCTCCGTC A	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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<b>Indices for entry on sample sheet</b>			
<b>Unique dual-index number</b>	<b>i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000)</b>	<b>i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000)</b>	<b>i7 bases for entry on sample sheet (all instruments)</b>
UDI 161	CTCCTCGAAG	CTTCGAGGAG	AAGACCTCTT
UDI 162	ATACCGCAGA	TCTGCGGTAT	GACCTCTTCT
UDI 163	CCTATCTGAT	ATCAGATAGG	TACTTCCTTG
UDI 164	GATCGGTTAC	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	CGTGAATAT
UDI 173	CCGAGACAT	ATGTCTCCGG	CAACCATCGG
UDI 174	GAGATACTG	CAGTTATCTC	CGGTCCATTC
UDI 175	TTGTAAGCGC	GCGCTTACAA	AGAAGAGCCA
UDI 176	CAAGAGGAGG	CCTCTCTTG	CTATGCAATG
UDI 177	AACCTTAGGA	TCCTAAGGTT	CACTGAACCG
UDI 178	CTGGCAACTC	GAGTTGCCAG	TACTGTGTGA
UDI 179	GAACCTGTG	CAACAAGTTC	GCATTCTGTT
UDI 180	TGTGCAAGAT	ATCTTGACACA	CTCCGCTAAG

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<b>Indices for entry on sample sheet</b>			
<b>Unique dual-index number</b>	<b>i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000)</b>	<b>i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000)</b>	<b>i7 bases for entry on sample sheet (all instruments)</b>
UDI 181	AATCGAGAGA	TCTCTCGATT	TCGCTTGAGA
UDI 182	AGCGTGTCAG	CTGACACGCT	AACTAGCCTT
UDI 183	CTTGGTGATT	AATCACCAAG	TTCGCTCAGG
UDI 184	GAAGCAGCAA	TTGCTGCTTC	CTCTACAACA
UDI 185	TTCCGTCGAC	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	TATCCTCCAG
UDI 199	GCGAGATATA	TATATCTCGC	TAAGTCGTTT
UDI 200	TGACTGCTAT	ATAGCAGTCA	TCCGGATTGA

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<b>Indices for entry on sample sheet</b>			
<b>Unique dual-index number</b>	<b>i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000)</b>	<b>i5 bases for entry on sample sheet (iSeq 100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000)</b>	<b>i7 bases for entry on sample sheet (all instruments)</b>
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	CCTTACGCCA	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	GTTCCAATAG	CTATTGGAAC	ATCACGCTTC
UDI 216	TGTCAGGCTC	GAGCCTGACA	TAGCTATGCA
UDI 217	CAACAGTGTT	AACACTGTTG	TGTTCTCAT
UDI 218	AAGAGAGGAA	TTCTCTCTT	CATACCTTCT
UDI 219	CGGTTGTAGC	GCTACAACCG	GCCTTCAATG
UDI 220	GCCTGAAGTG	CACCTCAGGC	CTTGACCAGC

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<b>Indices for entry on sample sheet</b>			
<b>Unique dual-index number</b>	<b>i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000)</b>	<b>i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000)</b>	<b>i7 bases for entry on sample sheet (all instruments)</b>
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	TGGATTGTTT	CCGAATATTC

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<b>Indices for entry on sample sheet</b>			
<b>Unique dual-index number</b>	<b>i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000)</b>	<b>i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000)</b>	<b>i7 bases for entry on sample sheet (all instruments)</b>
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	AGTGTGATA
UDI 247	CAGTCCAATT	AATTGGACTG	ATCCTTCGGT
UDI 248	AGAGGCTTGG	CCAAGCCTCT	GACAACGATT
UDI 249	CAGGCTCTCA	TGAGAGCCTG	GAACCGGTAG
UDI 250	GTTGCCTCTC	GAGAGCGAAC	AGCAATGAGC
UDI 251	TCGGACTAAT	ATTAGCCGA	CAAGACTCCA
UDI 252	CGAGATCTTC	GAAGATCTCG	ACCGTGTAGG
UDI 253	ATAACCGGAC	GTCCGGTTAT	AGGCACAGGT
UDI 254	CGTGTAGTTA	TAACTACACG	CGACAGATCG
UDI 255	GAACATAGGT	ACCTATGTC	ACGCGACAAC
UDI 256	TCTAACATCG	CGATGTTAGA	ACTTGCCTTA
UDI 257	AACGGTGGCA	TGCCACCGTT	CACCACTCAT
UDI 258	AGGACGGTGT	ACACCGTCTT	CTTCGTA ACT
UDI 259	CTGTGACCTG	CAGGTCACAG	CAGTATTCGG
UDI 260	GCTGTAACAA	TTGTTACAGC	CAGTCTGGAC

