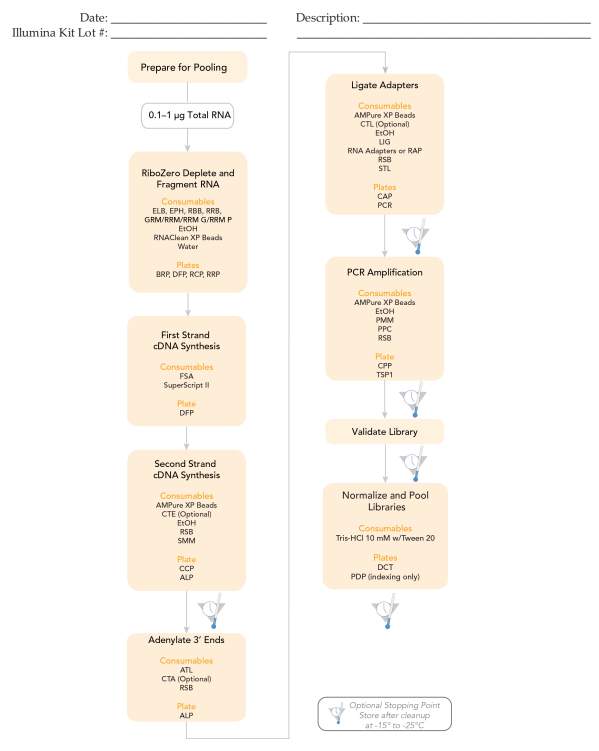
Experienced User Card and Lab Tracking Form

FOR RESEARCH USE ONLY





NOTE

Unless familiar with the HS protocol in the latest version of the *TruSeq Stranded Total RNA Sample Preparation Guide* (part # 15031048), new or less experienced users are advised to follow the protocol in the guide before using this Experienced User Card and Lab Tracking Form.





Consumables

TruSeq Stranded Total RNA Sample Prep HS Protocol

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| Date/Time: | Operator: | |
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Consumables

| Item | Lot Number |
|--|----------------|
| A-Tailing Control (CTA) | Lot #: |
| A-Tailing Mix (ATL) | Lot #: |
| Elute, Prime, Fragment High Mix (EPH) | Lot #: |
| Elution Buffer (ELB) | Lot #: |
| End Repair Control (CTE) | Lot #: |
| First Strand Synthesis Act D Mix (FSA) | Lot #: |
| Globin Removal Mix (GRM) | Lot #: |
| Ligation Control (CTL) | Lot #: |
| Ligation Mix (LIG) | Lot #: |
| PCR Master Mix (PMM) | Lot #: |
| PCR Primer Cocktail (PPC) | Lot #: |
| Resuspension Buffer (RSB) | Lot #: |
| rRNA Binding Buffer (RBB) | Lot #: |
| rRNA Removal Beads (RRB) | Lot #: |
| rRNA Removal Mix (RRM) | Lot #: |
| rRNA Removal Mix Gold (RRM G) | Lot #: |
| rRNA Removal Mix Plant (RRM P) | Lot #: |
| Second Strand Marking Master Mix (SMM) | Lot #: |
| Stop Ligation Buffer (STL) | Lot #: |
| 80% Ethanol | Date Prepared: |
| Adapter Indices or RAP | Lot Number |
| RNA Adapter Index 1 (AR001) | Lot #: |
| RNA Adapter Index 2 (AR002) | Lot #: |
| RNA Adapter Index 3 (AR003) | Lot #: |
| RNA Adapter Index 4 (AR004) | Lot #: |
| RNA Adapter Index 5 (AR005) | Lot #: |
| RNA Adapter Index 6 (AR006) | Lot #: |

| Date/Time: | Operator: |
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| Adapter Indices or RAP | Lot Number |
|---------------------------------|------------|
| RNA Adapter Index 7 (AR007) | Lot #: |
| RNA Adapter Index 8 (AR008) | Lot #: |
| RNA Adapter Index 9 (AR009) | Lot #: |
| RNA Adapter Index 10 (AR010) | Lot #: |
| RNA Adapter Index 11 (AR011) | Lot #: |
| RNA Adapter Index 12 (AR012) | Lot #: |
| RNA Adapter Index 13 (AR013) | Lot #: |
| RNA Adapter Index 14 (AR014) | Lot #: |
| RNA Adapter Index 15 (AR015) | Lot #: |
| RNA Adapter Index 16 (AR016) | Lot #: |
| RNA Adapter Index 18 (AR018) | Lot #: |
| RNA Adapter Index 19 (AR019) | Lot #: |
| RNA Adapter Index 20 (AR020) | Lot #: |
| RNA Adapter Index 21 (AR021) | Lot #: |
| RNA Adapter Index 22 (AR022) | Lot #: |
| RNA Adapter Index 23 (AR023) | Lot #: |
| RNA Adapter Index 24 (AR024) | Lot #: |
| RNA Adapter Index 25 (AR025) | Lot #: |
| RNA Adapter Index 27 (AR027) | Lot #: |
| RNA Adapter Plate, 96plex (RAP) | Lot #: |



Ribo-Zero™ Deplete and Fragment RNA

TruSeq Stranded Total RNA Sample Prep HS Protocol

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$\mathsf{Ribo}\text{-}\mathsf{Zero}^{^\mathsf{TM}}\,\mathsf{Deplete}\,\mathsf{and}\,\mathsf{Fragment}\,\mathsf{RNA}$

This process depletes rRNA from total RNA. After the rRNA is depleted, the remaining RNA is purified, fragmented, and primed for cDNA synthesis.

