

Addendum

GENECHIP® WHOLE TRANSCRIPT (WT) SENSE TARGET LABELING ASSAY MANUAL ADDENDUM

This addendum supplements the *GeneChip® Whole Transcript (WT) Sense Target Labeling Assay Manual* version 4 (P/N 701880 Rev 4). Two changes need to be made to the manual and are outlined in this addendum. For your convenience, in addition to the description listed below, new tables are provided that can be separated and included in the manual where the changes specifically occur.

! **IMPORTANT: It is critical that users note these changes to the protocol.**

Alteration One:

Chapter 2, Table 2.3 on page 15 (RiboMinus Reaction and Wash Volumes Table)

In the last row of Table 2.3 under *Concentration and Clean-up*, the Ethanol should be **80% Ethanol** and not 100% Ethanol, as listed in the manual. Please note that the first occurrence of Ethanol listed in the table under *Concentration and Clean-up* remains 100% Ethanol, and only the second wash uses 80% Ethanol.

Alteration Two:

Chapter 4, Table 4.1 on page 33 (First Cycle, Primer/Poly-A RNA Controls For 100ng of Total RNA)

The content in the originally published table is correct. However, researchers using more than 100 ng of starting material need to adjust the volumes of Diluted Poly-A RNA Controls (3rd dilution, 1:50) and RNase-free Water appropriately. This addendum contains a modified version of Table 4.1 which adds an additional column to display the appropriate volumes for 300 ng of total RNA.

Volumes for 300 ng starting material:

T7-(N) ₆ Primers, 2.5 µg/µL	2 µL (no change)
Diluted Poly-A RNA Controls (3rd dilution, 1:50)	6 µL (change)
RNase-free Water	12 µL (change)
Total Volume	20 µL (no change)

! **TIP: For your convenience, you may either manually edit your hard-copy version of the manual, or cut and paste the tables provided on page 2 of this addendum to the appropriate pages.**

Chapter 2, Table 2.3 on page 15

Table 2.3 RiboMinus Reaction and Wash Volumes


Total RNA Amount	1 µg	1 µg	2 µg
RNA Concentration Range	1 µg/µL	0.31 µg/µL to 1 µg/µL	≥ 0.62 µg/µL
RiboMinus Probe Hybridization			
Total RNA/Poly-A Controls Mix (from Procedure A)	3.0 µL	up to 5.2 µL	up to 5.2 µL
RiboMinus Probe (100 pmol/µL)	0.8 µL	0.8 µL	1.6 µL
Hybridization Buffer with Betaine (from Procedure B)	20 µL	30 µL	40 µL
Total Hybridization Volume	23.8 µL	36.0 µL	46.8 µL
Beads Preparation			
Magnetic Beads	50 µL	50 µL	100 µL
2X Wash with Water	50 µL	50 µL	100 µL
1X Wash with Hybridization Buffer with Betaine	50 µL	50 µL	100 µL
Re-suspension of Beads with Hybridization Buffer with Betaine	30 µL	20 µL	60 µL
rRNA Reduction			
Final Wash of Beads with Hybridization Buffer with Betaine	50 µL	50 µL	50 µL
Concentration and Clean-up			
Approximate Supernatant Volume	~100 µL	~100 µL	~150 µL
cRNA Binding Buffer	350 µL	350 µL	525 µL
100% Ethanol	250 µL	250 µL	375 µL
Total Volume Applied to Column	~700 µL	~700 µL	~1,050 µL*
cRNA Wash Buffer	500 µL	500 µL	500 µL
80% Ethanol	500 µL	500 µL	500 µL

* Multiple applications to column required (column capacity ≤ 800 µL).

Chapter 4, Table 4.1 on page 33

Table 4.1 First-cycle, Primer/Poly-A RNA Controls

Component	Total RNA Volume	
	100 ng	300 ng
T7-(N) ₆ Primers, 2.5 µg/µL	2 µL	2 µL
Diluted Poly-A RNA Controls (3rd dilution, 1:50)	2 µL	6 µL
RNase-free Water	16 µL	12 µL
Total Volume	20 µL	20 µL



**GeneChip® Whole Transcript (WT) Sense Target
Labeling Assay Manual**
Version 4

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Use of the GeneChip® WT cDNA Synthesis and Amplification Kit in accordance with the instructions provided is accompanied by a limited license to U.S. Patent Nos. 5,716,785; 5,891,636; 6,291,170; and 5,545,522. Users who do not purchase this Kit may be required to obtain a license under these patents or to purchase another licensed kit. Reagent products may also be covered by one or more of the following patents: U.S. Patent Nos. 6,864,059.

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Introduction

The Affymetrix GeneChip® Whole Transcript (WT) Sense Target Labeling Assay is designed to generate amplified and biotinylated sense-strand DNA targets from the entire expressed genome without bias. This assay and associated reagents have been optimized specifically for use with the GeneChip® ST Arrays where “ST” stands for “Sense Target,” and the probes on the arrays have been selected to be distributed throughout the entire length of each transcript.



NOTE: The WT Assay is not compatible with GeneChip arrays designed to focus on the 3' ends of the transcripts. For the 3' arrays, continue to follow the protocols detailed in the *GeneChip® Expression Analysis Technical Manual (P/N 900223)*.

This manual describes in detail two configurations of the WT Assay with key differences indicated in [Table 1.1](#). The 1 µg Total RNA Labeling Protocol starts with a ribosomal RNA (rRNA) reduction procedure where the 28S and 18S rRNA population is significantly reduced from the total RNA sample minimizing the background and thereby increasing the array detection sensitivity and specificity. The rRNA reduction becomes critical when a user is interested in high-sensitivity analysis of expression levels for both genes and exons using the GeneChip® Exon 1.0 ST Arrays. This is because exon probe sets contain a smaller number of probes and in some cases, selection of those probes is constrained by the limited size of the probe selection region. Therefore it is imperative to use the high-sensitivity assay for optimal performance.

The protocol has been optimized for the 1 µg input amount. However, a modest increase in cRNA yield has been observed by increasing the amount of total RNA used. Input amounts up to 2 µg show no adverse impact on array performance. However, at 2 µg, it is highly recommended to scale-up the RiboMinus reagents to insure efficient rRNA reduction. Failure to do so may have a small negative impact on sensitivity, particularly for analyses at the exon level.

On the other hand, analysis of the gene level benefits from a larger number of high-quality probes selected from the entire transcript, thus the advantage of the additional

rRNA reduction step is reduced. Therefore, the 100 ng Total RNA Labeling Protocol is acceptable for use with the GeneChip® Gene 1.0 ST Arrays.

When omitting the RiboMinus procedure, using 100 ng of total RNA as input has been shown to generate sufficient cRNA from a diverse set of RNA sources. However, for some RNAs that yield less cRNA, increasing the amount of total RNA used can result in a modest increase in cRNA made. Target generated from a range of 100 to 300 ng of input total RNA has demonstrated equivalent array performance.

For more information regarding the performances of the two protocols on Gene 1.0 ST Arrays refer to the *Whole Transcript Sense Target Labeling Assay Performance* white paper.

As outlined in [Figure 1.1](#), following the rRNA reduction procedure, the two protocols merge where double-stranded cDNA is synthesized with random hexamers tagged with a T7 promoter sequence. The double-stranded cDNA is subsequently used as a template and amplified by T7 RNA polymerase producing many copies of antisense cRNA. In the second cycle of cDNA synthesis, random hexamers are used to prime reverse transcription of the cRNA from the first cycle to produce single-stranded DNA in the sense orientation.

In order to reproducibly fragment the single-stranded DNA and improve the robustness of the assay, a novel approach is utilized where dUTP is incorporated in the DNA during the second-cycle, first-strand reverse transcription reaction. This single-stranded DNA sample is then treated with a combination of uracil DNA glycosylase (UDG) and apurinic/apyrimidinic endonuclease 1 (APE 1) that specifically recognizes the unnatural dUTP residues and breaks the DNA strand. DNA is labeled by terminal deoxynucleotidyl transferase (TdT) with the Affymetrix® proprietary DNA Labeling Reagent that is covalently linked to biotin.

Following the recommended procedures, sufficient target is anticipated to be generated for hybridization to a single array.

Follow the instructions closely for the most optimal results. As an Affymetrix GeneChip microarray user, your feedback is welcome. Please contact your technical support representative with any input on how we can improve this resource.

Table 1.1 Two Configurations of the WT Sense Target Labeling Assay

	1 µg Total RNA Labeling Protocol	100 ng Total RNA Labeling Protocol
Recommended Amount of Starting Material	1 µg Total RNA	100 ng Total RNA
Acceptable Range of Input Amount	1 – 2 µg	100 – 300 ng
Procedural Difference	Requires rRNA Reduction with RiboMinus Kit	Omits rRNA Reduction
Exon ST Arrays	Recommended	Not Optimal
Gene ST Arrays	Acceptable	Recommended

