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# Chapter 1 Overview

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## Introduction

This protocol explains how to prepare up to 96 libraries starting from genomic DNA (gDNA) using the Illumina<sup>®</sup> TruSeq<sup>®</sup> DNA Nano library prep workflow. The goal is to add adapter sequences to DNA fragment ends to create indexed libraries for single-read or paired-end sequencing.

The TruSeq DNA Nano Library Prep workflow protocol includes the following features.

- ▶ Streamlined workflow:
  - ▶ Size-selection beads and master-mixed reagents reduce reagent containers and pipetting.
  - ▶ Universal adapter to prepare DNA libraries for single-read, paired-end, and indexed sequencing.
  - ▶ One workflow with options for processing low sample (LS) and high sample (HS) numbers.
- ▶ Flexible throughput:
  - ▶ 24- and 96-sample workflow configurations accommodate a range of experiments.
  - ▶ Support for non-indexed sequencing and low-plexity pooling.
  - ▶ Optimized shearing for whole-genome resequencing with insert sizes of 350 bp or 550 bp.
- ▶ Inclusive components:
  - ▶ Library Prep components include library prep reagents excluding index adapters.
  - ▶ Index adapter components must be purchased separately. See [Supporting Information on page 22](#) for more details.

## DNA Input Recommendations

Quantify the input gDNA and assess the quality before starting library preparation. For best results, use the following input amounts.

Insert Size	Input gDNA
350 bp	100 ng
550 bp	200 ng

Lower input amounts result in low yield and increased duplicates.

## Quantify Input DNA

Quantify input DNA per the following recommendations:

- ▶ Successful library prep depends on accurate quantification of input DNA.
- ▶ Use fluorometric-based methods for quantification, such as Qubit or PicoGreen to provide accurate quantification for dsDNA. UV spectrophotometric based methods, such as the Nanodrop, measures any nucleotides present in the sample including RNA, dsDNA, ssDNA, and free nucleotides, which can give an inaccurate measurement of gDNA.
- ▶ Quantification methods depend on accurate pipetting methods. Do not use pipettes at the extremes of volume specifications. Make sure that pipettes are calibrated.

## Assess DNA Quality

Absorbance measurements at 260 nm are commonly used to assess DNA quality:

- ▶ The ratio of absorbance at 260 nm to absorbance at 280 nm is used as an indication of sample purity. Values from 1.8 through 2.0 indicate relatively pure DNA.
- ▶ The presence of RNA or small nucleic acid fragments, such as nucleotides, can compromise both absorbance measurements.
- ▶ Make sure that samples are free of contaminants.

## Positive Control

Use Coriell Human-1 DNA (NA18507) or Promega Human Genomic DNA (G3041) as a positive control sample for this protocol.

## Additional Resources

The following documentation is available for download from the Illumina website.

Resource	Description
<a href="#">Custom Protocol Selector</a>	A wizard for generating customized end-to-end documentation that is tailored to the library prep method, run parameters, and analysis method used for the sequencing run.
<a href="#">TruSeq DNA Nano Checklist (document # 1000000040813)</a>	Provides a checklist of the protocol steps, and is intended for experienced users.
<a href="#">Index Adapter Pooling Guide (document # 1000000041074)</a>	Provides pooling guidelines for preparing libraries for sequencing systems that require balanced index combinations. Review this guide before beginning library preparation.
<a href="#">Illumina Experiment Manager Guide (part # 15031335) and IEM TruSeq DNA, RNA, or ChIP Quick Reference Card (document # 15037152)</a>	Provides information about creating and editing sample sheets.
<a href="#">BaseSpace Sequence Hub help</a>	Provides information about BaseSpace <sup>®</sup> Sequence Hub, a data analysis tool.
<a href="#">Local Run Manager Software Guide (document #100000002702)</a>	Provides an overview of the Local Run Manager (LRM) software, instructions for using software features, and instructions for installing analysis modules on the instrument computer.

Visit the [TruSeq DNA Nano workflow support page](#) on the Illumina website for access to requirements and compatibility, additional documentation, software downloads, online training, frequently asked questions, and best practices.

# Chapter 2 Protocol

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## Introduction

This chapter describes the TruSeq DNA Nano Library Prep workflow protocol.

- ▶ Follow the steps in the order shown, using the specified volumes and incubation parameters.
- ▶ Before proceeding, confirm the delivered contents and make sure that you have the required equipment and consumables.
- ▶ Review Best Practices from the TruSeq DNA Nano Library Prep workflow support page on the Illumina website.

This protocol provides one workflow with variations for differences in sample numbers. [HS] and [LS] identify the appropriate option for your number of samples. Expect equivalent results from either option, but the HS option can yield more consistent results between samples.

**Table 1 Workflow Variations**

Workflow Variable	HS	LS
24-Sample Workflow	Process > 24 samples with index adapter tubes*	Process ≤ 24 samples with index adapter tubes*
96-Sample Workflow	Process > 24 samples with index adapter plate	Process ≤ 24 samples with index adapter plate
Plate Type	96-well Hard-Shell PCR plate 96-well midi plate	96-well 0.3 ml PCR plate 96-well midi plate
Incubation Equipment	Microheating systems	96-well thermal cycler
Mixing Method	Microplate shaker	Pipetting

\* Combine the Set A and Set B indexes to pool up to 24 libraries.

## Tips and Techniques

Unless a safe stopping point is specified in the protocol, proceed immediately to the next step.

### Avoiding Cross-Contamination

- ▶ When adding or transferring samples, change tips between *each sample*.
- ▶ Remove unused index adapter tubes from the working area.

## Sealing the Plate

- ▶ Always seal the 96-well plate before the following steps in the protocol:
  - ▶ Shaking steps
  - ▶ Vortexing steps
  - ▶ Centrifuge steps
  - ▶ Thermal cycling steps
- ▶ Apply the adhesive seal to cover the plate, and seal with a rubber roller.
- ▶ Microseal 'B' adhesive seals are effective at -40°C to 110°C. Use Microseal 'B' for shaking, centrifuging, and long-term storage.
- ▶ Microseal 'A' adhesive film is used for thermal cycling steps to prevent evaporation.

## Plate Transfers

- ▶ When transferring volumes between plates, transfer the specified volume from each well of a plate to the corresponding well of the other plate.

## Centrifugation

- ▶ Centrifuge at any step in the procedure to consolidate liquid or beads in the bottom of the well, and to prevent sample loss.