| Item | Quantity | Storage | Supplied By |
|--|---|----------------|-------------|
| Elute, Prime, Fragment High Mix (EPH) | 1 tube per 48 reactions | -15°C to -25°C | Illumina |
| Elution Buffer (ELB) | 1 tube per 48 reactions | 2°C to 8°C | Illumina |
| One of the following, depending on the kit you are using: • Globin Removal Mix (GRM) • rRNA Removal Mix (RRM) • rRNA Removal Mix - Gold (RRM G) • rRNA Removal Mix - Plant (RRM P) | 1 tube per 48 reactions | -15°C to -25°C | Illumina |
| Resuspension Buffer (RSB) | 1 tube | -15°C to -25°C | Illumina |
| rRNA Binding Buffer (RBB) | 1 tube per 48 reactions | -15°C to -25°C | Illumina |
| rRNA Removal Beads (RRB) | 1 tube per 48 reactions | 2°C to 8°C | Illumina |
| Barcode labels for: BRP (Bind rRNA Plate) DFP (Depleted RNA Fragmentation Plate) RCP (RNA Clean Up Plate) RRP (rRNA Removal Plate) | 1 label per plate | 15°C to 30°C | Illumina |
| 96-well HSP Plates | 2 | 15°C to 30°C | User |
| 96-well MIDI Plates | 2 | 15°C to 30°C | User |
| Freshly Prepared 70% Ethanol (EtOH) | 200 μl per sample | 15°C to 30°C | User |
| Microseal 'B' Adhesive Seals | 5 | 15°C to 30°C | User |
| RNAClean XP Beads | 99 µl per sample | 2°C to 8°C | User |
| RNase/DNase-free Eight-Tube Strips and Caps (if using multichannel pipettes) | 6 | 15°C to 30°C | User |
| RNase/DNase-free Reagent Reservoirs (if using multichannel pipettes) | 6 | 15°C to 30°C | User |
| Ultra Pure Water | Enough to dilute each total RNA sample to a final volume of 10 µl | 15°C to 30°C | User |

| Date/Time: | Operator: |
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| Make BRP | |
| | |
| [_] 1 | Dilute the total RNA with nuclease-free ultra pure water to a final volume of $10 \mu l$ in the new 96-well HSP plate labeled with the BRP barcode. |
| [_] 2 | Add 5 μ l of rRNA Binding Buffer to each well of the BRP plate. |
| [_] 3 | Add 5 µl of one of the following reagents to each well of the BRP plate, depending on the kit you are using: • Globin Removal Mix • rRNA Removal Mix • rRNA Removal Mix - Gold • rRNA Removal Mix - Plant |
| [_] 4 | Mix the contents of the BRP plate thoroughly as follows: |
| | a Seal the BRP plate with a Microseal 'B' adhesive seal. |
| [_] | |
| [_] 5 | Centrifuge the BRP plate to 280 × g for 1 minute. |
| [_] 6 | Return the following to -15°C to -25°C storage: • rRNA Binding Buffer |
| | One of the following, depending on the kit you are using: Globin Removal Mix rRNA Removal Mix rRNA Removal Mix - Gold rRNA Removal Mix - Plant |
| Incubate 1 | BRP |
| [_] 1 [_] [_] | Place the sealed BRP plate on the pre-programmed thermal cycler. Close the lid, then select and run the RNA Denaturation program. a Choose the pre-heat lid option and set to 100°C b 68°C for 5 minutes |
| [_] 2 | After the 5 minute incubation, place the BRP plate on the bench and incubate at room temperature for 1 minute. |
| Make RRP | |
| [_] 1 | Vortex the room temperature rRNA Removal Bead tube vigorously to resuspend the beads. |
| [_] 2 | Add 35 μl of rRNA Removal Beads to each well of the new 96-well MIDI plate labeled with the RRP barcode. |
| [_] 3 | Remove the adhesive seal from the BRP plate. |
| [_] 4 | Transfer the entire contents from each well of the BRP plate to the corresponding well of the RRP plate containing rRNA Removal Beads. |
| [_] 5 [_] [_] | Mix the contents of the RRP plate thoroughly as follows: a Seal the RRP plate with a Microseal 'B' adhesive seal. b Shake the RRP plate on a microplate shaker continuously at 1000 rpm for 1 minute. |



| Date/Time | : Operator: |
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| [_] 6 | Remove the adhesive seal from the RRP plate. |
| [_] 7 | Place the RRP plate on the magnetic stand at room temperature for 1 minute. |
| [_] 8 | Transfer all of the supernatant from each well of the RRP plate to the corresponding well of |
| [_] 0 | the new 96-well MIDI plate labeled with the RCP barcode. |
| [_] 9 | Place the RCP plate on the magnetic stand at room temperature for 1 minute. |
| [_] 10 | Return the rRNA Removal Beads to 2°C to 8°C storage. |
| Clean Up R | CP |
| [_] 1 | Vortex the RNAClean XP beads until they are well dispersed, then add 99 μ l of well-mixed RNAClean XP beads to each well of the RCP plate containing ribosomal depleted RNA. Mix thoroughly as follows: |
| 4 | NOTE If starting with degraded total RNA, add 193 µl of well-mixed RNAClean XP beads to each well of the RCP plate containing ribosomal depleted RNA. |
| | a Seal the RCP plate with a Microseal 'B' adhesive seal. |
| | b Shake the RCP plate on a microplate shaker at 1800 rpm for 2 minutes. |
| [_] 2 | Incubate the RCP plate at room temperature for 15 minutes. Start time: Stop time: |
| [_] 3 | Remove the adhesive seal from the RCP plate. |
| [_] 4 | Place the RCP plate on the magnetic stand at room temperature, for 5 minutes to make sure |
| [_] - | that all of the beads are bound to the side of the wells. |
| | Start time: Stop time: |
| [_] 5 | Remove and discard all of the supernatant from each well of the RCP plate. |
| [_] 6 | With the RCP plate on the magnetic stand, add 200 µl freshly prepared 70% EtOH to each well without disturbing the beads. |
| [_] 7 | Incubate the RCP plate at room temperature for 30 seconds, and then remove and discard all of the supernatant from each well. |
| [_] 8 | Let the RCP plate stand at room temperature for 15 minutes to dry, and then remove the RCP plate from the magnetic stand. Start time: Stop time: |
| [_] 9 | Centrifuge the thawed, room temperature Elution Buffer to 600 × g for 5 seconds. |
| [_] 10 [_ [_ | Add 11 µl Elution Buffer to each well of the RCP plate. Mix thoroughly as follows:] a Seal the RCP plate with a Microseal 'B' adhesive seal.] b Shake the RCP plate on a microplate shaker at 1800 rpm for 2 minutes. |
| [_] 11 | Incubate the RCP plate at room temperature for 2 minutes. Start time: Stop time: |
| [_] 12 | Centrifuge the RCP plate to 280 × g for 1 minute. |
| [_] 13 | |
| | |