Whole Transcript Sense Target Labeling Assay Schematic

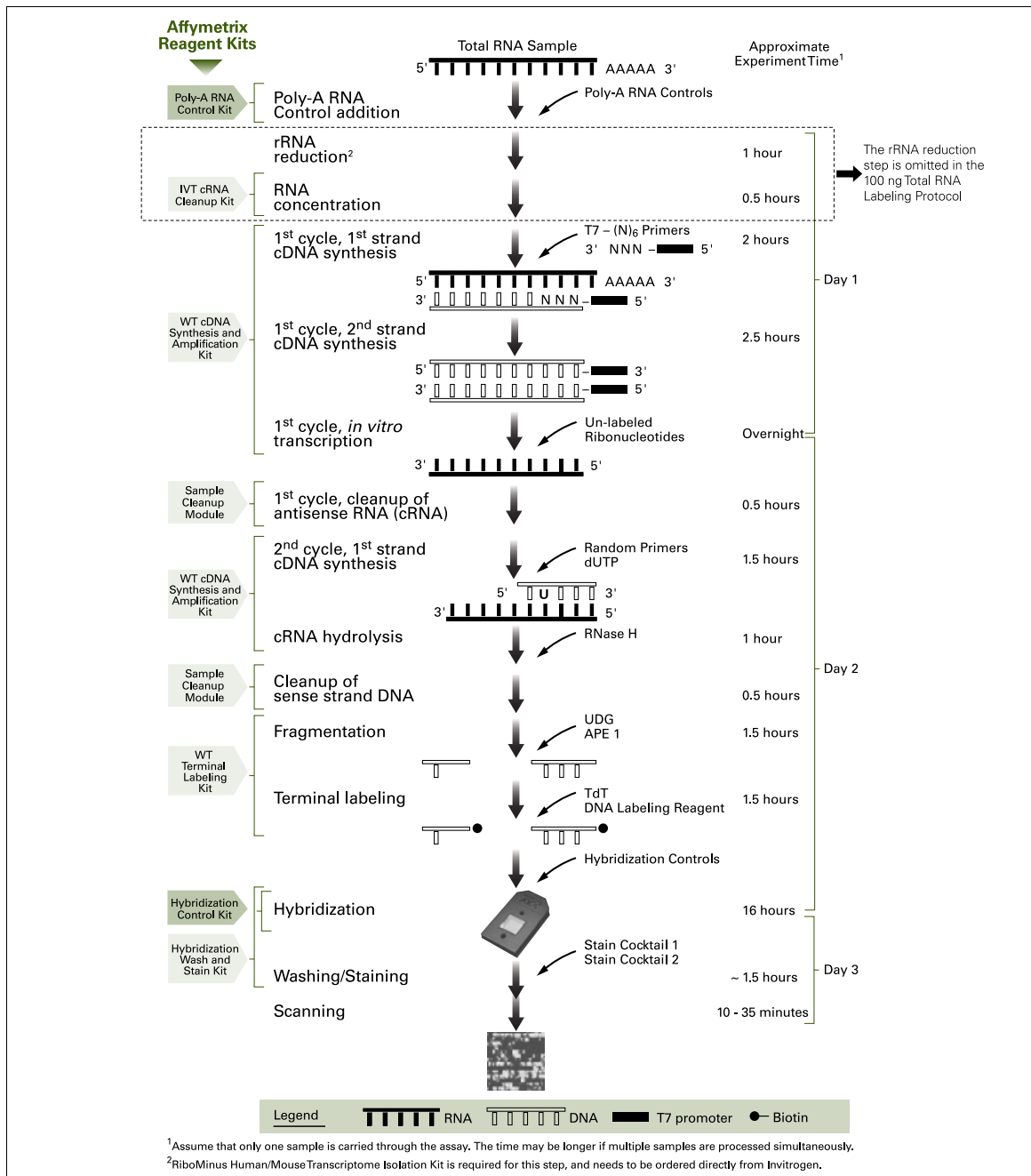


Figure 1.1 GeneChip® Whole Transcript Sense Target Labeling Assay

Materials

Necessary Reagents

Table 1.2 Necessary Reagents

Material	Source	P/N
rRNA Reduction (Not required for the 100 ng Total RNA Labeling Protocol)		
RiboMinus™ Transcriptome Isolation Kit (Human/Mouse)	Invitrogen	K1550-02 (49 Rxn)*
Magna-Sep™ Magnetic Particle Separator	Invitrogen	K1585-01
Betaine, 5M	Sigma-Aldrich	B-0300
Target Labeling		
GeneChip® WT Sense Target Labeling and Control Reagents Contains one of each of the following kits that can also be ordered individually:	Affymetrix	900652 (30 Rxn)
<ul style="list-style-type: none"> • GeneChip® Eukaryotic Poly-A RNA Control Kit (~100 Rxn) • GeneChip® WT cDNA Synthesis and Amplification Kit (30 Rxn) • GeneChip® WT Terminal Labeling Kit (30 Rxn) • GeneChip® Sample Cleanup Module (30 Rxn) • GeneChip® IVT cRNA Cleanup Kit (30 Rxn) • GeneChip® Hybridization Control Kit (30 Rxn) 		
cDNA Synthesis and Amplification		
GeneChip® Eukaryotic Poly-A RNA Control Kit Contains:	Affymetrix	900433 (~100 Rxn)
<ul style="list-style-type: none"> • Poly-A Control Stock • Poly-A Control Dil Buffer 		

* 49 reactions when using 1 µg of total RNA. The number of reactions per kit will be ~24 if using 2 µg of total RNA.

Table 1.2 Necessary Reagents (Continued)

Material	Source	P/N
GeneChip® WT cDNA Synthesis and Amplification Kit	Affymetrix	900673 (30 Rxn) or 900672 (10 Rxn)
Sub-kit 1: GeneChip® WT cDNA Synthesis Kit		
Contains:		
<ul style="list-style-type: none"> • T7-(N)₆ Primers, 2.5 µg/µL • 5X 1st Strand Buffer • DTT, 0.1M • dNTP, 10 mM • RNase Inhibitor • SuperScript™ II • MgCl₂, 1M • DNA Polymerase I • RNase H • Random Primers, 3 µg/µL • dNTP+dUTP, 10 mM • RNase-free Water 		
Sub-kit 2: GeneChip® WT cDNA Amplification Kit		
Contains:		
<ul style="list-style-type: none"> • 10X IVT Buffer • IVT NTP Mix • IVT Enzyme Mix • IVT Control 		
Fragmentation and Labeling		
GeneChip® WT Terminal Labeling Kit	Affymetrix	900671 (30 Rxn) or 900670 (10 Rxn)
Contains:		
<ul style="list-style-type: none"> • 10X cDNA Fragmentation Buffer • UDG, 10 U/µL • APE 1, 1,000 U/µL • 5X TdT Buffer • TdT, 30 U/µL • DNA Labeling Reagent, 5 mM • RNase-free Water 		
cDNA, cRNA Cleanup		
GeneChip® IVT cRNA Cleanup Kit	Affymetrix	900547 (30 Rxn)
Contains:		
<ul style="list-style-type: none"> • IVT cRNA Cleanup Spin Columns • IVT cRNA Binding Buffer • IVT cRNA Wash Buffer, 5 mL concentrate • RNase-free Water • 1.5 mL Collection Tubes (for elution) • 2 mL Collection Tubes 		

Table 1.2 Necessary Reagents (Continued)

Material	Source	P/N
GeneChip® Sample Cleanup Module Contains: <ul style="list-style-type: none"> • cDNA Cleanup Spin Columns • cDNA Binding Buffer • cDNA Wash Buffer, 6mL concentrate • cDNA Elution Buffer • IVT cRNA Cleanup Spin Columns • IVT cRNA Binding Buffer • IVT cRNA Wash Buffer, 5 mL concentrate • RNase-free Water • 1.5 mL Collection Tubes (for elution) • 2 mL Collection Tubes • 5X Fragmentation Buffer 	Affymetrix	900371 (30 Rxn)
Hybridization, Stain and Wash		
GeneChip® Hybridization Control Kit Contains: <ul style="list-style-type: none"> • 20X Hybridization Controls • 3 nM Control Oligo B2 	Affymetrix	900454 (30 Rxn) or 900457 (150 Rxn)
GeneChip® Hybridization, Wash, and Stain Kit (30 reactions) containing: Hybridization Module from Box 1 <ul style="list-style-type: none"> • Pre-Hybridization Mix • 2X Hybridization Mix • DMSO • Nuclease-free water Stain Module from Box 1 <ul style="list-style-type: none"> • Stain Cocktail 1 • Stain Cocktail 2 • Array Holding Buffer Wash Buffers A and B from Box 2 <ul style="list-style-type: none"> • Wash Buffer A (P/N 900721) • Wash Buffer B (P/N 900722) 	Affymetrix	900720 (30 Rxn)

Miscellaneous Reagents

Table 1.3 Miscellaneous Reagents

Materials	Source	P/N
Miscellaneous Reagents		
Absolute ethanol	Gold Shield Chemical Co.	N/A
RNA 6000 Nano Kit	Agilent	5067-1511
Gel-Shift Assay (Optional)		
Novex XCell SureLock Mini-Cell*	Invitrogen	EI0001
TBE Gel, 4-20%,1.0 mm, 12 well*	Invitrogen	EC62252
Novex Hi-Density TBE Sample Buffer (5X)	Invitrogen	LC6678
10X TBE Buffer	Cambrex	50843
SYBR Gold	Invitrogen	S-11494
10 bp DNA ladder and 100 bp DNA ladder	Invitrogen	10821-015 15628-019
ImmunoPure NeutrAvidin	Pierce	31000
PBS, pH 7.2	Invitrogen	20012-027

*Or equivalent.

Miscellaneous Supplies

Table 1.4 Miscellaneous Supplies*

Materials	Source	P/N
Miscellaneous Supplies		
1.5 mL RNase-free Microfuge Tubes*	Ambion	12400
1.5 mL Non-stick RNase-free Microfuge Tubes*	Ambion	12450
0.2 mL MicroAmp reaction tubes (8 tubes/strip)*	Applied Biosystems	N801-0580
MicroAmp caps for 8 strip tubes*	Applied Biosystems	N801-0535
Pipette for 25 mL*	VWR	53283-710
Pipet-aid*	VWR	53498-103
Tough-Spots®	USA Scientific	9185

* Or equivalent.

Instruments

Table 1.5 Instruments

Instruments	Manufacturer	P/N
NanoDrop ND-1000*	NanoDrop Technologies	N/A
GeneChip® Hybridization Oven 640	Affymetrix	800138 (110 v) 800139 (220 v)
Eppendorf Centrifuge*	Eppendorf	5417C
Tube-Strip PicoFuge*	Stratagene	400540
PicoFuge*	Stratagene	400550
GeneChip® Fluidics Station 450	Affymetrix	00-0079
GeneChip® Scanner 3000 7G	Affymetrix	00-0212 (North America) 00-0213 (International)
GeneChip® AutoLoader with External Barcode Reader (Optional)	Affymetrix	00-0090 (GCS 3000 7G S/N 501) 00-0129 (GCS 3000 7G S/N 502)
ABI GeneAmp PCR System 9700*	Applied Biosystems	N8050001
Bioanalyzer 2100	Agilent	G2940CA
Heating blocks*	VWR	13259-030
Pipette for 0.1 to 2 µL*	Rainin	L-2
Pipette for 2 to 20 µL*	Rainin	L-20
Pipette for 20 to 200 µL*	Rainin	L-200
Pipette for 100 to 1000 µL*	Rainin	L-1000

*Or equivalent.