## Handling Beads

- ▶ Do not freeze beads.
- ▶ Pipette bead suspensions slowly.
- ▶ Before use, allow the beads to come to room temperature.
- ▶ Immediately before use, vortex the beads until they are well dispersed. The color of the liquid must appear homogeneous. Vortex throughout protocol as necessary to keep homogenous.
- ▶ If beads are aspirated into pipette tips, dispense back to the plate on the magnetic stand, and wait until the liquid is clear (~2 minutes).
- ▶ When washing beads:
  - ▶ Use the specified magnetic stand for the plate.
  - ▶ Dispense liquid so that beads on the side of the wells are wetted.
  - ▶ Keep the plate on the magnetic stand until the instructions specify to remove it.
  - ▶ Do not agitate the plate while it is on the magnetic stand. Do not disturb the bead pellet.

## Library Prep Workflow

The following diagram illustrates the workflow using a TruSeq DNA Nano Library Prep workflow and Index Adapter components. Safe stopping points are marked between steps.

**Figure 1** TruSeq DNA Nano Workflow



## Prepare for Pooling

When pooling samples for sequencing, use IEM, LRM, or BaseSpace Prep Tab to record information about your samples before beginning library preparation.

- ▶ Use IEM to create and edit sample sheets for Illumina sequencing systems and analysis software.
- ▶ Use LRM and BaseSpace Prep Tab to organize samples, libraries, pools, and a run for Illumina sequencing systems and analysis software.

Review the planning steps in the *Index Adapter Pooling Guide (document # 1000000041074)* when preparing libraries that require balanced index combinations.

## Fragment DNA

This step fragments to an insert size of 350 bp or 550 bp. Covaris shearing generates double-stranded DNA (dsDNA) fragments with 3' or 5' overhangs.

## Consumables

- ▶ gDNA samples
  - ▶ [350 bp insert size] 100 ng per sample
  - ▶ [550 bp insert size] 200 ng per sample
- ▶ RSB (Resuspension Buffer)
- ▶ SPB (Sample Purification Beads)
- ▶ Barcode labels
  - ▶ CFP (Covaris Fragmentation Plate)
  - ▶ CSP (Clean Up Sheared DNA Plate)
  - ▶ DNA (DNA Plate)
  - ▶ IMP (Insert Modification Plate)
- ▶ Freshly prepared 80% ethanol (EtOH)
- ▶ Plates
  - ▶ [HS] 96-well midi plates (3)
  - ▶ [HS] 96-well Hard-Shell 0.3 ml PCR plate (1)
  - ▶ [LS] 96-well 0.3 ml PCR plates, semiskirted or skirtless (4)
- ▶ Covaris tubes (1 per sample)
- ▶ Microseal 'B' adhesive seal

## About Reagents

- ▶ Vortex SPB before each use.
- ▶ Vortex SPB frequently to make sure that beads are evenly distributed.
- ▶ Aspirate and dispense SPB slowly due to the viscosity of the solution.

## Preparation

- 1 Prepare the following consumables.

Item	Storage	Instructions
RSB	-25°C to -15°C	Thaw at room temperature. After the initial thaw, store at 2°C to 8°C.
SPB	2°C to 8°C	Let stand for 30 minutes to bring to room temperature.

- 2 Turn on and set up the Covaris instrument per manufacturer guidelines.
- 3 [HS] Calibrate the microplate shaker with a stroboscope and set to 1800 rpm.
- 4 Apply barcode labels to plates.

Barcode Label	Plate for HS	Plate for LS
DNA	Midi	PCR
CFP	Hard-Shell PCR	PCR
CSP	Midi	PCR
IMP	Midi	PCR

## Procedure

### Normalize gDNA

- 1 Quantify gDNA using a fluorometric-based method.
- 2 Normalize gDNA samples with RSB to a final volume of 52.5  $\mu$ l in the DNA plate.
  - ▶ 100 ng for a 350 bp insert size.
  - ▶ 200 ng for a 550 bp insert size.
- 3 [HS] Mix and centrifuge as follows.
  - a Shake at 1800 rpm for 2 minutes.
  - b Centrifuge at 280  $\times$  g for 1 minute.
- 4 [LS] Pipette to mix, and then centrifuge briefly.

### Fragment DNA

- 1 Transfer 52.5  $\mu$ l DNA samples to separate Covaris tubes. Use the wells of the CFP plate to hold the tubes upright.
- 2 Centrifuge at 280  $\times$  g for 5 seconds.
- 3 Fragment using the appropriate Covaris settings:

**Table 2 350 bp Insert**

Setting	M220	S220	S2	E210
Duty Cycle (%)	20	5		10
Intensity	—	—		5.0
Peak/Displayed Power (W)	50	175	23	14
Cycles/Burst			200	
Duration (seconds)	65	50		45
Mode	—		Frequency sweeping	
Temperature ( $^{\circ}$ C)	20		5.5–6	

**Table 3 550 bp Insert**

Setting	M220	S220	S2	E210
Duty Cycle (%)	20	5		10
Intensity	—	—		2.0
Peak/Displayed Power (W)	50	175	9	7
Cycles/Burst			200	
Duration (seconds)	45	25		45
Mode	—		Frequency sweeping	
Temperature ( $^{\circ}$ C)	20		5.5–6	

- 4 Centrifuge at 280  $\times$  g for 5 seconds.
- 5 Transfer 50  $\mu$ l sample from each Covaris tube to the corresponding well of the CSP plate.

## Clean Up Fragmented DNA

- 1 Vortex SPB until well-dispersed.
- 2 Add 80  $\mu$ l SPB to each well.
- 3 Mix thoroughly as follows.
  - ▶ [HS] Shake at 1800 rpm for 2 minutes.
  - ▶ [LS] Pipette up and down.
- 4 Incubate at room temperature for 5 minutes.
- 5 Centrifuge at 280  $\times$  g for 1 minute.
- 6 Place on a magnetic stand and wait until the liquid is clear (~8 minutes).
- 7 Remove and discard all supernatant from each well.
- 8 Wash two times as follows.
  - a Add 200  $\mu$ l fresh 80% EtOH to each well.
  - b Incubate on the magnetic stand for 30 seconds.
  - c Remove and discard all supernatant from each well.
- 9 Use a 20  $\mu$ l pipette to remove residual EtOH from each well.
- 10 Air dry on the magnetic stand for 5 minutes.
- 11 Add 62.5  $\mu$ l RSB to each well, and then remove from the magnetic stand.
- 12 Mix thoroughly as follows.
  - ▶ [HS] Shake at 1800 rpm for 2 minutes.
  - ▶ [LS] Pipette up and down.
- 13 Incubate at room temperature for 2 minutes.
- 14 Centrifuge at 280  $\times$  g for 1 minute.
- 15 Place on a magnetic stand and wait until the liquid is clear (2–5 minutes).
- 16 Transfer 60  $\mu$ l supernatant to the corresponding well of the IMP plate.