Incub

TruSeq Stranded Total RNA Sample Prep HS Protocol

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| [_] 14 | Place the RCP plate on the magnetic stand at room temperature for 5 minutes. Start time: Stop time: |
| [_] 15 | Return the Elution Buffer to 2°C to 8°C storage. |
| [_] 16 | Transfer 8.5 μl supernatant from the RCP plate to the new 96-well HSP plate labeled with the DFP barcode. |
| [_] 17 | Add 8.5 μl Elute, Prime, Fragment High Mix to each well of the DFP plate. Mix thoroughly as follows: |
| [_] | 1 |
| [_] 18 | Return the Elute, Prime, Fragment High Mix to -15°C to -25°C storage and the RNAClean XF Beads tube to 2°C to 8°C storage. |
| ate 1 E |)FP |
| [_] 1 [_] [_] [_] | b 94°C for 8 minutes |
| [_] 2 | Remove the DFP plate from the thermal cycler when it reaches 4°C and centrifuge briefly. |
| [_] 3 | Proceed immediately to Synthesize First Strand cDNA on page 9. |
| Co | pmments |



Synthesize First Strand cDNA

TruSeq Stranded Total RNA Sample Prep HS Protocol

Experienced User Card and Lab Tracking Form

| Date/Time: | Operator: | |
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Synthesize First Strand cDNA

This process reverse transcribes the cleaved RNA fragments that were primed with random hexamers into first strand cDNA using reverse transcriptase and random primers. The addition of Actinomycin D to the First Stand Synthesis Act D mix (FSA) prevents spurious DNA-dependent synthesis, while allowing RNA-dependent synthesis, improving strand specificity.

Consumables

| Item | Quantity | Storage | Supplied By |
|--|----------|----------------|-------------|
| First Strand Synthesis Act D Mix (FSA) | 1 tube | -15°C to -25°C | Illumina |
| Microseal 'B' Adhesive Seal | 1 | 15°C to 30°C | User |
| RNase/DNase-free Eight-Tube Strips and Caps (if using multichannel pipettes) | 1 | 15°C to 30°C | User |
| RNase/DNase-free Reagent Reservoirs (if using multichannel pipettes) | 1 | 15°C to 30°C | User |
| SuperScript II Reverse Transcriptase | 1 tube | -15°C to -25°C | User |

Add FSA

| [_] 1 | Remove the adhesive seal from the DFP plate. |
|---------------------|--|
| [_] 2 | Centrifuge the thawed First Strand Synthesis Mix Act D tube to $600 \times g$ for 5 seconds. |
| [_] 3 | Add 50 μ l SuperScript II to the First Strand Synthesis Act D Mix tube. Mix gently, but thoroughly and centrifuge briefly. If you are not using the entire contents of the First Strand Synthesis Act D Mix tube, add SuperScript II at a ratio of 1 μ l SuperScript II for each 9 μ l First Strand Synthesis Act D Mix. Label the First Strand Synthesis Mix Act D tube to indicate that the SuperScript II has been added. |
| [_] 4 [_] [_] | 1 |
| [_] 5 | Return the First Strand Synthesis Mix Act D tube to -15°C to -25°C storage immediately after use. |

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Date/Time: _______ Operator: _______

Incubate 2 DFP

[_] 1 Place the sealed DFP plate on the pre-programmed thermal cycler. Close the lid and select Synthesize 1st Strand.

[_] a Choose the pre-heat lid option and set to 100°C

[_] b 25°C for 10 minutes

[_] c 42°C for 15 minutes

[_] d 70°C for 15 minutes

[_] e Hold at 4°C

[_] 2 When the thermal cycler reaches 4°C, remove the DFP plate from the thermal cycler and proceed immediately to Synthesize Second Strand cDNA on page 11.

Comments



Synthesize Second Strand cDNA

TruSeq Stranded Total RNA Sample Prep HS Protocol

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Synthesize Second Strand cDNA

This process removes the RNA template and synthesizes a replacement strand, incorporating dUTP in place of dTTP to generate ds cDNA. The incorporation of dUTP quenches the second strand during amplification, because the polymerase does not incorporate past this nucleotide. AMPure XP beads are used to separate the ds cDNA from the second strand reaction mix. At the end of this process, you have blunt-ended cDNA.

Consumables

| Item | Quantity | Storage | Supplied By |
|---|-------------------------|----------------|-------------|
| (Optional) End Repair Control (CTE) | 1 tube per 48 reactions | 2°C to 8°C | Illumina |
| Resuspension Buffer (RSB) | 1 tube | 2°C to 8°C | Illumina |
| Second Strand Marking Master Mix (SMM) | 1 tube per 48 reactions | -15°C to -25°C | Illumina |
| Barcode labels for: • ALP (Adapter Ligation Plate) • CCP (cDNA Clean Up Plate) • IMP (Insert Modification Plate) | 1 label per plate | 15°C to 30°C | Illumina |
| 96-well MIDI Plates | 2 | 15°C to 30°C | User |
| AMPure XP Beads | 90 μl per sample | 2°C to 8°C | User |
| Freshly Prepared 80% Ethanol (EtOH) | 400 μl per sample | 15°C to 30°C | User |
| Microseal 'B' Adhesive Seals | 4 | 15°C to 30°C | User |
| RNase/DNase-free Eight-Tube Strips and Caps (if using multichannel pipettes) | 5 | 15°C to 30°C | User |
| RNase/DNase-free Reagent Reservoirs (if using multichannel pipettes) | 5 | 15°C to 30°C | User |