Suggested Workflow¹

Day 1

- Complete Chapter 2: rRNA Reduction – ~1.5 hours (for GeneChip Exon 1.0 ST Arrays)
- Complete Chapter 3 or 4, Procedures A – C: First Cycle: Synthesis of cDNA – ~3.5 hours
- Start Chapter 3, Procedure D: First Cycle: Synthesis of cRNA – 16 hours – Start on Day 1, finish on Day 2

Day 2

- Complete Chapter 3, Procedures D – I: Second Cycle: Synthesis of first-strand DNA, fragmentation, and labeling – 8 hours
- Start Chapter 5: Hybridization – 17 hours – Start on Day 2, finish on Day 3

Day 3

- Chapter 6: Array Washing, Staining, and Scanning – 2 hours

¹ Assumes that only one sample is carried through the assay. The estimated time required may be longer if multiple samples are processed simultaneously.

rRNA REDUCTION AND PREPARATION OF TOTAL RNA WITH DILUTED POLY-A RNA CONTROLS



NOTE: The rRNA reduction step may be omitted when following the 100 ng Total RNA Labeling Protocol to prepare targets for GeneChip Gene 1.0 ST Arrays. Proceed directly to [Chapter 4](#) and follow the 100 ng Total Labeling Protocol.

Procedure A: Preparation of Dilutions of Poly-A RNA Controls

This Procedure requires the use of the GeneChip® Poly-A RNA Control Kit.

The quality of the starting RNA sample is essential to the overall success of the analysis. Since the most appropriate protocol for the isolation of RNA can be source dependent, we recommend using a protocol that has been established for the tissues or cells being used. In the absence of an established protocol, using one of the commercially available kits designed for RNA isolation is suggested.



IMPORTANT: When using a commercial kit, follow the manufacturer's instructions for RNA isolation.

The following protocol requires a minimum of 1 μg of total RNA as starting material, and the concentration should not fall below 0.31 $\mu\text{g}/\mu\text{L}$. In other words, the 1 μg of total RNA should be suspended in a maximum of 3.2 μL of solution in volume. One to 2 μg of total RNA may be used, but the RNA concentration must be high enough because the maximum volume of total RNA is 3.2 μL .

1. The Poly-A RNA controls are provided as a concentrated stock of 4 different transcripts at staggered concentrations. Dilution buffer is supplied with the kit to prepare the appropriate dilutions based on [Table 2.1](#). Use non-stick RNase-free microfuge tubes for all dilutions. Keep the tubes on ice at all times.

Table 2.1 Poly-A RNA Control Stock Serial Dilutions

Starting Amount of Total RNA	Serial Dilutions			Volume into Sample
	First	Second	Third	
1 µg	1:20	1:50	1:50	2 µL
2 µg	1:20	1:50	1:25	2 µL

2. Add 2 µL of Poly-A RNA Control Stock to 38 µL of Poly-A Control Dil Buffer to make the First Dilution (1:20).
3. Mix and spin to collect the solution at the bottom of the tube.
4. Add 2 µL of the First Dilution to 98 µL of Poly-A Control Dil Buffer to make the Second Dilution (1:50).
5. Mix and spin to collect the solution at the bottom of the tube.
6. **When using 1 µg of total RNA as starting material**, add 2 µL of the Second Dilution to 98 µL of Poly-A Control Dil Buffer to make the Third Dilution (1:50).
When using 2 µg of total RNA as starting material, add 2 µL of the Second Dilution to 48 µL of Poly-A Control Dil Buffer to make the Third Dilution (1:25).
7. Mix and spin to collect the solution at the bottom of the tube. Add 2 µL of the appropriate Third Dilution to 1 µg or 2 µg of total RNA to make up the Total RNA/Poly-A RNA Controls Mix as described in [Table 2.1](#).

Procedure B: Preparation of Hybridization Buffer with Betaine

This Procedure requires the use of the RiboMinus Human/Mouse Transcriptome Isolation Kit that needs to be obtained directly from Invitrogen.

! **IMPORTANT:** The protocol for using the RiboMinus Kit for rRNA reduction for GeneChip® arrays has been modified from the manufacturer's original recommendation. Follow only the procedure described in this manual for optimal transition to the target labeling process immediately following the rRNA reduction steps.

1. Prepare the buffer by mixing these components as listed in [Table 2.2](#). Keep tube at room temperature.

Table 2.2 Hybridization Buffer with Betaine

Component	Starting Total RNA Amount	
	1 µg	2 µg
	Volume for 1 Rxn	Volume for 1 Rxn
Betaine, 5M	54 µL	84 µL
Invitrogen Hybridization buffer	126 µL	196 µL
Total Volume	180 µL*	280 µL*

* A 30 µL overfill is included in the Total Volume, sufficient for completing the RiboMinus procedure for a single total RNA sample.

Procedure C: RiboMinus Probe Hybridization

This Procedure requires the use of the RiboMinus Human/Mouse Transcriptome Isolation Kit that needs to be obtained directly from Invitrogen.

1. In a 0.2 mL strip tube, mix the following components in [Table 2.3](#). Keep the tubes on ice. If the total RNA is in 1 µg/µL of concentration, follow the instructions in the second column. If the total RNA sample is at a lower concentration of between 0.31 µg/µL to 1 µg/µL, then follow the protocol listed in the third column of [Table 2.3](#).



NOTE: It is recommended to prepare a master mix including the RiboMinus probe and the Hybridization Buffer with Betaine and then add the combined amount (see [Table 2.3](#)) to the tube containing the Total RNA/Poly-A RNA Control Mix.

Table 2.3 RiboMinus Reaction and Wash Volumes

Total RNA Amount	1 µg	1 µg	2 µg
RNA Concentration Range	1 µg/µL	0.31 µg/µL to 1 µg/µL	≥ 0.62 µg/µL
RiboMinus Probe Hybridization			
Total RNA/Poly-A Controls Mix (from Procedure A)	3.0 µL	up to 5.2 µL	up to 5.2 µL
RiboMinus Probe (100 pmol/µL)	0.8 µL	0.8 µL	1.6 µL
Hybridization Buffer with Betaine (from Procedure B)	20 µL	30 µL	40 µL
Total Hybridization Volume	23.8 µL	36.0 µL	46.8 µL
Beads Preparation			
Magnetic Beads	50 µL	50 µL	100 µL
2X Wash with Water	50 µL	50 µL	100 µL
1X Wash with Hybridization Buffer (with Betaine)	50 µL	50 µL	100 µL
Re-suspension of Beads with Hybridization Buffer (with Betaine)	30 µL	20 µL	60 µL
Final Wash of Beads with Hybridization Buffer (with Betaine)	50 µL	50 µL	50 µL
Concentration and Clean-up			
Approximate Supernatant Volume	~100 µL	~100 µL	~150 µL
cRNA Binding Buffer	350 µL	350 µL	525 µL
100% Ethanol	250 µL	250 µL	375 µL
Total Volume Applied to Column	~700 µL	~700 µL	~1,050 µL*
cRNA Wash Buffer	500 µL	500 µL	500 µL
100% Ethanol	500 µL	500 µL	500 µL

* Multiple applications to column required (column capacity ≤ 800 µL).

2. Flick the tube gently to mix, spin briefly and incubate at 70°C for 5 minutes in a thermal cycler.
3. Quench the reaction immediately by placing the tube on ice while preparing the magnetic beads.


Procedure D: Preparation of Beads

This Procedure requires the use of the RiboMinus Human/Mouse Transcriptome Isolation Kit that needs to be obtained directly from Invitrogen.

 **NOTE:** Procedures D and E require the use of a 37°C heat block and a 50°C heat block. Pre-heat the heat blocks prior to initiating these steps.

A heating block at 37°C is required. Use the volumes between parenthesis if the starting amount of total RNA is 2 µg.

1. Completely re-suspend the bottle containing magnetic beads by flicking it until no deposit is observed at the bottom of the bottle.
2. Pipet 50 µL (100 µL) of beads suspension into a 1.5 mL non-stick RNase-free tube. Steps 3 to 6 are done at room temperature.
3. Briefly spin and place the tube with the beads suspension on the magnetic stand for ~1 minute. With the tube remaining in the stand, gently aspirate and discard the supernatant.

 **NOTE:** Drying of the beads decreases the bead efficiency, therefore ensure that the beads are hydrated at all times. It is recommended to handle a limited number of tubes at one time to reduce the risk of drying the beads.

4. 1st Wash:

- A. Add 50 µL (100 µL) of RNase-free water to the beads and re-suspend them by flicking the tube.

 **NOTE:** If the beads are still attached to the wall of the tube, put the tube back into the magnetic stand and rotate the tube with quick motion until the beads are in suspension.

- B. Spin briefly. Place the tube on the magnetic stand for ~1 minute. With the tube remaining in the stand, gently aspirate and discard the supernatant.

5. **2nd Wash:** Add 50 µL (100 µL) of RNase-free water to the beads and re-suspend them by flicking the tube. Place the tube on the magnetic stand for ~1 minute. With the tube remaining in the stand, gently aspirate and discard the supernatant.

- 6. 3rd Wash:** Add 50 μL (100 μL) of the **Hybridization Buffer with Betaine** (from Procedure B) to the beads and re-suspend them by flicking the tube. Spin briefly. Place the tube on the magnetic stand for ~ 1 minute. With the tube remaining in the stand, gently aspirate and discard the supernatant.



NOTE: When the Hybridization Buffer with Betaine is added in the 3rd Wash, even after complete settling of the beads on the magnetic stand, a thin film of beads may coat the entire inside of the tube, so the solution may appear yellow. Carefully aspirate out the supernatant from the center of the tube, which should be clear in color.

- 7.** Re-suspend the beads in **Hybridization Buffer with Betaine**. Keep them at 37°C in a heating block for 1 to 2 minutes. Refer to [Table 2.3](#) for appropriate volume.



NOTE: If there is a large number of samples to be processed at the same time the beads can be prepared in a large batch. This can be done with up to a total of 10 samples, by increasing the volume of the beads and wash solutions proportionately, and carrying out the wash steps in a single tube. Make sure to gently flick the tube with the beads several times for a thorough mixing after each step. Use the magnetic stand (as described under Step 4) if the beads remain attached to the wall of the tube. Following the 3rd Wash (Procedure D, Step 6), the resuspended beads can then be aliquoted to individual tubes as described in Procedure D, Step 7 before proceeding to rRNA Reduction for individual samples.