## Repair Ends and Select Library Size

This step uses End Repair Mix 2 to convert the overhangs resulting from fragmentation into blunt ends. A 3' to 5' exonuclease activity removes the 3' overhangs. A 5' to 3' polymerase activity completes the 5' overhangs. After end repair, different ratios of Sample Purification Beads are used to select the appropriate library size.

### Consumables

- ▶ ERP 2 or ERP 3 (End Repair Mix)
- ▶ RSB (Resuspension Buffer)
- ▶ SPB (Sample Purification Beads)
- ▶ Barcode labels
  - ▶ ALP (Adapter Ligation Plate)
  - ▶ CEP (Clean Up End Repair Plate)
- ▶ Freshly prepared 80% ethanol (EtOH)

- ▶ PCR-grade water
- ▶ Tube
  - ▶ [≤ 6 samples] 1.7 ml microcentrifuge tube
  - ▶ [> 6 samples] 15 ml conical tube
- ▶ Plates
  - ▶ [HS] 96-well midi plates (2)
  - ▶ [LS] 96-well 0.3 ml PCR plates, semiskirted or skirtless (2)
- ▶ Microseal 'B' adhesive seals

## About Reagents

- ▶ Vortex SPB before each use.
- ▶ Vortex SPB frequently to make sure that beads are evenly distributed.
- ▶ Aspirate and dispense SPB slowly due to the viscosity of the solution.

## Preparation

- 1 Prepare the following consumables.

Item	Storage	Instructions
ERP 2 or ERP 3	-25°C to -15°C	Thaw at room temperature, and then set aside on ice. Return to storage after use.
RSB	2°C to 8°C	Let stand for 30 minutes to bring to room temperature.
SPB	2°C to 8°C	Let stand for 30 minutes to bring to room temperature.

- 2 [HS] Preheat the microheating system to 30°C.
- 3 [LS] Save the following ERP program on the thermal cycler:
  - ▶ Choose the preheat lid option and set to 100°C
  - ▶ 30°C for 30 minutes
  - ▶ Hold at 4°C
- 4 Label plates as follows.
  - ▶ Apply an ALP barcode label to a midi or PCR plate.
  - ▶ Apply a CEP barcode label to a midi or PCR plate.

## Procedure

### Convert Overhangs

- 1 Centrifuge ERP 2 at 600 × g for 5 seconds.
- 2 Add 40 µl ERP 2 or ERP 3 to each well.
- 3 [HS] Mix, centrifuge, and incubate as follows.
  - a Shake at 1800 rpm for 2 minutes.
  - b Centrifuge at 280 × g for 1 minute.
  - c Place on the 30°C microheating system, lid closed, for 30 minutes.
  - d Place on ice.

- [LS] Pipette to mix, centrifuge briefly, and then place on the thermal cycler and run the ERP program. Each well contains 100  $\mu$ l.

## Remove Large DNA Fragments

- Vortex SPB until well-dispersed.
- Using the following formulas, determine the appropriate volumes of SPB and PCR-grade water for diluting SPB.

The formulas include 15% excess for multiple samples.

**Table 4 Diluted SPB for a 350 bp Insert Size**

Reagent	Formula	Example Volume for 12 Samples	Your Calculation
SPB	# of samples $\times$ 109.25 $\mu$ l	1311 $\mu$ l	
PCR-grade water	# of samples $\times$ 74.75 $\mu$ l	897 $\mu$ l	

**Table 5 Diluted SPB for a 550 bp Insert Size**

Reagent	Formula	Example Volume for 12 Samples	Your Calculation
SPB	# of samples $\times$ 92 $\mu$ l	1104 $\mu$ l	
PCR-grade water	# of samples $\times$ 92 $\mu$ l	1104 $\mu$ l	

- Using your calculations from the previous step, dilute SPB with PCR-grade water.
  - ▶ For  $\leq$  6 samples, dilute in a new 1.7 ml microcentrifuge tube.
  - ▶ For  $>$  6 samples, dilute in a new 15 ml conical tube.
- Vortex diluted SPB until well-dispersed.
- Add 160  $\mu$ l diluted SPB to each well.
- Mix thoroughly as follows.
  - ▶ [HS] Shake at 1800 rpm for 2 minutes.
  - ▶ [LS] Pipette up and down.
- Incubate at room temperature for 5 minutes.
- Centrifuge at 280  $\times$  g for 1 minute.
- Place on a magnetic stand and wait until the liquid is clear (~5 minutes).
- Transfer 250  $\mu$ l supernatant to the corresponding well of the CEP plate.
- Discard remaining diluted SPB.

## Remove Small DNA Fragments

- Vortex undiluted SPB until well-dispersed.
- Add 30  $\mu$ l undiluted SPB to each well.
- Mix thoroughly as follows.
  - ▶ [HS] Shake at 1800 rpm for 2 minutes.
  - ▶ [LS] Pipette up and down.
- Incubate at room temperature for 5 minutes.

- 5 Centrifuge at 280 × g for 1 minute.
- 6 Place on a magnetic stand and wait until the liquid is clear (~5 minutes).
- 7 Remove and discard all supernatant from each well.
- 8 Wash two times as follows.
  - a Add 200 µl fresh 80% EtOH to each well.
  - b Incubate on the magnetic stand for 30 seconds.
  - c Remove and discard all supernatant from each well.
- 9 Use a 20 µl pipette to remove residual EtOH from each well.
- 10 Air dry on the magnetic stand for 5 minutes.
- 11 Add 20 µl RSB to each well, and then remove from the magnetic stand.
- 12 Mix thoroughly as follows.
  - ▶ [HS] Shake at 1800 rpm for 2 minutes.
  - ▶ [LS] Pipette up and down.
- 13 Incubate at room temperature for 2 minutes.
- 14 Centrifuge at 280 × g for 1 minute.
- 15 Place on a magnetic stand and wait until the liquid is clear (~5 minutes).
- 16 Transfer 17.5 µl supernatant to the corresponding well of the ALP plate.