Add SMM

- [] 1 Remove the adhesive seal from the DFP plate.
- [_] 2 Do one of the following:
 - If using the in-line control reagent:
 - Centrifuge the thawed End Repair Control tube to $600 \times g$ for 5 seconds.
 - Dilute the End Repair Control to 1/50 in Resuspension Buffer (For example, 2 μ l End Repair Control + 98 μ l Resuspension Buffer) before use.
 - Add 5 μl of diluted End Repair Control to each well of the DFP plate.
 - If not using the in-line control reagent, add 5 μ l of Resuspension Buffer to each well of the DFP plate.



| Date/Time: | Operator: |
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| [_] 3 | Centrifuge the thawed Second Strand Marking Master Mix to 600 × g for 5 seconds. |
| [_] 4 [_] [_] | Add 20 μ l of thawed Second Strand Marking Master Mix to each well of the DFP plate. Mix thoroughly as follows: a Seal the DFP plate with a Microseal 'B' adhesive seal. |
| [_] 5 | Return the Second Strand Marking Master Mix tube to -15°C to -25°C storage after use. |
| Incubate 3 [| OFP |
| [_] 1 | Place the sealed DFP plate on the pre-heated thermal cycler. Close the lid and incubate at 16°C for 1 hour. |
| | Start time: Stop time: |
| [_] 2 | Remove the DFP plate from the thermal cycler and place it on the bench. |
| [_] 3 | Remove the adhesive seal from the DFP plate. |
| [_] 4 | Let the DFP plate stand to bring it to room temperature. |
| Clean Up DI | FP |
| [_] 1 | Vortex the AMPure XP beads until they are well dispersed. |
| [_] 2 | Add 90 μl of well-mixed AMPure XP beads to each well of the new MIDI plate labeled with the CCP barcode. |
| [_] 3 [_] [_] | |
| [_] 4 | Incubate the CCP plate at room temperature for 15 minutes. |
| | Start time: Stop time: |
| [_] 5 | Centrifuge the CCP plate to 280 × g for 1 minute. |
| [_] 6 | Remove the adhesive seal from the CCP plate. |
| [_] 7 | Place the CCP plate on the magnetic stand at room temperature, for 5 minutes to make sure that all of the beads are bound to the side of the wells. Start time: Stop time: |
| [_] 8 | Remove and discard 135 µl supernatant from each well of the CCP plate. |
| [_] 9 | With the CCP plate on the magnetic stand, add 200 μl freshly prepared 80% EtOH to each well without disturbing the beads. |
| [_] 10 | Incubate the CCP plate at room temperature for 30 seconds, and then remove and discard all of the supernatant from each well. |
| [_] 11 | Repeat steps 9 and 10 one time for a total of two 80% EtOH washes. |
| [_] 12 | Let the CCP plate stand at room temperature for 15 minutes to dry, and then remove the CCP plate from the magnetic stand. |
| | Start time: Stop time: |



Synthesize Second Strand cDNA

TruSeq Stranded Total RNA Sample Prep HS Protocol

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| | 1 |
| [_] 15 | Incubate the CCP plate at room temperature for 2 minutes. Start time: Stop time: |
| [_] 16 | Centrifuge the CCP plate to 280 × g for 1 minute. |
| [_] 17 | Remove the adhesive seal from the CCP plate. |
| [_] 18 | Place the CCP plate on the magnetic stand at room temperature for 5 minutes. Start time: Stop time: |
| | Transfer 15 μ l supernatant (ds cDNA) from the CCP plate to the new MIDI plate labeled with the ALP barcode. |
| Q | SAFE STOPPING POINT If you do not plan to proceed immediately to <i>Adenylate 3' Ends</i> on page 15, you can safely stop the protocol here. If you are stopping, seal the ALP plate with a Microseal 'B' adhesive seal and store a -15°C to -25°C for up to 7 days. |
| Co | mments |
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Adenylate 3' Ends

TruSeq Stranded Total RNA Sample Prep HS Protocol

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| Date/Time: | Operator: |
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Adenylate 3' Ends

A single 'A' nucleotide is added to the 3' ends of the blunt fragments to prevent them from ligating to one another during the adapter ligation reaction. A corresponding single '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

| Item | Quantity | Storage | Supplied By |
|--|-------------------------------|----------------|-------------|
| (Optional) A-Tailing Control (CTA) | 1 tube per 48 reactions | -15°C to -25°C | Illumina |
| A-Tailing Mix (ATL) | 1 tube per 48 reactions | -15°C to -25°C | Illumina |
| Resuspension Buffer (RSB) | 1 tube | 2°C to 8°C | Illumina |
| Ice | As needed to place a plate on | -15°C to -25°C | User |
| Microseal 'B' Adhesive Seal | 1 | 15°C to 30°C | User |
| RNase/DNase-free Eight-Tube Strips and Caps (if using multichannel pipettes) | 3 | 15°C to 30°C | User |
| RNase/DNase-free Reagent Reservoirs (if using multichannel pipettes) | 3 | 15°C to 30°C | User |

Add ATL

- [] 1 Do one of the following:
 - If using the in-line control reagent:
 - Centrifuge the thawed A-Tailing Control tube to 600 × g for 5 seconds.
 - Dilute the A-Tailing Control to 1/100 in Resuspension Buffer (For example, 1 μl A-Tailing Control + 99 μl Resuspension Buffer) before use. Discard the diluted A-Tailing Control after use.
 - Add 2.5 μ l of diluted A-Tailing Control to each well of the ALP plate.
 - If not using the in-line control reagent, add $2.5~\mu l$ of Resuspension Buffer to each well of the ALP plate.