Procedure E: rRNA Reduction

This Procedure requires the use of the RiboMinus Human/Mouse Transcriptome Isolation Kit that needs to be obtained directly from Invitrogen.

Two heating blocks are required: one at 37°C, and the other at 50°C. Use the volumes between parenthesis if the starting amount of total RNA is 2 µg.

1. Transfer the ice-cooled hybridized sample prepared in Procedure C to the beads prepared in Procedure D, mix well, and briefly spin.
2. Incubate the tube with the mixture at 37°C for 10 minutes in a heating block. After 5 minutes of incubation, gently flick-mix the tube.
3. Briefly spin and place the tube in the magnetic stand for 1 to 2 minutes to obtain the rRNA-probe pellet.



NOTE: The supernatant contains the rRNA-Reduced Total RNA/Poly-A RNA Controls Mix.

4. With the tube in the magnetic stand, transfer the supernatant to a 1.5 mL non-stick RNase-free tube and leave on ice.
5. Wash the beads by re-suspending them in 50 µL (50 µL) of **Hybridization Buffer with Betaine** and incubate at 50°C for 5 minutes.
6. Place the tube in the magnetic stand for 1 to 2 minutes, transfer the supernatant, and combine with the supernatant in the tube from Procedure E, Step 4. The total volume of the rRNA-reduced sample is approximately 100 µL (150 µL).

Procedure F: Concentration

This Procedure requires the use of the GeneChip® IVT cRNA Cleanup Kit.



NOTE: This Procedure uses the IVT cRNA Cleanup Spin Columns in place of the purification columns as part of the RiboMinus Kit. Carefully follow the protocol described here for best results.

1. Proceed to the cleanup procedure using the IVT cRNA Cleanup Spin Columns from the IVT cRNA Cleanup Kit following the protocol described below. Use the volumes between parenthesis if the starting amount of total RNA is 2 µg.
2. If not already done, add 20 mL of Ethanol (100%) to the cRNA Wash Buffer supplied in the IVT cRNA Cleanup Kit.
3. Add 350 µL (525 µL) of cRNA Binding Buffer to each rRNA-reduced sample from Procedure E and vortex for 3 seconds.
4. Add 250 µL (375 µL) of 100% ethanol to each reaction and flick the tube to mix.
5. Apply the sample to the IVT cRNA Cleanup Spin Column sitting in a 2 mL Collection Tube.



IMPORTANT: If the starting amount of total RNA is 2 µg, the total supernatant volume will exceed the capacity of the column. Apply ~700 µL, centrifuge for 15 seconds at ≥ 8,000 x g. Discard the flow-through. Place the column back in to 2 mL collection tube and apply the remaining sample (~350 µL). Continue to Step 6.

6. Centrifuge for 15 seconds at ≥ 8,000 x g. Discard the flow-through.
7. Transfer the IVT cRNA Cleanup Spin Column to a new 2 mL Collection Tube. Add 500 µL of cRNA Wash Buffer and centrifuge for 15 seconds at ≥ 8,000 x g. Discard the flow-through.
8. Wash again with 500 µL of 80% (v/v) ethanol. Centrifuge for 15 seconds at ≥ 8,000 x g and discard the flow-through.
9. Open column cap and spin at ≤ 25,000 x g (maximum speed) for 5 minutes with the cap left open.
10. Transfer the IVT cRNA Cleanup Spin Column to a new 1.5 mL Collection Tube and add 11 µL of RNase-free Water directly to the membrane. Spin at ≤ 25,000 x g (maximum speed) for 1 minute.
11. The eluted rRNA-Reduced Total RNA/Poly-A RNA Controls Mix is ~9.8 µL. Keep sample on ice. Proceed with Procedure G (Analysis with Bioanalyzer) and [Chapter 3](#), Procedure A.



NOTE: Store eluted rRNA-reduced total RNA at -80°C if not proceeding immediately to Procedure G (Analysis with Bioanalyzer) and [Chapter 3](#), Procedure A.

Procedure G: Analysis with Bioanalyzer

1. Use 1 μL of the concentrated sample to check its quality by running the Eukaryotic Total RNA Nano Assay in the Bioanalyzer. Please see the Reagent Kit Guide provided with the RNA 6000 Nano LabChip Kit for instructions. Based on the Bioanalyzer results, on average, >100 ng of rRNA-reduced sample may be recovered from 1 μg of total RNA starting material but the recovery rate may vary depending on the tissue type. See [Figure 2.1](#) for an example of the results.



NOTE: It is recommended to analyze 100 ng (1 μL) of total RNA sample without RiboMinus Kit treatment as a control.

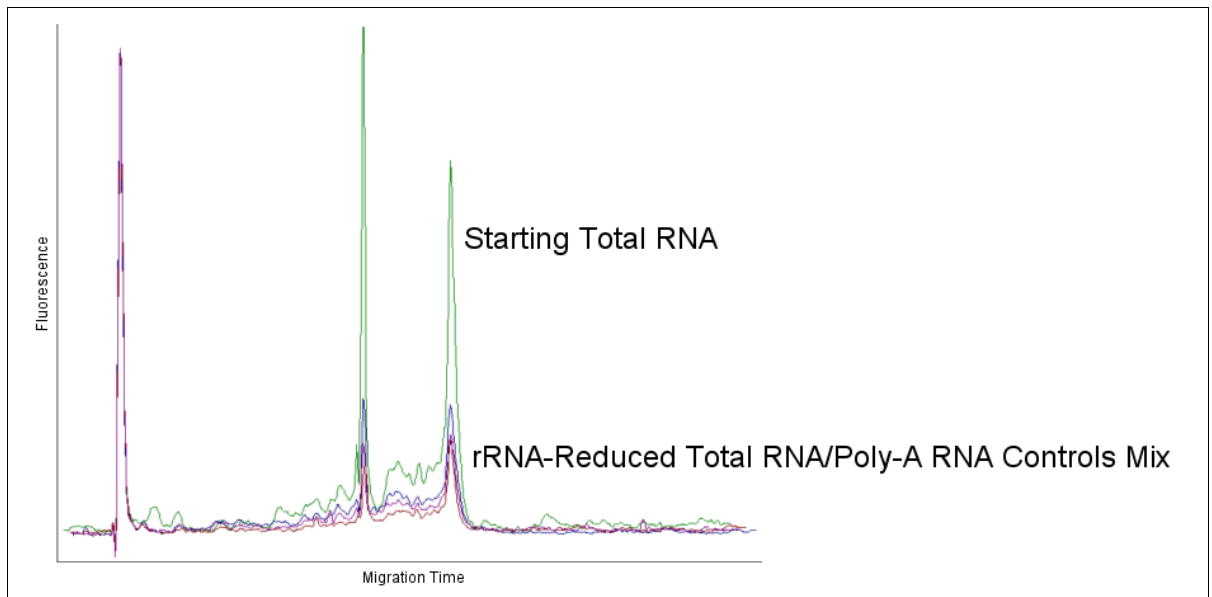


Figure 2.1 Bioanalyzer profile of Human Brain total RNA before and after rRNA reduction with RiboMinus Kit (treated samples are in triplicates)

1 μg TOTAL RNA TARGET LABELING PROTOCOL



NOTE: The 1 μg Total RNA Labeling Protocol is recommended for use with GeneChip Exon 1.0 ST Arrays. To prepare targets for GeneChip Gene 1.0 ST Arrays, follow the 100 ng Total RNA Labeling Protocol in [Chapter 4](#).

Procedure A: Preparation of rRNA-Reduced Total RNA/Poly-A RNA Controls/T7-(N)₆ Primers Mix

This Procedure requires the use of the GeneChip[®] WT cDNA Synthesis Kit.

1. Dilute the T7-(N)₆ Primers, 2.5 $\mu\text{g}/\mu\text{L}$ stock 1:5 with RNase-free water to make up a 500 ng/ μL working solution. Keep on ice. Prepare fresh every time.
2. Mix the diluted T7-(N)₆ Primers with the Concentrated rRNA-Reduced Sample from Chapter 2, Procedure F as listed in [Table 3.1](#), using 0.2 mL strip tubes on ice.

Table 3.1 First-cycle, Preparation of rRNA-reduced Total RNA/Poly-A RNA Controls/T7-(N)₆ Primers

Component	Volume in 1 Rxn
rRNA-Reduced Total RNA/Poly-A RNA Controls Mix	4 μL
Diluted T7-(N) ₆ Primers, 500 ng/ μL	1 μL
Total Volume	5 μL

3. Flick the tube to mix, spin down the tube, and incubate at:
 - 70°C for 5 minutes
 - 4°C for at least 2 minutes
4. Spin down and place on ice for use in Procedure B.

Procedure B: First-Cycle, First-Strand cDNA Synthesis

This Procedure requires the use of the GeneChip® WT cDNA Synthesis Kit.

1. Prepare the First-Cycle, First-Strand Master Mix as shown in [Table 3.2](#). Add the SuperScript™ II enzyme to the master mix last and proceed immediately to aliquot into the tubes from Procedure A, Step 4, as described in Step 2 below.

Table 3.2 First-Cycle, First-Strand Master Mix

Component	Volume in 1 Rxn
5X 1 st Strand Buffer	2 µL
DTT, 0.1M	1 µL
dNTP Mix, 10 mM	0.5 µL
RNase Inhibitor	0.5 µL
SuperScript II	1 µL
Total Volume	5 µL

2. Add 5 µL of the First-Cycle, First-Strand Master Mix to the tube containing the Concentrated rRNA-Reduced Total RNA/Poly-A RNA Controls/ T7-(N)₆ Primers Mix from Procedure A, flick-mix, and spin-down. The total reaction volume is 10 µL.
3. Incubate the reaction at:
 - 25°C for 10 minutes
 - 42°C for 60 minutes
 - 70°C for 10 minutes
4. Cool the reaction to 4°C for at least 2 minutes before immediately continuing to the First-Cycle, Second-Strand cDNA Synthesis.



NOTE: Keeping the reaction at 4°C longer than 10 minutes may result in reduced cRNA yields.

Procedure C: First-Cycle, Second-Strand cDNA Synthesis

This Procedure requires the use of the GeneChip® WT cDNA Synthesis Kit.