### SAFE STOPPING POINT

If you are stopping, seal the plate and store at -25°C to -15°C for up to 7 days.

## Adenylate 3' Ends

One adenine (A) nucleotide is added to the 3' ends of the blunt fragments to prevent them from ligating to each other during adapter ligation. A corresponding thymine (T) nucleotide on the 3' end of the adapter provides a complementary overhang for ligating the adapter to the fragment. This strategy ensures a low rate of chimera (concatenated template) formation.

### Consumables

- ▶ ATL or ATL 2 (A-Tailing Mix)
- ▶ RSB (Resuspension Buffer)
- ▶ Microseal 'B' adhesive seals

### Preparation

- 1 Prepare the following consumables.

Item	Storage	Instructions
ATL or ATL 2	-25°C to -15°C	Thaw at room temperature. Return to storage after use.
RSB	2°C to 8°C	Let stand for 30 minutes to bring to room temperature.

- 2 [HS] Preheat two microheating systems, one to 37°C and another to 70°C.
- 3 [LS] Save the following ATAIL70 program on the thermal cycler:

- ▶ Choose the preheat lid option and set to 100°C
- ▶ 37°C for 30 minutes
- ▶ 70°C for 5 minutes
- ▶ 4°C for 5 minutes
- ▶ Hold at 4°C

## Procedure

- 1 Centrifuge ATL or ATL 2 at 600 × g for 5 seconds.
- 2 Add 12.5 µl ATL or ATL 2 to each well.
- 3 Mix thoroughly as follows.
  - ▶ [HS] Shake at 1800 rpm for 2 minutes.
  - ▶ [LS] Pipette up and down.
- 4 Centrifuge at 280 × g for 1 minute.
- 5 [HS] Incubate as follows.
  - a Place on the 37°C microheating system, lid closed, for 30 minutes.
  - b Move to the 70°C microheating system, lid closed, for 5 minutes.
  - c Place on ice for 5 minutes.
- 6 [LS] Incubate as follows.
  - a Place on the thermal cycler and run the ATAIL70 program. Each well contains 30 µl.
  - b Centrifuge at 280 × g for 1 minute.

## Ligate Adapters

This process ligates index adapters to the ends of the DNA fragments, which prepares them for hybridization onto a flow cell.

Index adapters must be ordered separately from the Library Prep components. For information on compatible index adapters, see [Supporting Information on page 22](#).

## Consumables

- ▶ DNA Adapters (tubes or index adapter plate)
- ▶ LIG 2 (Ligation Mix)
- ▶ RSB (Resuspension Buffer)
- ▶ SPB (Sample Purification Beads)
- ▶ STL (Stop Ligation Buffer)
- ▶ Barcode labels
  - ▶ CAP (Clean Up ALP Plate)
  - ▶ Index Adapter Components
  - ▶ PCR (Polymerase Chain Reaction Plate)
- ▶ Freshly prepared 80% ethanol (EtOH)
- ▶ Plates
  - ▶ [HS] 96-well midi plate (1)
  - ▶ [HS] 96-well Hard-Shell 0.3 ml PCR plate (1)

- ▶ [LS] 96-well 0.3 ml PCR plates, semiskirted or skirtless (2)
- ▶ [HS] Microseal 'B' adhesive seals

## About Reagents

- ▶ Do not remove LIG 2 from storage until instructed to do so in the procedure.
- ▶ Return LIG 2 to storage immediately after use.
- ▶ Vortex SPB before each use.
- ▶ Vortex SPB frequently to make sure that beads are evenly distributed.
- ▶ Aspirate and dispense SPB slowly due to the viscosity of the solution.

## Preparation

- 1 Prepare the following consumables.

Item	Storage	Instructions
DNA Adapters	-25°C to -15°C	Thaw at room temperature for 10 minutes. Return to storage after use.
STL	-25°C to -15°C	Thaw at room temperature. Return to storage after use.
RSB	2°C to 8°C	Let stand for 30 minutes to bring to room temperature.
SPB	2°C to 8°C	Let stand for 30 minutes to bring to room temperature.

- 2 [HS] Preheat a microheating system to 30°C.
- 3 [LS] Save the following LIG program on the thermal cycler:
  - ▶ Choose the preheat lid option and set to 100°C
  - ▶ 30°C for 10 minutes
  - ▶ Hold at 4°C
- 4 Label plates as follows.
  - ▶ Apply a CAP barcode label to a midi or PCR plate.
  - ▶ Apply a PCR barcode label to a Hard-Shell PCR or PCR plate.

## Procedure

### Add Index Adapters

- 1 [HS] Prepare the appropriate Index Adapter Plate as follows.
  - a Remove the tape seal.
  - b Centrifuge at 280 × g for 1 minute.
  - c Remove the plastic cover. If you are not processing the entire plate, save the cover.
  - d Apply the index adapter plate barcode label.
- 2 [LS] Centrifuge the adapter tubes at 600 × g for 5 seconds.
- 3 Remove LIG 2 from -25°C to -15°C storage.
- 4 In the order listed, add the following reagents to each well:
  - ▶ RSB (2.5 µl)
  - ▶ LIG 2 (2.5 µl)

- ▶ DNA adapters (2.5  $\mu$ l)
- 5 Mix thoroughly as follows.
  - ▶ [HS] Shake at 1800 rpm for 2 minutes.
  - ▶ [LS] Pipette up and down.
- 6 Centrifuge at 280  $\times$  g for 1 minute.
- 7 Incubate as follows.
  - ▶ [HS] Place on the 30°C microheating system, lid closed, for 10 minutes. Set aside on ice.
  - ▶ [LS] Place on the thermal cycler and run the LIG program.  
Each well contains 37.5  $\mu$ l.
- 8 Centrifuge the STL at 600  $\times$  g for 5 seconds.
- 9 Add 5  $\mu$ l STL to each well.
- 10 Mix thoroughly as follows.
  - ▶ [HS] Shake at 1800 rpm for 2 minutes.
  - ▶ [LS] Pipette up and down.
- 11 Centrifuge at 280  $\times$  g for 1 minute.

## Clean Up Ligated Fragments

Steps 1 through 14 are performed one time using the Round 1 volumes, then repeated using the Round 2 volumes.