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<b>Indices for entry on sample sheet</b>			
<b>Unique dual-index number</b>	<b>i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000)</b>	<b>i5 bases for entry on sample sheet (iSeq 100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000)</b>	<b>i7 bases for entry on sample sheet (all instruments)</b>
UDI 261	TACGGACGTC	GACGTCCGTA	TACCGTTCTA
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	TCTGGAAGT
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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<b>Indices for entry on sample sheet</b>			
<b>Unique dual-index number</b>	<b>i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000)</b>	<b>i5 bases for entry on sample sheet (iSeq 100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000)</b>	<b>i7 bases for entry on sample sheet (all instruments)</b>
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	GTGTCTTATT	TCGACCGCAT
UDI 289	ATGTGCCTCA	TGAGGCACAT	TAACCTAGGT
UDI 290	CGACTCCGTT	AACGGAGTCG	AACTCATGCG
UDI 291	GCTGTTGTGG	CCACAACAGC	CCGGATGAAC
UDI 292	TACCAATCAC	GTGATTGGTA	CGTTGCCGTA
UDI 293	ATGTCTTACG	CGTAAGACAT	GCTCTACGGT
UDI 294	CGCAACAATA	TATTGTTGCG	TGCATTGGCG
UDI 295	GAACGAAGAC	GTCTTCGTTT	CGATTGTGAC
UDI 296	TCGAGGACGT	ACGTCTCTGA	GACTGCACTA
UDI 297	ATTATGAGCG	CGCTCATAAT	GTTAACTGCT
UDI 298	CGCGTTATAA	TTATAACGCG	TCGGACCTTG
UDI 299	GCGTGTCATGT	ACATGCACGC	TGCAGCAAGC
UDI 300	TAAGCGGCTC	GAGCCGCTTA	CACATGCGAA

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<b>Indices for entry on sample sheet</b>			
<b>Unique dual-index number</b>	<b>i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000)</b>	<b>i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000)</b>	<b>i7 bases for entry on sample sheet (all instruments)</b>
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	TGGCGACCTA	TAGGTCGCCA	CTTGCTCACA
UDI 309	AGAAGCGACA	TGTCGCTTCT	CAACACCTGT
UDI 310	CAGGATAATC	GATTATCCTG	CAATTGCTCG
UDI 311	GCTCCTACAG	CTGTAGGAGC	CATAGACAAC
UDI 312	TCAACAGGT	ACCTGTTGAA	TTGGTGCTA
UDI 313	CCTCGTCCAT	ATGGACGAGG	TATGTCCTGT
UDI 314	AGCGTTGGTT	AACCAACGCT	GCCAAATCGT
UDI 315	CATTGAACA	TGTTCAATG	TAGGCGATCG
UDI 316	GCTTACCGAC	GTCGGTAAGC	ATGAGTG TAC
UDI 317	TTAGCTTAGG	CCTAAGCTAA	CCGAAGGATA
UDI 318	CCGACACACA	TGTGTGTCGG	AGTCCACTGT
UDI 319	ATTCGCTGAT	ATCAGCGAAT	GCGGCTAATT
UDI 320	CCAAGAGGCA	TGCCTCTGG	TCTAACTCAG

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<b>Indices for entry on sample sheet</b>			
<b>Unique dual-index number</b>	<b>i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000)</b>	<b>i5 bases for entry on sample sheet (iSeq 100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000)</b>	<b>i7 bases for entry on sample sheet (all instruments)</b>
UDI 321	GACGCAGTTC	GAACTGCGTC	CAAGCTGAGC
UDI 322	TGGAACTCGG	CCGAGTTCCA	CCAGAGCACA
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	CGGTGGTTCT
UDI 330	CGGTCGATTC	GAATCGACCG	CTGTGCTTGG
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