1. Make a fresh dilution of 17.5 mM MgCl₂ each time. Mix 2 µL of 1M MgCl₂ with 112 µL of RNase-free water.
2. Prepare the First-Cycle, Second-Strand Master Mix as described in [Table 3.3](#). Add the RNase H and DNA Polymerase I enzymes to the master mix last and proceed immediately to aliquot into the tubes from Procedure B, Step 4 as described in Step 3 below.

Table 3.3 First-Cycle, Second-Strand Master Mix

Component	Volume in 1 Rxn
RNase-free Water	4.8 µL
MgCl ₂ , 17.5 mM	4.0 µL
dNTP Mix, 10 mM	0.4 µL
DNA Polymerase I	0.6 µL
RNase H	0.2 µL
Total Volume	10.0 µL

3. Add 10 µL of the First-Cycle, Second-Strand Master Mix to the reaction tube from the First-Strand cDNA Synthesis Reaction in Procedure B for a total reaction volume of 20 µL. Flick or gently vortex the tubes and spin down.
4. Incubate the reaction in a thermal cycler at:
 - 16°C for 120 minutes without heated lid
 - 75°C for 10 minutes with heated lid
5. Cool the sample for at least 2 minutes at 4°C before immediately proceeding to the next Procedure: First-Cycle, cRNA Synthesis and Cleanup.



NOTE: Keeping the reaction at 4°C longer than 10 minutes may result in reduced cRNA yields.

Procedure D: First-Cycle, cRNA Synthesis and Cleanup

This Procedure requires the use of the GeneChip® WT cDNA Amplification Kit and the GeneChip® Sample Cleanup Module.

1. In a separate tube, assemble the IVT Master Mix at room temperature as listed in [Table 3.4](#). Add the IVT Enzyme Mix to the master mix last and proceed immediately to aliquot into the tubes from Procedure C, Step 5, as described in Step 5 below.



NOTE: If a white precipitate is still present in the 10X IVT Buffer after thawing, incubate the tube at 37°C until the precipitate gets dissolved. Do not assemble the reaction on ice since the spermidine in the 10X IVT Buffer can lead to precipitation of the template DNA.

Table 3.4 First-Cycle, IVT Master Mix

Component	Volume in 1 Rxn
10X IVT Buffer	5.0 µL
IVT NTP Mix	20.0 µL
IVT Enzyme Mix	5.0 µL
Total Volume	30.0 µL

2. Transfer 30 µL of the IVT Master Mix to each First-Cycle cDNA Synthesis Reaction sample from Procedure C to a final volume of 50 µL. Flick-mix the solution, and briefly spin in a microfuge.
3. Incubate the reaction for 16 hours at 37°C. To prevent condensation that may result from water bath-style incubators, incubations are best performed in oven incubators for even temperature distribution, or in a thermal cycler. Hold at 4°C.
4. Proceed to the cleanup procedure for cRNA using the cRNA Cleanup Spin Columns from the GeneChip Sample Cleanup Module following the protocol described below. Store the sample at –80°C if not purifying the cRNA immediately.
5. If not already done, add 20 mL of Ethanol (100%) to the cRNA Wash Buffer supplied in the GeneChip Sample Cleanup Module.
6. Add 50 µL of RNase-free water to each IVT reaction to a final volume of 100 µL.
7. Add 350 µL of cRNA Binding Buffer to each sample and vortex for 3 seconds.
8. Add 250 µL of 100% ethanol to each reaction and flick-mix.
9. Apply the sample to the IVT cRNA Cleanup Spin Column sitting in a 2 mL Collection Tube.
10. Centrifuge for 15 seconds at $\geq 8,000 \times g$. Discard the flow-through.

11. Transfer the IVT cRNA Cleanup Spin Column to a new 2 mL Collection Tube. Add 500 µL of cRNA Wash Buffer to column and centrifuge for 15 seconds at $\geq 8,000 \times g$. Discard the flow-through.
12. Wash again with 500 µL of 80% (v/v) Ethanol. Centrifuge for 15 seconds at $\geq 8,000 \times g$ and discard the flow-through.
13. Open the column cap and spin at $\leq 25,000 \times g$ (maximum speed) for 5 minutes with the caps open.
14. Transfer the IVT cRNA Cleanup Spin Column to a new 1.5 mL Collection Tube and add 15 µL of RNase-free water directly to the membrane. Incubate at room temperature for 5 minutes. Spin at $\leq 25,000 \times g$ (maximum speed) for 1 minute.
15. Elute a second time by pipetting the flow-through in the Collection Tube (~13.5 µL) back onto the Spin Column membrane. Place the Spin Column back into the Collection Tube and incubate at room temperature for 5 minutes. Spin at $\leq 25,000 \times g$ (maximum speed) for 1 minute.
16. The eluted cRNA is ~ 13.5 µL. Determine the cRNA yield by spectrophotometric UV measurement at 260 nm, 280 nm, and 320 nm:
Concentration of cRNA ($\mu\text{g}/\mu\text{L}$) = $[A_{260} - A_{320}] \times 0.04 \times \text{dilution factor}$
 $\mu\text{g of cRNA} = \text{eluate in } \mu\text{L} \times \text{concentration of cRNA in } \mu\text{g}/\mu\text{L}$
The NanoDrop® ND-1000 can also be used to measure the concentration.

NOTE: The average yield may vary depending on the type of tissue used and the quality of the RiboMinus rRNA reduction step.

NOTE: Store eluted cRNA at -80°C if not proceeding immediately to Procedure E.

NOTE: If cRNA concentration is too low to obtain 8 to 10 µg of cRNA in 6.5 µL, cRNA can be concentrated using a SpeedVac.

Procedure E: Second-Cycle, First-Strand cDNA Synthesis

This Procedure requires the use of the GeneChip® WT cDNA Synthesis Kit.

1. Mix cRNA sample from Procedure D with the Random Primers in a strip tube, as listed in [Table 3.5](#) below.

Table 3.5 Second-Cycle, cRNA/Random Primers Mix

Component	Volume in 1 Rxn
cRNA, 10 µg*	variable
Random Primers (3 µg/µL)	1.5 µL
RNase-free water	up to 8 µL
Total Volume	8.0 µL

*For some samples that generate high yield in the second-cycle cDNA synthesis reaction, 8 µg may be used to obtain 5 µg of single-stranded DNA target. The starting amount of cRNA varies depending on the type of tissue used.

2. Flick-mix and spin down the tubes.
3. Incubate the Second-Cycle, cRNA/Random Primers Mix at:
 - 70°C for 5 minutes
 - 25°C for 5 minutes
4. Cool the samples at 4°C for at least 2 minutes.
5. In a separate tube, prepare the Second-Cycle, Reverse Transcription Master Mix as described in [Table 3.6](#). Add the SuperScript™ II enzyme to the master mix last and proceed immediately to aliquot into tubes from Step 4 as described in Step 6, [on page 27](#).

Table 3.6 Second-Cycle, First-Strand cDNA Synthesis Master Mix

Component	Volume in 1 Rxn
5X 1 st Strand Buffer	4.0 µL
DTT, 0.1M	2.0 µL
dNTP+dUTP, 10 mM	1.25 µL
SuperScript II	4.75 µL
Total Volume	12.0 µL

- 6.** Transfer 12 μ L of the Second-Cycle, First-Strand cDNA Synthesis Master Mix to the Second-Cycle, cRNA/Random Primers Mix from Procedure E, Step 4 for a total reaction volume of 20 μ L. Mix thoroughly by gently flicking the tubes a few times and centrifuge briefly.
- 7.** Incubate the reactions at:
 - 25°C for 10 minutes
 - 42°C for 90 minutes
 - 70°C for 10 minutes
 - 4°C for at least 2 minutes

Procedure F: Hydrolysis of cRNA and Cleanup of Single-Stranded DNA

This Procedure requires the use of the GeneChip® WT cDNA Synthesis Kit and the GeneChip® Sample Cleanup Module.

1. Add 1 μL of RNase H to each of the samples and incubate at:
 - 37°C for 45 minutes
 - 95°C for 5 minutes
 - 4°C for 2 minutes
2. Proceed to the cleanup step using the cDNA Cleanup Spin Columns from the GeneChip Sample Cleanup Module following the protocol as described below. Store the sample at -20°C if not purifying the Single-Stranded DNA immediately.
3. If not already done, add 24 mL of Ethanol (100%) to the cDNA Wash Buffer supplied in the GeneChip Sample Cleanup Module.
4. Add 80 μL of RNase-free water to each sample, followed by 370 μL of cDNA Binding Buffer, and vortex for 3 seconds.
5. Apply the entire sample (the total volume is 471 μL) to a cDNA Spin Column sitting in a 2 mL Collection Tube.
6. Spin at $\geq 8,000 \times g$ for 1 minute. Discard the flow-through.
7. Transfer the cDNA Cleanup Spin Column to a new 2 mL Collection Tube and add 750 μL of cDNA Wash Buffer to the column. Spin at $\geq 8,000 \times g$ for 1 minute and discard the flow-through.
8. Open the cap of the cDNA Cleanup Spin Column and spin at $\leq 25,000 \times g$ for 5 minutes with the caps open. Discard the flow-through, and place the column in a 1.5 mL collection tube.
9. Pipet 15 μL of the cDNA Elution Buffer directly to the column membrane and incubate at room temperature for 1 minute. Then, spin at $\leq 25,000 \times g$ for 1 minute.
10. Repeat the elution step by pipetting another 15 μL of the cDNA Elution Buffer directly to the column membrane and incubate at room temperature for 1 minute. Then, spin at $\leq 25,000 \times g$ for 1 minute.
11. The total volume of the eluted Single-Stranded DNA is $\sim 28 \mu\text{L}$ total. Take 2 μL from each sample to determine the yield by spectrophotometric UV measurement at 260 nm, 280 nm, and 320 nm:

Concentration of Single-Stranded DNA ($\mu\text{g}/\mu\text{L}$) = $[A_{260} - A_{320}] \times 0.033 \times \text{dilution factor}$
 μg of DNA = eluate in μL \times concentration of DNA in $\mu\text{g}/\mu\text{L}$

Each tube should have $\geq 5.5 \mu\text{g}$ of Single-Stranded DNA.

The NanoDrop® ND-1000 can also be used to measure the concentration.