- 1 Add the appropriate volume of SPB to each well.
  - ▶ **Round 1** —42.5  $\mu$ l
  - ▶ **Round 2** —50  $\mu$ l
- 2 Mix thoroughly as follows.
  - ▶ [HS] Shake at 1800 rpm for 2 minutes.
  - ▶ [LS] Pipette up and down.
- 3 Incubate at room temperature for 5 minutes.
- 4 Centrifuge at 280  $\times$  g for 1 minute.
- 5 Place on a magnetic stand and wait until the liquid is clear (2–5 minutes).
- 6 Remove and discard all supernatant from each well.
- 7 Wash two times as follows.
  - a Add 200  $\mu$ l fresh 80% EtOH to each well.
  - b Incubate on the magnetic stand for 30 seconds.
  - c Remove and discard all supernatant from each well.
- 8 Use a 20  $\mu$ l pipette to remove residual EtOH from each well.
- 9 Air dry on the magnetic stand for 5 minutes.
- 10 Add the appropriate volume of RSB to each well.
  - ▶ **Round 1** —52.5  $\mu$ l
  - ▶ **Round 2** —27.5  $\mu$ l
- 11 Remove from the magnetic stand, and then mix thoroughly as follows.
  - ▶ [HS] Shake at 1800 rpm for 2 minutes.

- ▶ [LS] Pipette up and down.
- 12 Incubate at room temperature for 2 minutes.
- 13 Centrifuge at 280 × g for 1 minute.
- 14 Place on a magnetic stand and wait until the liquid is clear (2–5 minutes).
- 15 Transfer 50 µl supernatant to the corresponding well of the CAP plate.
- 16 Repeat steps 1 through 14 using the new plate and the **Round 2** volumes.
- 17 Transfer 25 µl supernatant to the corresponding well of the PCR plate.

### SAFE STOPPING POINT

If you are stopping, seal the plate and store at -25°C to -15°C for up to 7 days.

## Enrich DNA Fragments

This step uses PCR to selectively enrich DNA fragments that have adapter molecules on both ends and to amplify the amount of DNA in the library. PCR is performed with PCR Primer Cocktail, which anneals to adapter ends. Minimize the number of PCR cycles to avoid skewing representation of the library.



### NOTE

Fragments without adapters cannot hybridize to the primers on the surface of the flow cell. Fragments with an adapter on one end can hybridize to the primers, but cannot form clusters.

### Consumables

- ▶ EPM (Enhanced PCR Mix)
- ▶ PPC (PCR Primer Cocktail)
- ▶ RSB (Resuspension Buffer)
- ▶ SPB (Sample Purification Beads)
- ▶ TSP 1 (Target Sample Plate) barcode label
- ▶ Freshly prepared 80% ethanol (EtOH)
- ▶ Plates
  - ▶ [HS] 96-well Hard-Shell 0.3 ml PCR plate (1)
  - ▶ [LS] 96-well 0.3 ml PCR plate, semiskirted or skirtless (1)
- ▶ [HS] Microseal 'A' film
- ▶ Microseal 'B' adhesive seals

### About Consumables

- ▶ Use Microseal 'A' when sealing the plate before placing it on the thermal cycler. Use Microseal 'B' for other steps that require a sealed plate.
- ▶ Vortex SPB before each use.
- ▶ Vortex SPB frequently to make sure that beads are evenly distributed.
- ▶ Aspirate and dispense SPB slowly due to the viscosity of the solution.

## Preparation

- 1 Prepare the following consumables.

Item	Storage	Instructions
PPC	-25°C to -15°C	Thaw at room temperature. Invert to mix, then centrifuge at 600 × g for 1 minute. Do not vortex. Return to storage after use.
EPM	-25°C to -15°C	Thaw on ice. Invert to mix, then centrifuge at 600 × g for 1 minute. Do not vortex. Return to storage after use.
RSB	2°C to 8°C	Let stand for 30 minutes to bring to room temperature.
SPB	2°C to 8°C	Let stand for 30 minutes to bring to room temperature.

- 2 Save the following PCRNano program on the thermal cycler:
  - ▶ Choose the preheat lid option and set to 100°C
  - ▶ 95°C for 3 minutes
  - ▶ Eight cycles of:
    - ▶ 98°C for 20 seconds
    - ▶ 60°C for 15 seconds
    - ▶ 72°C for 30 seconds
  - ▶ 72°C for 5 minutes
  - ▶ Hold at 4°C
- 3 Apply the TSP1 barcode label to a Hard-Shell PCR or PCR plate.

## Procedure

### Amplify DNA Fragments

- 1 Place the plate on ice and add 5 µl PPC to each well.
- 2 Add 20 µl EPM to each well.
- 3 Mix thoroughly as follows.
  - ▶ [HS] Shake at 1600 rpm for 2 minutes.
  - ▶ [LS] Pipette up and down.
- 4 Centrifuge at 280 × g for 1 minute.
- 5 Place on the thermal cycler and run the PCRNano program.  
Each well contains 50 µl.

### Clean Up Amplified DNA

- 1 Centrifuge at 280 × g for 1 minute.
- 2 Vortex SPB until well-dispersed.
- 3 Add the appropriate volume of SPB to each well:

Adapter Type	SPB Volume
Adapter tubes	50 µl
Index Adapter Plate	47.5 µl

- 4 Mix thoroughly as follows.
  - ▶ [HS] Shake at 1800 rpm for 2 minutes.
  - ▶ [LS] Pipette up and down.
- 5 Incubate at room temperature for 5 minutes.
- 6 Centrifuge at  $280 \times g$  for 1 minute.
- 7 Place on a magnetic stand and wait until the liquid is clear (2–5 minutes).
- 8 Remove and discard all supernatant from each well.
- 9 Wash two times as follows.
  - a Add 200  $\mu$ l fresh 80% EtOH to each well.
  - b Incubate on the magnetic stand for 30 seconds.
  - c Remove and discard all supernatant from each well.
- 10 Use a 20  $\mu$ l pipette to remove residual EtOH from each well.
- 11 Air-dry on the magnetic stand for 5 minutes.
- 12 Add 32.5  $\mu$ l RSB to each well, and then remove from the magnetic stand.
- 13 Mix thoroughly as follows.
  - ▶ [HS] Shake at 1800 rpm for 2 minutes.
  - ▶ [LS] Pipette up and down.
- 14 Incubate at room temperature for 2 minutes.
- 15 Centrifuge at  $280 \times g$  for 1 minute.
- 16 Place on a magnetic stand and wait until the liquid is clear (2–5 minutes).
- 17 Transfer 30  $\mu$ l supernatant to the corresponding well of the TSP1 plate.

#### SAFE STOPPING POINT

If you are stopping, seal the plate and store at  $-25^{\circ}\text{C}$  to  $-15^{\circ}\text{C}$  for up to 7 days.