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<b>Indices for entry on sample sheet</b>			
<b>Unique dual-index number</b>	<b>i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000)</b>	<b>i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000)</b>	<b>i7 bases for entry on sample sheet (all instruments)</b>
UDI 341	GTAGACGCTC	GAGCGTCTAC	CCTATCCATC
UDI 342	TCGTCCGTCA	TGACGGACGA	ACCGCGAGTA
UDI 343	CCGTGATAGG	CCTATCACGG	AAGTTCTGGT
UDI 344	AGGATGACCT	AGGTCATCCT	ACAGGTATCG
UDI 345	CCTCGAGTAC	GTA CTGAGG	ATGACGGATT
UDI 346	GTC ACTGAGG	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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<b>Indices for entry on sample sheet</b>			
<b>Unique dual-index number</b>	<b>i5 bases for entry on sample sheet (MiSeq, HiSeq 2000/2500, NovaSeq 6000)</b>	<b>i5 bases for entry on sample sheet (iSeq100, MiniSeq, NextSeq, HiSeqX, HiSeq 3000/4000)</b>	<b>i7 bases for entry on sample sheet (all instruments)</b>
UDI 361	ATGTGGAGGT	ACCTCCACAT	CGCGTTATCG
UDI 362	CCATGGCTCA	TGAGCCATGG	GTAGCCTCCT
UDI 363	CCAATCACGC	GCGTGATTGG	ACTAGACACT
UDI 364	TTAGATCCAG	CTGGATCTAA	CGATTCGTTG
UDI 365	AGGAATATCG	CGATATTCCT	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	CTGGATAAGT
UDI 376	CTAGACCGCA	TGCGGTCTAG	ATAGCTTACG
UDI 377	TATGGCTTGT	ACAAGCCATA	GTCCATGAGT
UDI 378	CGTTGTTCCT	AGGAACAACG	ACTCCAGTCG
UDI 379	CCGACATTAG	CTAATGTCGG	TCTCAGCACG
UDI 380	TGTGAAGGCA	TGCCTCACA	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 6 and Figure 7. The index motives used in the QIAseq Combinatorial Dual-Index Kits are listed in Table 10. To make sequencing preparation more convenient, you can download Illumina-compatible sample sheets for different sequencing instruments at [www.qiagen.com](http://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 6. QIAseq CDI Y-Adapter Plate (96) layout (CDI 1–96).

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

**Figure 7. QIAseq CDI Y-Adapter Plate (24) layout (CDI 1–24).**

**Table 10. 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	GTA CTGAC	GTCAGTAC	D708	TAATGCGC
			D709	CGGCTATG
			D710	TCCGCGAA
			D711	TCTCGCGC
			D712	AGCGATAG

# Appendix D: Adapter Plate Preparation for Ultralow Input Ligation

This appendix describes how to prepare the following QIAGEN adapter plates for Ultralow Input Ligation:

- QIAseq CDI Y-Adapter Plate (24), mat. no. 1100697
- QIAseq CDI Y-Adapter Plate (96), mat. no. 1096117
- QIAseq UDI Y-Adapter Plate (24), mat. no. 1118061
- QIAseq UDI Y-Adapter Plate A (96), mat. no. 1118062
- QIAseq UDI Y-Adapter Plate B (96), mat. no. 1118063
- QIAseq UDI Y-Adapter Plate C (96), mat. no. 1118064
- QIAseq UDI Y-Adapter Plate D (96), mat. no. 1118065

Prepare Adapters for Ultralow Input Ligation by thawing and diluting in nuclease-free water prior to Adapter Ligation setup. QIAGEN recommends the following dilution factors to achieve an optimum molar ratio between fragmented sample DNA and adapter oligos:

**Table 11. Adapter dilution factors**

Sample DNA Amount	QIAseq Adapter Plate Dilution
10–99 pg	1:1000 dilution
100–999 pg	1:100 dilution
1–9 ng	1:10 dilution
10–100 ng	No dilution

1. Thaw the Adapter Plate on ice. Vortex and spin down thawed adapters before use.
2. Remove the clear protective adapter plate lid and carefully pierce only the foil seal for each adapter well to be used. Use a fresh pipette tip to pierce each well to avoid cross-contamination.
3. Dilute the adapters according to Table 10 to ensure a correct molar ratio with fragmented sample DNA. If necessary, use fresh PCR tubes or 96-well PCR plate for dilutions.

**To achieve a 1:10 Adapter Plate dilution:**

- 3a. Add 90  $\mu$ l nuclease-free water to each adapter well to be used. With a multi-channel pipettor set to 30  $\mu$ l, pipet up and down 5–6 times to mix. 1:10 diluted adapter is ready for use.