NOTE: Eluted single-stranded cDNA can be stored overnight at -20°C if not proceeding immediately to Procedure G.

Procedure G: Fragmentation of Single-Stranded DNA

This Procedure requires the use of the GeneChip® WT Terminal Labeling Kit.

1. Set up fragmentation reaction in 0.2 mL strip tubes using [Table 3.7](#).

Table 3.7 Fragmentation Master Mix

Component	Volume/Amount in 1 Rxn
Single-Stranded DNA	5.5 µg
RNase-free Water	up to 31.2 µL
Total Volume	31.2 µL

2. Prepare the Fragmentation Master Mix using [Table 3.8](#).

Table 3.8 Fragmentation Master Mix

Component	Volume in 1 Rxn
RNase-free Water	10 µL
10X cDNA Fragmentation Buffer	4.8 µL
UDG, 10 U/µL	1.0 µL
APE 1, 1,000 U/µL	1.0 µL
Total Volume	16.8 µL

3. Add 16.8 µL of the above Fragmentation Master Mix to the samples prepared in Step 1. Flick or gently vortex the tubes and spin down.
4. Incubate the reactions at:
 - 37°C for 60 minutes
 - 93°C for 2 minutes
 - 4°C for at least 2 minutes
5. Flick-mix, spin down the tubes, and transfer 45 µL of the sample to a new 0.2 mL strip tube. The remainder of the sample can be used for size analysis using a Bioanalyzer. Please see the Reagent Kit Guide that comes with the RNA 6000 Nano LabChip Kit for detailed instructions. The range in peak size of the fragmented samples should be approximately 40 to 70 nt. See [Figure 3.1](#) as an example of typical results on fragmented samples.
6. If the samples are not labeled immediately, store the fragmented Single-Stranded DNA at -20°C.

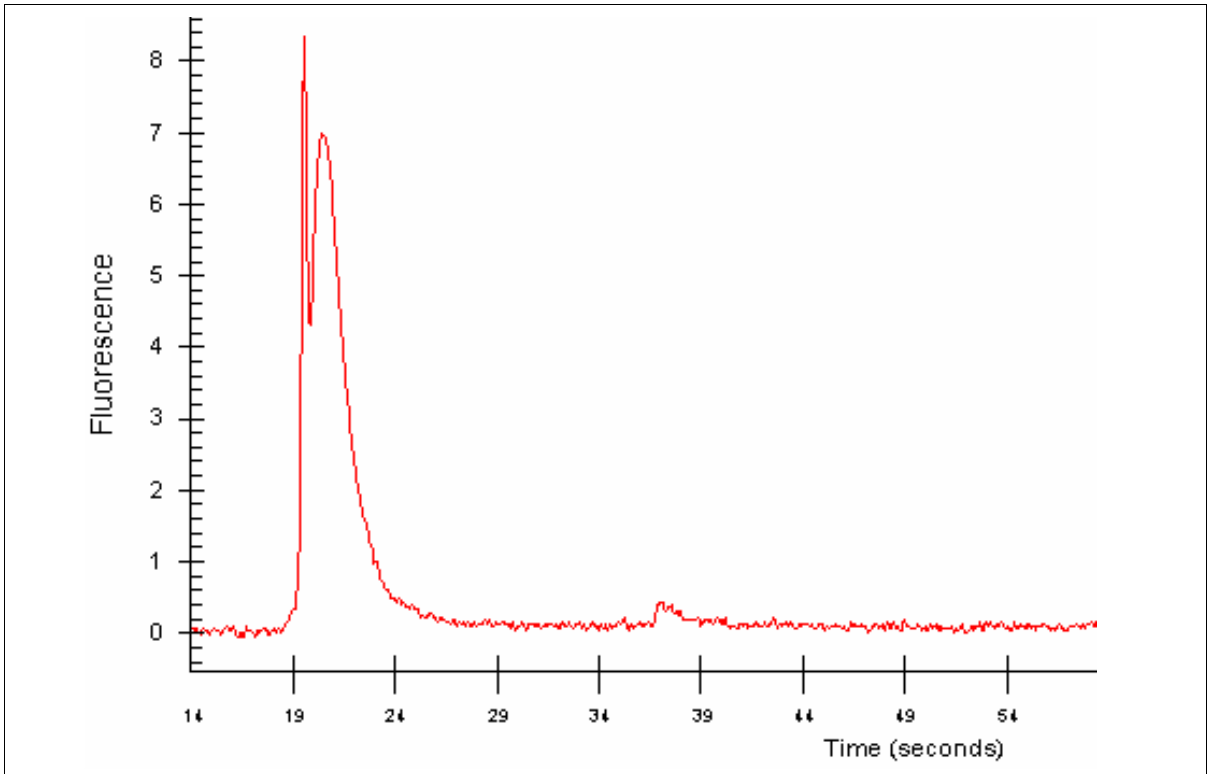


Figure 3.1 Bioanalyzer profile of Fragmented Single-Stranded DNA from Human Brain

Procedure H: Labeling of Fragmented Single-Stranded DNA

This Procedure requires the use of the GeneChip® WT Terminal Labeling Kit.

1. Prepare the labeling reactions as listed in [Table 3.9](#). A master mix using the 5X TdT Buffer, TdT and DNA Labeling reagent can be prepared just before aliquoting 15 µL into the 0.2 mL strip tubes containing the 45 µL of Fragmented Single-Stranded DNA.

Table 3.9 Labeling Reaction

Component	Volume in 1 Rxn
Fragmented Single-Stranded DNA (from Procedure G)	45 µL
5X TdT Buffer	12 µL
TdT	2 µL
DNA Labeling Reagent, 5 mM	1 µL
Total Volume	60 µL

2. After adding the labeling reagents to the fragmented DNA samples, flick-mix and spin them down.
3. Incubate the reactions at:
 - 37°C for 60 minutes
 - 70°C for 10 minutes
 - 4°C for at least 2 minutes
4. Remove 2 µL of each sample for Gel-shift analysis (optional) as described in [Appendix B](#), to assess the labeling efficiency.

100 ng Total RNA Labeling Protocol



NOTE: The 100 ng Total RNA Labeling Protocol is recommended to be used with the GeneChip Gene 1.0 ST Arrays. For optimal sensitivity on the GeneChip Exon 1.0 ST Arrays, the 1 µg Total RNA Labeling Protocol ([Chapter 2](#)) should be followed.

Procedure A: Preparation of dilutions of Poly-A RNA Controls

Follow the same procedure as described in [Chapter 2, Procedure A on page 11](#).

Procedure B: Preparation of T7-(N)₆ Primers/Poly-A RNA Controls

1. Prepare a fresh 250 ng/µL T7-(N)₆ Primers dilution (from a 2.5 µg/µL stock) by adding the concentrated T7-(N)₆ Primers to the diluted Poly-A RNA controls using a non-stick RNase-free microfuge tube as follows:

Table 4.1 First-cycle, Primer/Poly-A RNA Controls

Component	Volume
T7-(N) ₆ Primers, 2.5 µg/µL	2 µL
Diluted Poly-A RNA Controls (3rd dilution, 1:50)	2 µL
RNase-free Water	16 µL
Total Volume	20 µL

2. Flick-mix the solution, spin down, and place on ice.

Procedure C: Preparation of total RNA/T7-(N)₆ Primers/Poly-A RNA Controls

1. Mix total RNA and the T7-(N)₆ Primers/Poly-A RNA Controls solution as listed in [Table 4.2](#).

Table 4.2 First-Cycle, Total RNA/Primer/Poly-A RNA Controls

Component	Volume in 1 Rxn
Total RNA, 100 – 300 ng	variable
T7-(N) ₆ Primers/Poly-A RNA Controls Solution	2 µL
RNase-free Water	up to 5 µL
Total Volume	5 µL

2. Flick-mix, spin down the tube, and incubate for 5 minutes at 70°C. Then, cool the sample for at least 2 minutes at 4°C and spin down.
3. Place on ice for use in the section below.

Follow the same protocol as described in [Chapter 3, Procedures B-H \(starting on page 22\)](#) for the remainder of the Whole Transcript Sense Target Labeling Assay.

Hybridization

This Procedure requires the use of the GeneChip® Hybridization, Wash and Stain Kit. Three heating blocks are required: one at 65°C, one at 99°C, and the third one at 45°C.

1. Prepare the Hybridization Cocktail in a 1.5 mL RNase-free microfuge tube as shown in [Table 5.1](#).

Table 5.1 Hybridization Cocktail

Component	Volume for One 49/64 Format Array	Volume for One 169 Format Array	Final Concentration
Fragmented and Labeled DNA Target (from Chapter 3)	~60.0* μL	27 μL	~25 ng/ μL
Control Oligonucleotide B2 (3 nM)	3.7 μL	1.7 μL	50 pM
20X Eukaryotic Hybridization Controls (<i>bioB</i> , <i>bioC</i> , <i>bioD</i> , <i>cre</i>)	11 μL	5 μL	1.5, 5, 25 and 100 pM, respectively
2X Hybridization Mix	110 μL	50 μL	1X
DMSO	15.4 μL	7 μL	7%
Nuclease-free Water	up to 220.0 μL	up to 100	
Total Volume	220.0 μL	100 μL	

* This volume is 58 μL if a portion of the sample was set aside for Gel-shift analysis.



IMPORTANT: It is imperative that frozen stocks of 20X GeneChip® Eukaryotic Hybridization Controls are heated to 65°C for 5 minutes to completely resuspend the cRNA before aliquoting.

2. Flick or gently vortex the tubes and spin down.

3. Heat the Hybridization Cocktail at 99°C for 5 minutes. Cool to 45°C for 5 minutes, and centrifuge at maximum speed for 1 minute.
4. Equilibrate the GeneChip ST Array to room temperature immediately before use. Label the array with the name of the sample that will be hybridized.
5. Inject the appropriate amount (see [Table 5.2](#)) of the specific sample into the array through one of the septa (see [Figure 5.1](#) for location of the septa on the array).