## Check Libraries

### Quantify Libraries

Achieving high-quality data on Illumina sequencing systems requires optimum cluster density across every lane of the flow cell. Optimizing cluster densities requires accurate quantification of DNA libraries.

- 1 Quantify libraries with a fluorometric method that uses dsDNA binding dyes or qPCR.
- 2 [Optional] If you are using the KAPA Library Quantification Kit – Illumina/Universal, follow the KAPA instructions with the KAPA standard. To calculate the library concentration in nM, make the following insert size adjustments:
  - ▶ For 350 bp libraries, use 470 bp for the average fragment length.
  - ▶ For 550 bp libraries, use 670 bp for the average fragment length.

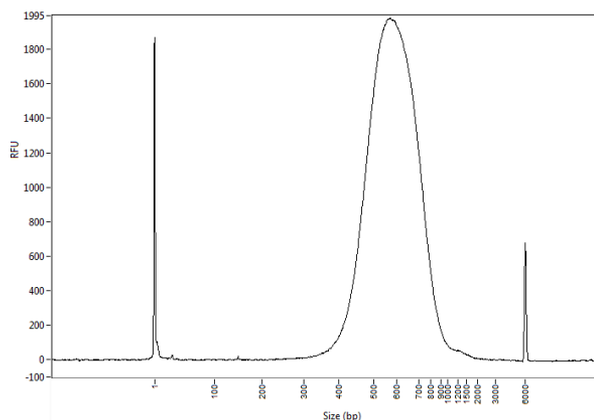
Quantification of TruSeq DNA Nano Library Prep libraries has been validated with the KAPA Library Quantification Kit – Illumina/Universal. The data sheet for this kit is available on the KAPA Biosystems website.

## Check Library Quality

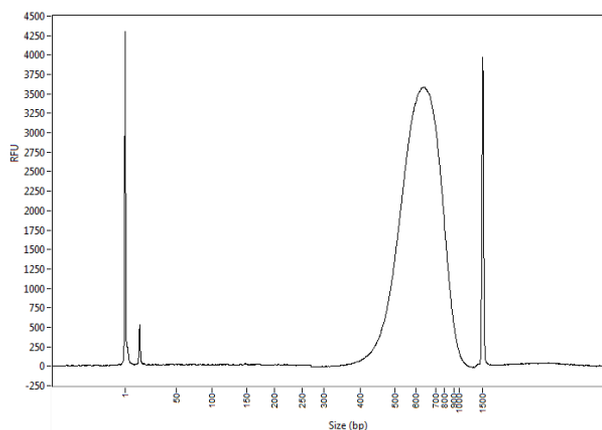
Verify fragment size by checking the library size distribution. Run on an Advanced Analytical Fragment Analyzer with the HS-NGS High Sensitivity 474 kit or an Agilent Technology 2100 Bioanalyzer using a High Sensitivity DNA kit.

- 1 If you are using a High Sensitivity DNA chip, dilute the DNA library 1:10 with water.
- 2 Run the library on the Advanced Analytical Fragment Analyzer or Agilent Technology 2100 Bioanalyzer:
  - ▶ For a High Sensitivity DNA or NGS Kit, run 1  $\mu$ l diluted DNA library.
  - ▶ For a Bioanalyzer DNA 7500 chip, run 1  $\mu$ l undiluted DNA library.

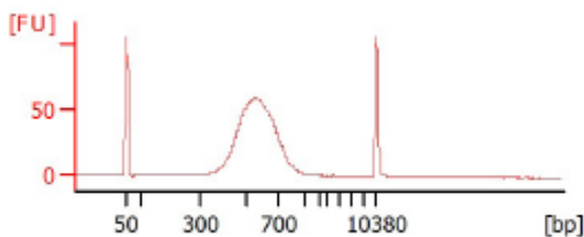
**Figure 2** Example Distribution of 350bp Library Run on Fragment Analyzer Using High Sensitivity NGS Kit

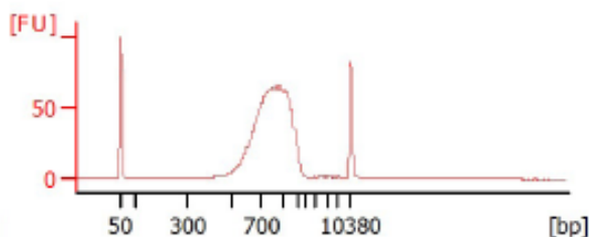


**Figure 3** Example Distribution of 550bp Library Run on Fragment Analyzer Using High Sensitivity NGS Kit



**Figure 4** Example Distribution of 350bp Library Run on Bioanalyzer Using High Sensitivity DNA Kit



**Figure 5** Example Distribution of 550bp Library Run on Bioanalyzer Using High Sensitivity DNA Kit

## Normalize and Pool Libraries

This step prepares DNA template for cluster generation. Non-indexed DNA libraries are normalized to 10 nM in the DCT plate. Indexed DNA libraries are normalized to 10 nM in the DCT plate and then pooled in equal volumes in the PDP plate.



### NOTE

For best practice, perform normalization and pooling directly prior to sequencing. To minimize index hopping, do not store libraries in the pooled form. For more information, see *Minimize index hopping in multiplexed runs* on the Illumina website.

## Consumables

- ▶ Barcode labels
  - ▶ DCT (Diluted Cluster Template)
  - ▶ PDP (Pooled DCT Plate) (for pooling only)
- ▶ Tris-HCl 10 mM, pH8.5 with 0.1% Tween 20
- ▶ Plates
  - ▶ [LS] 96-well 0.3 ml PCR plate, semiskirted or skirtless (1) (for pooling  $\leq$  40 samples)
- ▶ Microseal 'B' adhesive seals

## Preparation

- 1 Apply a DCT barcode label to a midi plate.
- 2 [For pooling only] Apply a PDP barcode label to a midi plate ( $>$  40 samples) or PCR plate ( $\leq$  40 samples).

## Procedure

### Normalize Libraries

- 1 Transfer 10  $\mu$ l library to the corresponding well of the DCT plate.
- 2 Normalize the library concentration to 10 nM using Tris-HCl 10 mM, pH8.5 with 0.1% Tween 20.
- 3 Mix thoroughly as follows.
  - ▶ [HS] Shake at 1000 rpm for 2 minutes.
  - ▶ [LS] Pipette up and down.
 Depending on the yield quantification data of each library, the final volume of each well can vary from 10–400  $\mu$ l.