**To achieve a 1:100 Adapter Plate dilution:**

- 3a. Add 90  $\mu$ l nuclease-free water to each adapter well to be used. With a multi-channel pipettor set to 30  $\mu$ l, pipet up and down 5–6 times to mix.
- 3b. Using a multi-channel pipettor, transfer 10  $\mu$ l 1:10 diluted adapter to a fresh 96-well plate. Carefully track the location of each adapter barcode in the new plate.
- 3c. Add 90  $\mu$ l nuclease-free water to each well in the 96-well PCR plate. With a multi-channel pipettor set to 30  $\mu$ l, pipet up and down 5–6 times to mix. 1:100 diluted adapter is ready for use.

**To achieve a 1:1000 Adapter Plate dilution:**

- 3a. Add 90  $\mu$ l nuclease-free water to each adapter well to be used. With a multi-channel pipettor set to 30  $\mu$ l, pipet up and down 5–6 times to mix.
- 3b. In a fresh 96-well PCR plate, add 198  $\mu$ l nuclease-free water.
- 3c. Using a multi-channel pipettor, remove 2  $\mu$ l 1:10 diluted adapter and add to the PCR tubes or wells in the 96-well PCR plate that already contain 198  $\mu$ l nuclease-free water. Carefully track the location of each adapter barcode in the new plate.
- 3d. With a multi-channel pipettor set to 30  $\mu$ l, pipet up and down 5–6 times to mix. 1:1000 diluted adapter is now ready for use.

- 
4. After adapter ligation, replace the adapter plate lid and freeze unused adapters at  $-15$  to  $-35^{\circ}\text{C}$ . If desired, residual diluted adapter can be removed and discarded before plate storage.
  5. Do not attempt to reuse diluted adapter due to the risk of barcode cross-contamination and lower than expected adapter concentration after storage of very dilute oligos.

# Appendix E: GeneRead Adapter Preparation for Ultralow Input Ligation

This appendix describes how to prepare the following QIAGEN adapter products for Ultralow Input Ligation:

- GeneRead Adapter I Set A 12-plex (144), cat. no 180985
- GeneRead Adapter I Set B 12-plex (144), cat. no 180986

Adapter preparation includes thawing and diluting in nuclease-free water prior to adapter ligation setup. QIAGEN recommends the following dilution factors to achieve an optimum molar ratio between fragmented sample DNA and adapter oligos:

**Table 12. Adapter dilution factors**

Sample DNA amount	GeneRead Adapter I dilution
10–99 pg	1:1000
100–999 pg	1:100
1–9 ng	1:10
10–100 ng	No dilution

Adapter Preparation using the GeneRead Adapter I Set A (cat. no. 180985) or GeneRead Adapter I Set B (cat. no. 180496)

**Note:** When using GeneRead Adapters, open one tube at a time and change gloves between pipetting the different barcode adapters to avoid cross-contamination.

1. Thaw the GeneRead Adapter tubes ice. Vortex and spin down thawed adapters before use.



2. Transfer 2  $\mu$ l of each adapter to be used to a new PCR tube or wells in a 96-well PCR plate. Carefully track the location of each adapter barcode in the new plate.
3. Re-cap each GeneRead Adapter tube immediately after use. Freeze unused adapters at  $-15$  to  $-35^{\circ}\text{C}$ .
4. Dilute the adapters in the PCR tubes or plate according to Table 11 to ensure a correct molar ratio with fragmented sample DNA.

**To achieve a 1:10 GeneRead Adapter dilution:**

- 4a. Add 18  $\mu$ l nuclease-free water to each adapter tube or well to be used. Pipet up and down 5–6 times to mix. 1:10 diluted adapter is ready for use.

**To achieve a 1:100 GeneRead Adapter dilution:**

- 4a. Add 18  $\mu$ l nuclease-free water to each adapter tube or well to be used. Pipet up and down 5–6 times to mix.
- 4b. Using a multi-channel pipettor, remove and discard 10  $\mu$ l 1:10 diluted adapter.
- 4c. Add 90  $\mu$ l nuclease-free water to each adapter tube or well to be used. Pipet up and down 5–6 times to mix. 1:100 diluted adapter is ready for use.