Table 5.2 Probe Array Cartridge Volumes for Hybridization Cocktail

Array Format	Volume
49 (Standard)	200 µL
64	200 µL
169	80 µL

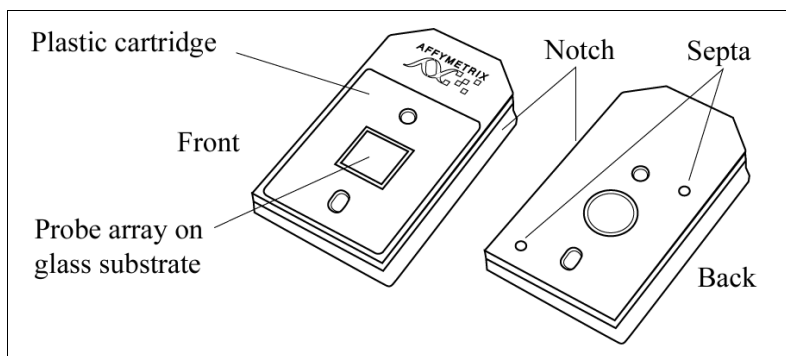


NOTE: It is necessary to use two pipette tips when filling the probe array cartridge: one for filling and the second to allow venting of air from the hybridization chamber.



NOTE: Ensure that the bubble inside the hyb chamber floats freely upon rotation to allow the hybridization cocktail to make contact with all portions of the array.

6. Place array in 45°C hybridization oven, at 60 rpm, and incubate for 17 hours ± 1 hour. During the latter part of the array hybridization, commence preparation of the reagents required immediately after completion of hybridization.

**Figure 5.1** GeneChip® Probe Array

Procedure A: Entering Experiment Information

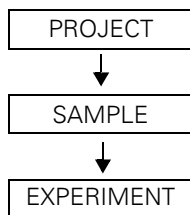
To wash, stain, and scan a probe array, an experiment must first be registered in GeneChip® Operating Software (GCOS). Please follow the instructions detailed in the “Setting Up an Experiment” section of the GCOS User’s Guide.

The fields of information required are:

- Experiment Name
- Probe Array Type
- Sample Name
- Sample Type
- Project

Sample templates, experiment templates, and array barcodes can also be employed in GCOS to standardize and simplify the registration process. Please see the GCOS User’s Guide for more information.

The Project, Sample Name, and Experiment Name fields establish a sample hierarchy that organizes GeneChip data in GCOS. In terms of the organizational structure, the Project is at the top of the hierarchy, followed by Sample Name and then Experiment Name.



Procedure B: Preparing the Fluidics Station

This Procedure requires the use of the GeneChip® Hybridization, Wash and Stain Kit.

The GeneChip® Fluidics Station 450/250 is used to wash and stain the GeneChip® ST Arrays. It is operated using GCOS.

Setting Up the Fluidics Station

1. Turn on the Fluidics Station using the toggle switch on the lower left side of the machine.
2. Select **Run** → **Fluidics** from the menu bar.
The Fluidics Station dialog box appears with a drop-down list for selecting the experiment name for each of the fluidics station modules. A second drop-down list is accessed for choosing the Protocol for each of the fluidics station modules.



NOTE: Refer to the *Fluidics Station User's Guide* for instructions on connecting and addressing multiple fluidics stations.

Priming the Fluidics Station

Priming ensures that the lines of the fluidics station are filled with the appropriate buffers and the fluidics station is ready for running fluidics station protocols.

Priming should be done:

- when the fluidics station is first started.
 - when wash solutions are changed.
 - before washing, if a shutdown has been performed.
 - if the LCD window instructs the user to prime.
1. To prime the fluidics station, select **Protocol** in the Fluidics Station dialog box.
 2. Choose **Prime_450** for the respective modules in the Protocol drop-down list.
 3. Ensure that the designated Fluidics Station Wash A and Wash B media bottles are clean. Transfer **Wash Buffer A** and **Wash Buffer B** from the kit to the clean, empty Fluidics Station bottles.
 4. Select the **All Modules** check box, then click **Run**.

Procedure C: Probe Array Washing and Staining

This Procedure requires the use of the GeneChip® Hybridization, Wash and Stain Kit.

After 17 hours \pm 1 hour of hybridization remove the array from the hybridization oven. Vent the array by inserting a clean pipette tip into one of the septa, and extract the hybridization cocktail with a pipettor through the remaining septum. Refill the probe array completely with the appropriate volume of Wash Buffer A, as given in [Table 6.1](#).

Table 6.1 Probe Array Cartridge Volumes for Wash Buffer A and Array Holding Buffer

Array	Volume
49 Format	250 μ L
64 Format	250 μ L
169 Format	100 μ L



NOTE: If necessary, at this point, the probe array can be stored at 4°C for up to 3 hours before proceeding with washing and staining. Equilibrate the probe array to room temperature before washing and staining.

The wash and stain procedure takes approximately 90 minutes to complete.

Preparing the Staining Reagents

Prepare the following reagents. Volumes given are sufficient for one probe array.

1. Remove **Stain Cocktail 1**, **Stain Cocktail 2**, and **Array Holding Buffer** from the Stain Module, Box 1.
2. Gently tap the bottles to mix well.
3. Aliquot the following reagents:
 - A. 600 μL of **Stain Cocktail 1** into a 1.5 mL amber microcentrifuge vial.
 - B. 600 μL of **Stain Cocktail 2** into a 1.5 mL (clear) microcentrifuge vial.
 - C. 800 μL of **Array Holding Buffer** into a 1.5 mL (clear) microcentrifuge vial.
4. Spin down all vials to remove the presence of any air bubbles.



NOTE: Stain Cocktail 1 is light-sensitive. Please be sure to use amber microcentrifuge vials when aliquoting.

Fluidics Protocols

Table 6.2 Fluidics Protocols for the GeneChip® ST Arrays

Fluidics Station 450 FS450_0001 and FS450_0007	
Post Hyb Wash #1	10 cycles of 2 mixes/cycle with Wash Buffer A at 30°C
Post Hyb Wash #2	6 cycles of 15 mixes/cycle with Wash Buffer B at 50°C
Stain	Stain the probe array for 5 minutes in SAPE solution at 35°C
Post Stain Wash	10 cycles of 4 mixes/cycle with Wash Buffer A at 30°C
2nd Stain	Stain the probe array for 5 minutes in antibody solution at 35°C
3rd Stain	Stain the probe array for 5 minutes in SAPE solution at 35°C
Final Wash	15 cycles of 4 mixes/cycle with Wash Buffer A at 35°C.
Holding Buffer	Fill the probe array with Array Holding Buffer.
<ul style="list-style-type: none"> • Wash Buffer A = non-stringent wash buffer • Wash Buffer B = stringent wash buffer 	

Table 6.3 Fluidics Scripts for GeneChip® ST Array Types

Array Format	Fluidics Script Protocol
49 Format	FS450_0001
64 Format	FS450_0001
169 Format	FS450_0007

Washing and Staining the Probe Array on Fluidics Station 450

1. In the Fluidics Station dialog box on the workstation, select the correct experiment name from the drop-down **Experiment** list.
The Probe Array Type appears automatically.
2. In the **Protocol** drop-down list, select FS450_0001 or FS450_0007 to control the washing and staining steps. Refer to [Table 6.3](#) for information on determining the correct Protocol for your array.
3. Choose **Run** in the Fluidics Station dialog box to begin the washing and staining. Follow the instructions on the LCD window on the fluidics station.
If you are unfamiliar with inserting and removing probe arrays from the fluidics station modules, please refer to the appropriate *Fluidics Station User's Guide or Quick Reference Card* (P/N 08-0093 for the FS-450/250 fluidics station).
4. Insert the appropriate probe array into the designated module of the fluidics station while the cartridge lever is in the down, or **EJECT** position. When finished, verify that the cartridge lever is returned to the up, or **ENGAGE** position.
5. Remove any microcentrifuge vial remaining in the sample holder of the fluidics station module(s) being used.
6. Follow the instructions on the LCD window on the fluidics station by placing the three experiment sample vials (the microcentrifuge vials) into the sample holders 1, 2, and 3 on the fluidics station.
 - A. Place one vial containing 600 µL **Stain Cocktail 1** in sample holder 1.
 - B. Place one vial containing 600 µL **Stain Cocktail 2** in sample holder 2.
 - C. Place one vial containing 800 µL of **Array Holding Buffer** in sample holder 3.
 - D. Press down on the needle lever to snap needles into position and to start the run. The run begins. The Fluidics Station dialog box at the workstation terminal and the LCD window display the status of the washing and staining as the protocol progresses.
7. When the protocol is complete, the LCD window displays the message **EJECT & INSPECT CARTRIDGE**.
8. Remove the probe arrays from the fluidics station modules by first pressing down the cartridge lever to the eject position.
9. Check the probe array window for large bubbles or air pockets.
 - If the probe array has no large bubbles, it is ready to scan on the GeneChip® Scanner 3000 7G. Pull up on the cartridge lever to engage washblock and proceed to [Scanning on page 45](#).
 - If bubbles are present, do the following:
Return the probe array to the probe array holder. Follow instructions on the LCD window. Engage the washblock by gently pushing up on the cartridge lever to the engaged, or closed, position.

The fluidics station will drain the probe array and then fill it with a fresh volume of **Array Holding Buffer**. When it is finished, the LCD window will display **EJECT & INSPECT CARTRIDGE**. Again, remove the probe array and inspect it for bubbles. If no bubbles are present, it is ready to scan. Pull up on the lever to close the washblock and proceed to *Scanning on page 45*.

If attempt to fill the probe array without bubbles is unsuccessful, the array should be filled manually with **Array Holding Buffer** using a micropipette with volumes listed in [Table 6.1](#). Excessive washing will result in a loss of signal intensity.

- 10.** If you do not scan the arrays right away, keep the probe arrays at 4°C and in the dark until ready for scanning.
- 11.** If there are no more arrays to wash, shut down the fluidics station following the procedure outlined in the section, *Shutting Down the Fluidics Station on page 44*.

Shutting Down the Fluidics Station

1. After removing a probe array from the probe array holder, the LCD window displays the message **ENGAGE WASHBLOCK**.
2. Gently lift up the cartridge lever to engage, or close, the washblock.
 - The fluidics station automatically performs a Cleanout procedure. The LCD window indicates the progress of the Cleanout procedure.
3. When the fluidics station LCD window indicates **REMOVE VIALS**, the Cleanout procedure is complete.
4. Remove the sample microcentrifuge vial(s) from the sample holder(s).
5. If no other arrays are to be processed, place wash lines into a bottle filled with deionized water.
6. Choose **Shutdown_450** for all modules from the drop-down **Protocol** list in the Fluidics Station dialog box. Click the **Run** button for all modules.