- 4 Centrifuge at  $280 \times g$  for 1 minute.
- 5 Do the following:
  - ▶ To pool libraries, proceed to *Pool Libraries*.
  - ▶ Libraries that are not pooled must be diluted and denatured before proceeding to cluster generation. For more information, see the Dilute and Denature guide for your Illumina platform.

## Pool Libraries

The pooling procedure depends on the number of libraries being pooled 2–24, 25–48, or 49–96.

### Pool 2–24 Libraries

- 1 Transfer 10  $\mu$ l of each normalized library to one well of the PDP plate.
- 2 Mix thoroughly as follows.
  - ▶ [HS] Shake at 1800 rpm for 2 minutes.
  - ▶ [LS] Pipette up and down.
- 3 Centrifuge at  $280 \times g$  for 1 minute.
- 4 Proceed to cluster generation.  
For instructions, see the system guide for your Illumina instrument.

### Pool 25–48 Libraries

- 1 Transfer 5  $\mu$ l of each column of normalized library to column 1 of the PDP plate.
- 2 Mix thoroughly as follows.
  - ▶ [HS] Shake at 1800 rpm for 2 minutes.
  - ▶ [LS] Pipette up and down.
- 3 Centrifuge at  $280 \times g$  for 1 minute.
- 4 Transfer the contents from each well of column 1 to well A2.
- 5 Mix thoroughly as follows.
  - ▶ [HS] Shake at 1800 rpm for 2 minutes.
  - ▶ [LS] Pipette up and down.
- 6 Centrifuge at  $280 \times g$  for 1 minute.
- 7 Proceed to cluster generation.  
For instructions, see the system guide for your Illumina instrument.

### Pool 49–96 Libraries

- 1 Transfer 5  $\mu$ l of each column of normalized library to column 1 of the PDP plate.
- 2 Mix thoroughly as follows.
  - ▶ [HS] Shake at 1800 rpm for 2 minutes.
  - ▶ [LS] Pipette up and down.
- 3 Centrifuge at  $280 \times g$  for 1 minute.
- 4 Transfer the contents of each well of column 1 to a 1.7 ml microcentrifuge tube.
- 5 Mix thoroughly as follows.

- ▶ [HS] Shake at 1800 rpm for 2 minutes or vortex the tube.
  - ▶ [LS] Pipette up and down.
- 6 Centrifuge at  $280 \times g$  for 1 minute.
  - 7 Proceed to cluster generation.  
For instructions, see the system guide for your Illumina instrument.

## SAFE STOPPING POINT

If you are stopping, seal the plate or cap the tube and store at  $-25^{\circ}\text{C}$  to  $-15^{\circ}\text{C}$  for up to 7 days.

# Supporting Information

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## Product Contents

Make sure that you have all reagents identified in this section before starting the protocol.

The following library prep and index adapter components are available to order through Illumina to support the TruSeq DNA Nano Library Prep workflow.

From Illumina, order one catalog number for the library prep component and one catalog number for the index adapter component depending on the number of samples for your experiment.

Library Prep Component	Catalog #
TruSeq DNA Nano Library Prep Library Prep (24 Samples)	20015964
TruSeq DNA Nano Library Prep Library Prep (96 Samples)	20015965

Index Adapter Component	Catalog #
IDT for Illumina - TruSeq DNA UD Indexes (24 Indexes, 96 Samples)	20020590
IDT for Illumina - TruSeq DNA UD Indexes (96 Indexes, 96 Samples)	20022370
TruSeq DNA Single Indexes (12 indexes, 24 samples) Set A	20015960
TruSeq DNA Single Indexes (12 indexes, 24 samples) Set B	20015961
TruSeq DNA Combinatorial Dual Indexes (96 indexes, 96 samples)	20015949

## TruSeq DNA Nano Library Prep Library Prep (24 Samples)

This workflow contains two boxes: Box 1 and an SPB (Sample Purification Beads) box.

### Box 1, Store at -25°C to -15°C

Quantity	Reagent	Description
1	RSB	Resuspension Buffer
1	ERP 2 or ERP 3	End Repair Mix
1	ATL or ATL 2	A-Tailing Mix
1	LIG 2	Ligation Mix 2
1	STL	Stop Ligation Buffer
1	PPC	PCR Primer Cocktail
1	EPM	Enhanced PCR Mix

### SPB Box, Store at 2°C to 8°C

Quantity	Reagent	Description
1	SPB	Sample Purification Beads

## TruSeq DNA Nano Library Prep Library Prep (96 Samples)

This workflow contains two boxes: Box 1 and an SPB (Sample Purification Beads) box.

### Box 1, Store at -25°C to -15°C

This box also contains plate barcode labels.

Quantity	Reagent	Description
2	RSB	Resuspension Buffer
2	ERP 2 or ERP 3	End Repair Mix
2	ATL or ATL 2	A-Tailing Mix
2	LIG 2	Ligation Mix 2
2	STL	Stop Ligation Buffer
2	PPC	PCR Primer Cocktail
2	EPM	Enhanced PCR Mix

### SPB Box, Store at 2°C to 8°C

Quantity	Reagent	Description
4	SPB	Sample Purification Beads

## Consumables and Equipment

Make sure that you have the required user-supplied consumables and equipment before starting the protocol. Items that are unique to the HS or LS workflow are indicated.

The protocol has been optimized and validated using the items listed. Comparable performance is not guaranteed when using alternate consumables and equipment.

## Consumables

Consumable	Supplier
1.7 ml microcentrifuge tubes	General lab supplier
15 ml conical tubes	General lab supplier
10 µl barrier pipette tips	General lab supplier
10 µl multichannel pipettes	General lab supplier
10 µl single channel pipettes	General lab supplier
20 µl barrier pipette tips	General lab supplier
20 µl multichannel pipettes	General lab supplier
20 µl single channel pipettes	General lab supplier
200 µl barrier pipette tips	General lab supplier
200 µl multichannel pipettes	General lab supplier
200 µl single channel pipettes	General lab supplier
1000 µl barrier pipette tips	General lab supplier