| [_] 2 | Ad | d 12.5 μl of thawed A-Tailing Mix to each well of the ALP plate. Mix thoroughly as |
|-------|-----|--|
| | fol | lows: |
| [_ |] a | Seal the ALP plate with a Microseal 'B' adhesive seal. |
| [_ |] b | Shake the ALP plate on a microplate shaker at 1800 rpm for 2 minutes. |
| [_] 3 | Cer | ntrifuge the ALP plate to 280 × g for 1 minute. |



| Date/Time: | Operator: |
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| ncubate 1 / | ALP |
| [_] 1 | Place the sealed ALP plate on the pre-heated microheating system 1. Close the lid and incubate at 37°C for 30 minutes. |
| | Start time: Stop time: |
| [_] 2 | Immediately after the 37°C incubation, remove the ALP plate from system 1 and place the plate on the pre-heated microheating system 2. Close the lid and incubate at 70°C for 5 minutes. |
| | Start time: Stop time: |
| [_] 3 | Set the microheating system 1 to 30°C in preparation for <i>Ligate Adapters</i> . |
| [_] 4 | Immediately remove the ALP plate from the microheating system 2 and place the plate on ice for 1 minute. |
| [_] 5 | Proceed immediately to Ligate Adapters on page 17. |
| Co | omments |
| | |



Ligate Adapters

TruSeq Stranded Total RNA Sample Prep HS Protocol

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| Date/Time: | Operator |
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| Date/Time | Operator |

Ligate Adapters

This process ligates indexing adapters to the ends of the ds cDNA, preparing them for hybridization onto a flow cell.

Consumables

| Item | Quantity | Storage | Supplied By |
|--|---|----------------|-------------|
| (Optional) Ligation Control (CTL) | 1 tube per 48 reactions | -15°C to -25°C | Illumina |
| Choose from the following depending on the kit you are using: TruSeq Stranded Total RNA LT Sample Prep Kit contents: RNA Adapter Indices (AR001–AR016, AR018–AR023, AR025, AR027) TruSeq Stranded Total RNA HT Sample Prep Kit contents: RAP (RNA Adapter Plate) | 1 tube of each index being used, per column of 8 reactions or 1 RAP | -15°C to -25°C | Illumina |
| Ligation Mix (LIG) | 1 tube per 48 reactions | -15°C to -25°C | Illumina |
| Resuspension Buffer (RSB) | 1 tube | 2°C to 8°C | Illumina |
| Stop Ligation Buffer (STL) | 1 tube per 48 reactions | -15°C to -25°C | Illumina |
| Barcode labels for: CAP (Clean Up ALP Plate) PCR (Polymerase Chain Reaction Plate) RAP (RNA Adapter Plate) (if using the HT kit) | 1 label per plate | 15°C to 30°C | Illumina |
| 96-well HSP Plate | 1 | 15°C to 30°C | User |
| 96-well MIDI Plate | 1 | 15°C to 30°C | User |
| AMPure XP Beads | 92 µl per sample | 2°C to 8°C | User |
| Freshly Prepared 80% Ethanol (EtOH) | 800 μl per sample | 15°C to 30°C | User |
| Microseal 'B' Adhesive Seals | 7 | 15°C to 30°C | User |
| RNase/DNase-free Eight-Tube Strips and Caps (if using multichannel pipettes) | 4–28 | 15°C to 30°C | User |
| RNase/DNase-free Reagent Reservoirs (if using multichannel pipettes) | 4–28 | 15°C to 30°C | User |

Experienced User Card and Lab Tracking Form

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Add LIG

- [] 1 Do one of the following:
 - If using RNA Adapter tubes, centrifuge the thawed tubes to 600 × g for 5 seconds.
 - If using a RAP:

| _ | Thaw the plate for 10 m | inutes at room | temperature or | n the benchtop. | Visually |
|---|---------------------------|-------------------|-----------------|-----------------|----------|
| | inspect the wells to make | ke sure that they | y all are thawe | d. | |
| | Start time: | Sto | on time: | | |

- Remove the adapter plate tape seal.
- Centrifuge the plate to 280 × g for 1 minute to collect all of the adapter to the bottom of the well.
- Remove the plastic cover. Save the cover if you are not processing the entire plate at one time.
- If it is the first time using this RAP, apply the RAP barcode label to the plate.



NOTE

- The RAP is single-use for each well.
- Illumina recommends that the RAP does not undergo more than 4 freeze-thaw cycles.
- [_] 2 Centrifuge the Ligation Control (if using Ligation Control) and Stop Ligation Buffer tubes to 600 × g for 5 seconds.
- [] 3 Immediately before use, remove the Ligation Mix tube from -15°C to -25°C storage.
- [_] 4 Remove the adhesive seal from the ALP plate.
- [] 5 Do one of the following:
 - If using the in-line control reagent:
 - Dilute the Ligation Control to 1/100 in Resuspension Buffer (For example, 1 μl Ligation Control + 99 μl Resuspension Buffer) before use. Discard the diluted Ligation Control after use.
 - Add 2.5 μl of diluted Ligation Control to each well of the ALP plate.
 - If not using the in-line control reagent, add $2.5~\mu l$ of Resuspension Buffer to each well of the ALP plate.
- [] 6 Add 2.5 µl of Ligation Mix to each well of the ALP plate.
- [] 7 Return the Ligation Mix tube to -15°C to -25°C storage immediately after use.
- [_] 8 Do one of the following:
 - If using RNA Adapter tubes, add 2.5 μ l of the thawed RNA Adapter Index to each well of the ALP plate.
 - If using a RAP:
 - Place the RAP on the benchtop so that the part number barcode, on the long side of the plate, is facing you and the clipped corner is on the lower left.