**To achieve a 1:1000 GeneRead Adapter dilution:**

- 4a. Add 18  $\mu$ l nuclease-free water to each adapter well to be used. Pipet up and down 5–6 times to mix.
  - 4b. Using a multi-channel pipettor, transfer 2  $\mu$ l 1:10 diluted adapter to a fresh tube or well. Carefully track the location of each adapter barcode in the new plate.
  - 4c. Add 198  $\mu$ l nuclease-free water to the 2  $\mu$ l 1:10 adapter. Pipet up and down 5–6 times to mix. 1:1000 diluted adapter is ready for use.
5. Discard residual diluted adapter.
  6. Do not attempt to reuse diluted adapter due to the risk of barcode cross-contamination and lower than expected adapter concentration after storage of very dilute oligos.

# Ordering Information

<b>Product</b>	<b>Contents</b>	<b>Cat. no.</b>
QIAseq Ultralow Input Library Kit (12)	For 12 reactions: Buffers and reagents for Ultralow Input End Polishing, Ultralow Input Ligation and HiFi library amplification. 12-plex adapters sold separately. For use with Illumina instruments.	180492
QIAseq Ultralow Input Lib UDI-A Kit (96)	For 96 reactions: Buffers and reagents for Ultralow Input End Polishing, Ultralow Input Ligation and HiFi library amplification. Includes 96-plex Adapter Plate with individually pierceable foil sealed wells. For use with Illumina instruments.	180497
QIAseq Ultralow Input Lib UDI-B Kit (96)	For 96 reactions: Buffers and reagents for Ultralow Input End Polishing, Ultralow Input Ligation and HiFi library amplification. Includes 96-plex Adapter Plate with individually pierceable foil sealed wells. For use with Illumina instruments.	180498
QIAseq Ultralow Input Lib UDI-C Kit (96)	For 96 reactions: Buffers and reagents for Ultralow Input End Polishing, Ultralow Input Ligation and HiFi library amplification. Includes 96-plex Adapter Plate with individually pierceable foil sealed wells. For use with Illumina instruments.	180499

<b>Product</b>	<b>Contents</b>	<b>Cat. no.</b>
QIAseq Ultralow Input Lib UDI-D Kit (96)	For 96 reactions: Buffers and reagents for Ultralow Input End Polishing, Ultralow Input Ligation and HiFi library amplification. Includes 96-plex Adapter Plate with individually pierceable foil sealed wells. For use with Illumina instruments.	180500
QIAseq Ultralow Input Lib CDI Kit (96)	For 96 reactions: Buffers and reagents for Ultralow Input End Polishing, Ultralow Input Ligation and HiFi library amplification. Includes 96-plex Adapter Plate with individually pierceable foil sealed wells. For use with Illumina instruments.	180501
<b>Related products</b>		
GeneRead Adapter I Set A 12-plex (144)	For 144 reactions: 12 barcoded adapters for ligation to DNA library, for use with Illumina instruments	180985
GeneRead Adapter I Set B 12-plex (144)	For 144 reactions: 12 barcoded adapters for ligation to DNA library, for use with Illumina instruments	180986
GeneRead Size Selection Kit (50)	For 50 reactions: Spin columns and buffers	180514
QIAseq Library Quant Array 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	333304

<b>Product</b>	<b>Contents</b>	<b>Cat. no.</b>
<b>QIAseq Y-Adapter Kits for Illumina</b>		
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
<b>For genomic DNA purification</b>		
QIAamp DNA Mini Kit	For 50 DNA preps: 50 QIAamp Mini Spin Columns, QIAGEN Proteinase K, Reagents, Buffers, Collection Tubes (2 ml)	51304
GeneRead DNA FFPE Kit	For 50 preps: QIAamp MinElute Columns, Collection Tubes, Deparaffinization Solution, Uracil-N-Glycosylase, RNase-Free Water, RNase A, and Buffers	180134

Product	Contents	Cat. no.
QIAamp DNA FFPE Tissue Kit	For 50 DNA preps: 50 QIAamp MinElute Columns, Proteinase K, Buffers, Collection Tubes (2 ml)	56404
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

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# Document Revision History

Date	Changes
08/2021	Removed the QIAseq Ultralow Input Library Kit (96) (cat. no. 180495) product for discontinuation. 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. Corrected typo error in Table 2, changed 100 pg to 10 pg in row 1 (10 pg is indicated in the table title). Corrected typo in Figure 7: changed entries in row H (from 501/701, 501/702, 501/703, to 508/701, 508/702, 508/703).

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