Consumable	Supplier
1000 µl multichannel pipettes	General lab supplier
1000 µl single channel pipettes	General lab supplier
96-well storage plates, round well, 0.8 ml (midi plate)	Thermo Fisher Scientific, part # AB-0859
Adhesive seal roller	General lab supplier
Ethanol 200 proof (absolute) for molecular biology (500 ml)	Sigma-Aldrich, part E7023
Ice bucket	General lab supplier
[Optional] KAPA Library Quantification Kit - Illumina/Universal	KAPA Biosystems, part # KK4824
Microseal 'B' adhesive seals	Bio-Rad, part # MSB-1001
microTUBE AFA Fiber 6x16mm with: <ul style="list-style-type: none"> <li>• Crimp-Cap or</li> <li>• Pre-Slit Snap-Cap (for use with Covaris M220)</li> </ul>	Covaris, part # <ul style="list-style-type: none"> <li>• 520052 or</li> <li>• 520045</li> </ul>
PCR-grade water	General lab supplier
Qubit assay tubes or Axygen PCR-05-C tubes	Thermo Fisher Scientific, catalog # Q32856 or VWR, part # 10011 -830
RNase/DNase-free 8-tube strips and caps	General lab supplier
RNase/DNase-free multichannel reagent reservoirs, disposable	VWR, part # 89094-658
Tris-HCl 10 mM, pH 8.5	General lab supplier
Tween 20	Sigma-Aldrich, part # P7949

## Additional Consumables for HS Workflow

Consumable	Supplier
96-well Hard-Shell 0.3 ml PCR plate	Bio-Rad, part # HSP-9601

## Additional Consumables for LS Workflow

Consumable	Supplier
96-well 0.3 ml skirtless PCR plates or Twin.tec 96 well PCR plates	E&K Scientific, part # 480096 or Eppendorf, part # 951020303

## Equipment

Equipment	Supplier
[Optional] Fragment Analyzer™	Advanced Analytical, catalog # FSV2CE2F
[Optional] 2100 Bioanalyzer Desktop System	Agilent Technologies, part # G2940CA
One of the following Covaris systems: <ul style="list-style-type: none"> <li>• S2</li> <li>• S220</li> <li>• E210</li> <li>• M220</li> </ul>	Covaris M220, part # 500295*
Magnetic stand-96	Thermo Fisher Scientific, catalog # AM10027
Microplate centrifuge	General lab supplier
Vortexer	General lab supplier
qPCR system	General lab supplier

\* Contact Covaris for all other models.

## Additional Equipment for HS Workflow

Equipment	Supplier
High-Speed Microplate Shaker	WWR, catalog # • 13500-890 (110 V/120 V) or • 14216-214 (230 V)
SciGene TruTemp Heating System <sup>1</sup>	Illumina, catalog # • SC-60-503 (110 V) or • SC-60-504 (220 V)
Midi plate insert for heating system <sup>2</sup>	Illumina, catalog # BD-60-601
Stroboscope	General lab supplier

<sup>1</sup> Two systems are recommended to support successive heating procedures.

<sup>2</sup> Two inserts are recommended to support successive heating procedures.

## Thermal Cyclers

The following table lists the recommended specifications for the thermal cycler. If your lab has a thermal cycler that is not listed, validate it before starting the protocol.

Thermal Cycler	Temp Mode	Lid Temp	Vessel Type
Bio-Rad DNA Engine Tetrad 2	Calculated	Heated, constant at 100°C	Plate
MJ Research PTC-225 DNA Engine Tetrad	Calculated	Heated, constant at 100°C	Plate
Bio-Rad S1000	N/A	Heated, constant at 100°C	Plate

## qPCR Systems

The following table lists the validated qPCR systems for the TruSeq DNA Nano Library Prep protocol.

Equipment	Supplier
CFX96 Touch Real-Time PCR Detection System*	Bio-Rad, part # 185-5195
Mx3000P qPCR System	Agilent, part # 401511

\* Use CFX Manager software version 3.0 with Cq Determination mode: Single Threshold; Baseline Setting: Baseline Subtracted Curve Fit and Apply Fluorescent Drift Correction for data analysis. This setting can correct for abnormalities in fluorescence intensity of the standard curve caused by the instrument. For software installation, contact Bio-Rad.

## Index Adapter Sequences

For information on index adapter sequences, see [Illumina Adapter Sequences \(document # 1000000002694\)](#) which provides information regarding the nucleotide sequences that comprise Illumina oligonucleotides used in Illumina sequencing technologies.

## Acronyms

Acronym	Definition
ALP	Adapter Ligation Plate
ATL	A-Tailing Mix
CAP	Clean Up ALP Plate
CEP	Clean Up End Repair Plate
CFP	Covaris Fragmentation Plate
CSP	Clean Up Sheared DNA Plate
DCT	Diluted Cluster Template Plate
DNA	Customer Sample DNA Plate
ERP	End Repair Mix
HS	High Sample
IEM	Illumina Experiment Manager
IMP	Insert Modification Plate
LIG	Ligation Mix
LRM	Local Run Manager
LS	Low Sample
PDP	Pooled Dilution Plate
RSB	Resuspension Buffer
SPB	Sample Purification Beads
STL	Stop Ligation Buffer
TSP1	Target Sample Plate 1

# Technical Assistance

For technical assistance, contact Illumina Technical Support.

Website: [www.illumina.com](http://www.illumina.com)  
Email: [techsupport@illumina.com](mailto:techsupport@illumina.com)

## Illumina Customer Support Telephone Numbers

Region	Toll Free	Regional
North America	+1.800.809.4566	
Australia	+1.800.775.688	
Austria	+43 800006249	+43 19286540
Belgium	+32 80077160	+32 34002973
China	400.635.9898	
Denmark	+45 80820183	+45 89871156
Finland	+358 800918363	+358 974790110
France	+33 805102193	+33 170770446
Germany	+49 8001014940	+49 8938035677
Hong Kong	800960230	
Ireland	+353 1800936608	+353 016950506
Italy	+39 800985513	+39 236003759
Japan	0800.111.5011	
Netherlands	+31 8000222493	+31 207132960
New Zealand	0800.451.650	
Norway	+47 800 16836	+47 21939693
Singapore	+1.800.579.2745	
Spain	+34 911899417	+34 800300143
Sweden	+46 850619671	+46 200883979
Switzerland	+41 565800000	+41 800200442
Taiwan	00806651752	
United Kingdom	+44 8000126019	+44 2073057197
Other countries	+44.1799.534000	

**Safety data sheets (SDSs)**—Available on the Illumina website at [support.illumina.com/sds.html](http://support.illumina.com/sds.html).

**Product documentation**—Available for download in PDF from the Illumina website. Go to [support.illumina.com](http://support.illumina.com), select a product, then select **Documentation & Literature**.



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