Ligate Adapter

TruSeq Stranded Total RNA Sample Prep HS Protocol

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| Date/Time: | Operator: |
| | Do one of the following to pierce the foil seal: If using the entire plate at one time, use the bottom of a clean 96-well semi-skirted PCR plate to pierce a hole in all of the well seals simultaneously. Gently but firmly, press the clean plate over the foil seal. If using only part of the plate, use the bottom of a clean eight-tube strip, with caps attached, to pierce holes in the seals of the wells that will be used for ligation. Repeat with a new, clean eight-tube strip, with caps attached, for each row or column of adapters that will be used for ligation. Using an eight-tip multichannel pipette, transfer 2.5 µl of the appropriate thawed RNA Adapter from the RAP well to each well of the ALP plate. |
| [_] 9 [_] [_] | |
| [_] 10 | Centrifuge the ALP plate to 280 × g for 1 minute. |
| Incubate 2 A | ALP |
| [_] 1 | Place the sealed ALP plate on the pre-heated microheating system. Close the lid and incubate at 30°C for 10 minutes. Start time: Stop time: |
| [_] 2 | Remove the ALP plate from the microheating system. |
| Add STL | |
| [_] 1 | Remove the adhesive seal from the ALP plate. |
| [_] 2 [_] [_] | • |
| [_] 3 | Centrifuge the ALP plate to 280 × g for 1 minute. |
| Clean Up AL | _P |
| [_] 1 | Remove the adhesive seal from the ALP plate. |
| [_] 2 | Vortex the AMPure XP Beads for at least 1 minute or until they are well dispersed. |
| [_] 3 | Add 42 μl of mixed AMPure XP Beads to each well of the ALP plate. Mix thoroughly as follows: |
| [_] [_] | a Seal the ALP plate with a Microseal 'B' adhesive seal.b Shake the ALP plate on a microplate shaker at 1800 rpm for 2 minutes. |
| [_] 4 | Incubate the ALP plate at room temperature for 15 minutes. |

Centrifuge the ALP plate to 280 × g for 1 minute.

Remove the adhesive seal from the ALP plate.

Stop time: _

[_] 5 [_] 6 Start time: __

| ate/11m | e: Operator: |
|---------|--|
| [_] 7 | Place the ALP plate on the magnetic stand at room temperature for 5 minutes or until the liquid is clear. Start time: Stop time: |
| [_] 8 | Remove and discard 79.5 µl supernatant from each well of the ALP plate. |
| [_] 9 | With the ALP plate on the magnetic stand, add 200 μ l freshly prepared 80% EtOH to each well without disturbing the beads. |
| [_] 1 | Incubate the ALP plate at room temperature for 30 seconds, and then remove and discard al of the supernatant from each well. |
| [_] 1 | Repeat steps 9 and 10 one time for a total of two 80% EtOH washes. |
| [_] 1 | With the ALP plate on the magnetic stand, let the samples air-dry at room temperature for 15 minutes. |
| | Start time: Stop time: |
| | Remove the ALP plate from the magnetic stand. |
| | 4 Add 52.5 μl Resuspension Buffer to each well of the ALP plate. Mix thoroughly as follows: [] a Seal the ALP plate with a Microseal 'B' adhesive seal. [] b Shake the ALP plate on a microplate shaker at 1800 rpm for 2 minutes. |
| [_] 1 | 5 Incubate the ALP plate at room temperature for 2 minutes. Start time: Stop time: |
| [_] 1 | 6 Centrifuge the ALP plate to 280 × g for 1 minute. |
| [_] 1 | 7 Remove the adhesive seal from the ALP plate. |
| [_] 1 | Place the ALP plate on the magnetic stand at room temperature for 5 minutes or until the liquid is clear. Start time: Stop time: |
| [_] 1 | 9 Transfer 50 μ l supernatant from each well of the ALP plate to the corresponding well of the new MIDI plate labeled with the CAP barcode. |
| [_] 2 | 0 Vortex the AMPure XP Beads until they are well dispersed. |
| | Add 50 μl of mixed AMPure XP Beads to each well of the CAP plate for a second cleanup. Mix thoroughly as follows: [] a Seal the CAP plate with a Microseal 'B' adhesive seal. [] b Shake the CAP plate on a microplate shaker at 1800 rpm for 2 minutes. |
| [_] 2 | 2 Incubate the CAP plate at room temperature for 15 minutes. |
| | Start time: Stop time: |
| [_] 2 | 3 Centrifuge the CAP plate to 280 × g for 1 minute. |
| [_] 2 | 4 Remove the adhesive seal from the CAP plate. |
| [_] 2 | liquid is clear. |
| | Start time: Stop time: |
| - 1-12 | 6 Remove and discard 95 ul supernatant from each well of the CAP plate. |



| Date/Time: | Operator: |
|----------------------|---|
| [_] 27 | With the CAP plate on the magnetic stand, add 200 μl freshly prepared 80% EtOH to each well. |
| [_] 28 | Incubate the CAP plate at room temperature for 30 seconds, and then remove and discard all of the supernatant from each well. |
| [_] 29 | Repeat steps 27 and 28 one time for a total of two 80% EtOH washes. |
| [_] 30 | With the CAP plate on the magnetic stand, let the samples air-dry at room temperature for 15 minutes. |
| | Start time: Stop time: |
| [_] 31 | Remove the CAP plate from the magnetic stand. |
| [_] 32 [_] [_] | |
| [_] 33 | Incubate the CAP plate at room temperature for 2 minutes. Start time: Stop time: |
| [_] 34 | Centrifuge the CAP plate to 280 × g for 1 minute. |
| [_] 35 | Remove the adhesive seal from the CAP plate. |
| [_] 36 | Place the CAP plate on the magnetic stand at room temperature for 5 minutes or until the liquid is clear. |
| | Start time: Stop time: |
| [_] 37 | Transfer 20 μl supernatant from each well of the CAP plate to the corresponding well of the new HSP plate labeled with the PCR barcode. |
| 7 | SAFE STOPPING POINT If you do not plan to proceed immediately to <i>Enrich DNA Fragments</i> on page 23, you can safely stop the protocol here. If you are stopping, seal the PCR plate with a Microseal 'B' adhesive seal and store at -15°C to -25°C for up to seven days. |
| Co | omments |



Enrich DNA Fragments

TruSeq Stranded Total RNA Sample Prep HS Protocol

Experienced User Card and Lab Tracking Form

| Date/Time: | Operator: | |
|------------|-----------|--|

Enrich DNA Fragments

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

Consumables

| Item | Quantity | Storage | Supplied By |
|---|-------------------------|----------------|-------------|
| PCR Master Mix (PMM) | 1 tube per 48 reactions | -15°C to -25°C | Illumina |
| PCR Primer Cocktail (PPC) | 1 tube per 48 reactions | -15°C to -25°C | Illumina |
| Resuspension Buffer (RSB) | 1 tube | 2°C to 8°C | Illumina |
| Barcode labels for: CPP (Clean Up PCR Plate) barcode label TSP1 (Target Sample Plate) barcode label | 1 label per plate | 15°C to 30°C | Illumina |
| 96-well HSP Plate | 1 | 15°C to 30°C | User |
| 96-well MIDI Plate | 1 | 15°C to 30°C | User |
| AMPure XP Beads | 50 µl per sample | 2°C to 8°C | User |
| Freshly Prepared 80% Ethanol (EtOH) | 400 μl per sample | 15°C to 30°C | User |
| Microseal 'A' Film | 1 | 15°C to 30°C | User |
| Microseal 'B' Adhesive Seals | 3 | 15°C to 30°C | User |
| RNase/DNase-free Eight-Tube Strips and Caps (if using multichannel pipettes) | 5 | 15°C to 30°C | User |
| RNase/DNase-free Reagent Reservoirs (if using multichannel pipettes) | 5 | 15°C to 30°C | User |

Make PCR

| | 1 | Add 5 | μl c | of thawed | PCR | Primer | Cocktail | to | each | well | of | the | PCR | pla | ate. |
|--|---|-------|------|-----------|-----|--------|----------|----|------|------|----|-----|-----|-----|------|
|--|---|-------|------|-----------|-----|--------|----------|----|------|------|----|-----|-----|-----|------|

- [] 2 Add 25 µl of thawed PCR Master Mix to each well of the PCR plate.
 - [_] a Seal the PCR plate with a Microseal 'A' film.
 - [_] b Shake the PCR plate on a microplate shaker at 1600 rpm for 20 seconds.
- [_] 3 Centrifuge the PCR plate to 280 × g for 1 minute.



| • | |
|----------------------------|--|
| Amp PCR | |
| [_] 1 [_] [_] [_] | Place the sealed PCR plate on the pre-programmed thermal cycler. Close the lid, then select and run PCR to amplify the plate. a |
| Clean Up Po | CR |
| [_] 1 | Remove the adhesive seal from the PCR plate. |
| [_] 2 | Vortex the AMPure XP Beads until they are well dispersed. |
| [_] 3 | Do one of the following, depending on the adapter type used: If using the RNA Adapter tubes, add 50 µl of the mixed AMPure XP Beads to each well of the new MIDI plate labeled with the CPP barcode. If using the RAP, add 47.5 µl of the mixed AMPure XP Beads to each well of the new MIDI plate labeled with the CPP barcode. |
| | Transfer the entire contents from each well of the PCR plate to the corresponding well of the CPP plate containing 50 µl of mixed AMPure XP Beads. Mix thoroughly as follows: a Seal the CPP plate with a Microseal 'B' adhesive seal. b Shake the CPP plate on a microplate shaker at 1800 rpm for 2 minutes. |
| [_] 5 | Incubate the CPP plate at room temperature for 15 minutes. Start time: Stop time: |
| [_] 6 | Centrifuge the CPP plate to 280 × g for 1 minute. |
| [_] 7 | Remove the adhesive seal from the CPP plate. |
| [_] 8 | Place the CPP plate on the magnetic stand at room temperature for 5 minutes or until the liquid is clear. Start time: Stop time: |
| [_] 9 | Remove and discard 95 µl supernatant from each well of the CPP plate. |
| [_] 10 | With the CPP plate on the magnetic stand, add 200 μl freshly prepared 80% EtOH to each well without disturbing the beads. |
| [_] 11 | Incubate the CPP plate at room temperature for 30 seconds, and then remove and discard all of the supernatant from each well. |
| [_] 12 | Repeat steps 10 and 11 one time for a total of two 80% EtOH washes. |
| [_] 13 | With the CPP plate on the magnetic stand, let the samples air-dry at room temperature for 15 minutes, and then remove the plate from the magnetic stand. Start time: Stop time: |



| Date/Time: _ | Operator: |
|-----------------------------------|--|
| [_] 14 <i>A</i> [_] a [_] b | 1 |
| | Incubate the CPP plate at room temperature for 2 minutes. Start time: Stop time: |
| [_] 16 C | Centrifuge the CPP plate to 280 × g for 1 minute. |
| [_] 17 F | Remove the adhesive seal from the CPP plate. |
| li | Place the CPP plate on the magnetic stand at room temperature for 5 minutes or until the iquid is clear. Start time: Stop time: |
| | Fransfer 30 μ l supernatant from each well of the CPP plate to the corresponding well of the new HSP plate labeled with the TSP1 barcode. |
| V | SAFE STOPPING POINT If you do not plan to proceed immediately to <i>Validate Library</i> on page 27, you can safely stop the protocol here. If you are stopping, seal the TSP1 plate with a Microseal 'B' adhesive seal and store at -15°C to -25°C for up to 7 days. |
| Con | nments |



Validate Library

TruSeq Stranded Total RNA Sample Prep HS Protocol

| • | nced User Card and Lab Tracking Form Operator: |
|-----------|--|
| Date/1 | mre Operator |
| Validat | te Library |
| | Illumina recommends performing the following procedures for quality control analysis on your sample library and quantification of the DNA library templates. |
| Quantify | Libraries |
| | To achieve the highest quality data on Illumina sequencing platforms, it is important to create optimum cluster densities across every lane of the flow cell. Optimizing cluster densities requires accurate quantitation of DNA library templates. Quantify your libraries using qPCR according to the Illumina Sequencing Library qPCR Quantification Guide (part # 11322363). |
| Quality C | Control |
| [_] | Load 1 μ l of the resuspended construct on an Agilent Technologies 2100 Bioanalyzer using a DNA-specific chip such as the Agilent DNA 1000. |
| [_] | 2 Check the size and purity of the sample. The final product should be a band at approximately 260 bp. |
| | |

Comments

Validate Library

TruSeq Stranded Total RNA Sample Prep HS Protocol



| Experienced | l User | Card | and | Lab | Tracking | Form |
|-------------|--------|------|-----|-----|----------|------|
|-------------|--------|------|-----|-----|----------|------|

| Date/Time: | Operator: |
|-------------|-----------|
| Date/ Hille | Operator |

Normalize and Pool Libraries

This process describes how to prepare DNA templates for cluster generation. Indexed DNA libraries are normalized to 10 nM in the DCT plate and then pooled in equal volumes in the PDP plate. DNA libraries not intended for pooling are normalized to 10 nM in the DCT plate.

Consumables

| Item | Quantity | Storage | Supplied By |
|--|--|--------------|-------------|
| Barcode labels for: • DCT (Diluted Cluster Template) • PDP (Pooled DCT Plate) (for pooling only) | 1 label per plate | 15°C to 30°C | Illumina |
| 96-well HSP Plate (for pooling only) | 1 | 15°C to 30°C | User |
| 96-well MIDI Plate | 1 | 15°C to 30°C | User |
| Microseal 'B' Adhesive Seals | 5 | 15°C to 30°C | User |
| Tris-HCl 10 mM, pH8.5 with 0.1% Tween 20 | Enough to normalize the concentration of each sample library to 10 nM | 15°C to 30°C | User |

Make DCT

- [$_$] 1 Transfer 10 μ l of sample library from each well of the TSP1 plate to the corresponding well of the new MIDI plate labeled with the DCT barcode.
- [_] 2 Normalize the concentration of sample library in each well of the DCT plate to 10 nM using Tris-HCl 10 mM, pH 8.5 with 0.1% Tween 20.
- [_] 3 Mix the DCT plate as follows:
 - $[\]$ a Seal the DCT plate with a Microseal 'B' adhesive seal.
 - [] b Shake the DCT plate on a microplate shaker at 1000 rpm for 2 minutes.
- [_] 4 Centrifuge the DCT plate to 280 × g for 1 minute.
- [_] 5 Remove the adhesive seal from the DCT plate.
- [_] 6 Depending on the type of library you want to generate, do one of the following:
 - For non-pooled libraries, the protocol stops here. Do one of the following:
 - Proceed to cluster generation.
 - Seal the DCT plate with a Microseal 'B' adhesive seal and store at -15°C to -25°C.
 - For pooled libraries, proceed to Make PDP (for pooling only).

Make PDP (for pooling only)

Determine the number of samples to be combined together for each pool.



| ate/Time: | Operator: |
|---------------------|---|
| [_] 2 | Do one of the following: If pooling 2–24 samples: — Transfer 10 μl of each normalized sample library to be pooled from the DCT plate to one well of the new HSP plate labeled with the PDP barcode. — The total volume in each well of the PDP plate should be 10X the number of combined sample libraries and 20–240 μl (2–24 libraries). If pooling 25–96 samples: — Using a multichannel pipette, transfer 5 μl of each normalized sample library in column 1 from the DCT plate to column 1 of the new HSP plate labeled with the PDP barcode. — Transfer 5 μl of each normalized sample library in column 2 from the DCT plate to column 1 of the PDP plate. — Repeat the transfer for as many times as there are remaining columns in the DCT plate. The result is a PDP plate with pooled samples in column 1. Mix the PDP plate as follows: — Seal the PDP plate with a Microseal 'B' adhesive seal. — Shake the PDP plate on a microplate shaker at 1800 rpm for 2 minutes. — Centrifuge the PDP plate to 280 × g for 1 minute. — Remove the adhesive seal from the PDP plate. — Combine the contents of each well of column 1 into well A2 of the PDP plate for the |
| [_] 3 [_] [_] | |
| [_] 4 | Centrifuge the PDP plate to $280 \times g$ for 1 minute. |
| [_] 5 | Do one of the following: Proceed to cluster generation. Store the sealed PDP plate at -15°C to -25°C. |
| Cc | omments |
| | |



Technical Assistance

For technical assistance, contact Illumina Technical Support.

Table 1 Illumina General Contact Information

| Illumina Website | www.illumina.com |
|------------------|--------------------------|
| Email | techsupport@illumina.com |

Table 2 Illumina Customer Support Telephone Numbers

| Region | Contact Number | Region | Contact Number |
|---------------|----------------|-----------------|-----------------|
| North America | 1.800.809.4566 | Italy | 800.874909 |
| Austria | 0800.296575 | Netherlands | 0800.0223859 |
| Belgium | 0800.81102 | Norway | 800.16836 |
| Denmark | 80882346 | Spain | 900.812168 |
| Finland | 0800.918363 | Sweden | 020790181 |
| France | 0800.911850 | Switzerland | 0800.563118 |
| Germany | 0800.180.8994 | United Kingdom | 0800.917.0041 |
| Ireland | 1.800.812949 | Other countries | +44.1799.534000 |

MSDSs

Material safety data sheets (MSDSs) are available on the Illumina website at www.illumina.com/msds.

Product Documentation

Product documentation in PDF is available for download from the Illumina website. Go to www.illumina.com/support, select a product, then click **Documentation & Literature**.





Illumina San Diego, California 92122 U.S.A. +1.800.809.ILMN (4566) +1.858.202.4566 (outside North America) techsupport@illumina.com

www.illumina.com