The Shutdown protocol is critical to instrument reliability. Refer to the appropriate *Fluidics Station User's Guide* for more information.
7. After Shutdown protocol is complete, flip the ON/OFF switch of the fluidics station to the OFF position.



IMPORTANT: To maintain the cleanliness of the fluidics station and obtain the highest quality image and data possible, the bleach protocol is highly recommended. Please refer to the *GeneChip® Fluidics Station 450/250 User's Guide* (P/N 08-0092) available at www.affymetrix.com.

Scanning

The GeneChip® Scanner 3000 7G is also controlled by GeneChip® Operating Software (GCOS). The probe array is scanned after the wash protocols are complete. Make sure the laser is warmed up prior to scanning by turning it on at least 10 minutes prior to use. If probe array was stored at 4°C, warm to room temperature before scanning. Refer to the GCOS online help and the appropriate scanner user's manual for more information on scanning.

▲ WARNING: The scanner uses a laser and is equipped with a safety interlock system. Defeating the interlock system may result in exposure to hazardous laser light.

You must have read, and be familiar with, the operation of the scanner before attempting to scan a probe array. Please refer to the GeneChip® Scanner 3000 7G quick reference card or user's manual.

Handling the Probe Array

Before you scan the probe array, follow the directions in this section on handling the probe array. If necessary, clean the glass surface of the probe array with a non-abrasive towel or tissue before scanning. Do not use alcohol to clean glass.

Before scanning the probe array cartridge, apply Tough-Spots to each of the two septa on the probe array cartridge to prevent the leaking of fluids from the cartridge during scanning.

! IMPORTANT: Apply the spots just before scanning.

1. On the back of the probe array cartridge, clean excess fluid from around septa.

- Carefully apply one Tough-Spots to each of the two septa. Press to ensure that the spots remain flat. If the Tough-Spots do not apply smoothly, that is, if you observe bumps, bubbles, tears, or curled edges, do not attempt to smooth out the spot. Remove the spot and apply a new spot. See [Figure 7.1](#).

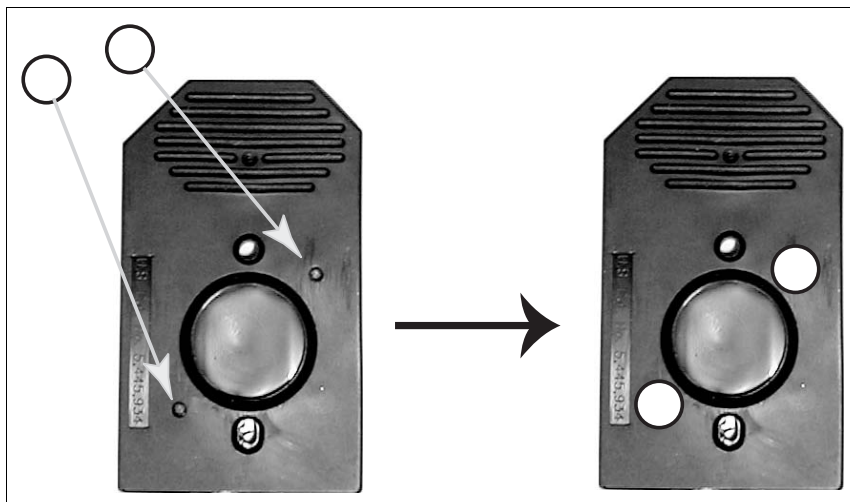


Figure 7.1 Applying Tough-Spots® to the probe array cartridge

- Insert the cartridge into the scanner and test the autofocus to ensure that the Tough-Spots do not interfere with the focus. If you observe a focus error message, remove the spot and apply a new spot. Ensure that the spots lie flat.

Scanning the Probe Array

1. Select **Run** → **Scanner** from the menu bar. Alternatively, click the **Start Scan** icon in the tool bar.
 - The Scanner dialog box appears with a drop-down list of experiments that have not been run.
2. Select the experiment name that corresponds to the probe array to be scanned. A previously run experiment can also be selected by using the **Include Scanned Experiments** option box. After selecting this option, previously scanned experiments appear in the drop-down list.
3. Once the experiment has been selected, click the **Start** button.
 - A dialog box prompts you to load an array into the scanner.
4. Open the sample door on the scanner and insert the probe array into the holder. Do not force the probe array into the holder. Close the sample door of the scanner.
5. Click **OK** in the Start Scanner dialog box.
 - The scanner begins scanning the probe array and acquiring data. When **Scan in Progress** is selected from the **View** menu, the probe array image appears on the screen as the scan progresses.

WT Sense Target Labeling Assay

1. What is the basic principle of the ribosomal RNA reduction procedure using the RiboMinus Human/Mouse Transcriptome Isolation Kit?

Four biotinylated LNA RiboMinus probes are designed to specifically bind to the abundant 18S and 28S rRNA species (2 probes each for 18S and 28S rRNA). Following hybridization of the biotinylated probes to the rRNA molecules in the total RNA sample, the rRNA is efficiently removed from the sample by the addition of the RiboMinus Magnetic Beads that are coated with streptavidin. The unbound fraction represents the RNA with rRNA species reduced. The sample is then concentrated before target labeling using the IVT cRNA Cleanup Kit. Consult the handbook included in the RiboMinus Kit from Invitrogen for more details.

2. Why is Betaine added to the RiboMinus Hybridization Buffer?

Betaine increases the hybridization stringency. It equalizes the GC T_m and AT T_m , so the background, non-specific, non-rRNA hybridization to the RiboMinus probes due to high GC content may be reduced.

3. Why do you choose to use an rRNA-reduction strategy but not a poly-A mRNA-specific selection protocol?

It has been shown that a portion of the transcripts in total RNA do not necessarily contain Poly-A tails; therefore, they will be excluded by a poly-A RNA positive selection technique. The rRNA reduction approach will also make the protocol more robust in handling smaller amounts of total RNA samples as little as 1 μ g.

4. What is the recommendation on how the total RNA samples should be prepared for this assay?

A standard preparation method should be used, as recommended for the current GeneChip® Human Genome U133 Arrays. The quality assessment metrics, including the Bioanalyzer trace and O.D. ratios, should remain unchanged.

5. Does genomic DNA contamination in the sample interfere with the results, and how do I monitor the degree of its effect?

By titrating genomic DNA back into the total RNA samples and monitoring the deterioration of the array data, it was determined during development of the assay that a moderate amount of genomic DNA contamination will only have minimum effect on the array results. Therefore, routine RNA isolation techniques coupled with DNase treatment should yield sufficiently high-quality sample for analysis on the GeneChip ST arrays.

6. What starting material is needed for the assay?

One μg of total RNA per sample is the recommended starting quantity if following the standard protocol with an up-front rRNA removal procedure. Less total RNA (100 to 300 ng) can be used with an alternative protocol without the rRNA reduction process, however, it is anticipated that the array detection sensitivity and specificity will be compromised at the exon level. Follow recommendations on the optimal assay for each array, as described in [Chapter 1](#).

7. Can I use more than 1 μg (or 100 ng) of total RNA?

While, 1 μg of total RNA is the recommended amount when employing the 1 μg Total RNA Labeling Protocol a range of 1 to 2 μg can be used. Higher amount of input total RNA typically result in modest increases in cRNA yield. However, when using 2 μg of total RNA it is highly recommended to scale-up the RiboMinus reagents appropriately when carrying out the rRNA reduction step. When employing the 100 ng Total RNA Labeling Protocol, 100 to 300 ng of total RNA may be used as input.

8. What is the typical cRNA yield after the IVT reaction in the first cycle?

Starting with 1 μg of total RNA, following the standard protocol, $\geq 8 \mu\text{g}$ of cRNA is routinely obtained; cRNA yields will depend mainly on the cell line or tissue used.

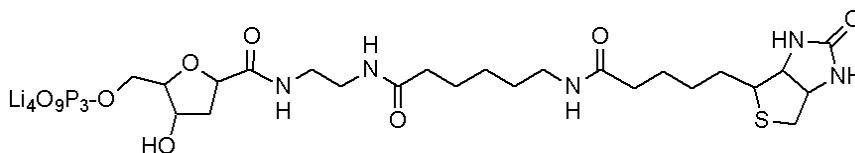
Starting with 100 ng of total RNA without the RiboMinus rRNA removal step, typically, 15 to 40 μg of cRNA can be generated after cRNA cleanup.

9. What is the basic principle of the single-stranded DNA fragmentation and labeling procedure?

Using cRNA generated from the IVT reaction at the end of the first cycle of the assay as a template, single-stranded DNA is synthesized using random primers and the dUTP + dNTP mix. The resulting single-stranded DNA (ss-DNA) containing the unnatural uracil base is then treated with Uracil DNA Glycosylase, which specifically removes the uracil residue from the ss-DNA molecules. In the same reaction, the APE 1 enzyme then cleaves the phosphodiester backbone where the base is missing, leaving a 3'-hydroxyl and a 5'-deoxyribose phosphate terminus.

10. What is the basic component in the DNA Labeling Reagent?

The key labeling molecule in the DNA Labeling Reagent is Biotin Allonamide Triphosphate. See the structure below:



11. What is the expected length of the fragmented DNA target?

On a Bioanalyzer, the fragmented single-stranded DNA target should have a peak centered around 40 to 70 bases with the majority of the fragments ranging from 20 bases to 200 bases.

12. Are there any safe stopping points in the assay? What are recommended storage conditions?

There are a few safe stopping points in the assay, including:

- After RNA Clean-up/Concentration following the RiboMinus rRNA reduction step, before proceeding to first cycle, first strand cDNA synthesis [Store rRNA-reduced total RNA at -80°C].
- After IVT reaction and the cRNA cleanup step in the first cycle, before proceeding to the second cycle of reverse transcription [Store cRNA at -80°C].
- After reverse transcription and the single-stranded cDNA cleanup step in the second cycle, before fragmentation and labeling [Store single stranded cDNA at -20°C].
- After fragmentation and labeling, before hybridization [Store labeled cDNA at -20°C].

13. How much single-stranded DNA target do you need to hybridize to one array?

It is recommended to hybridize approximately 5 µg or 2 µg of fragmented and labeled DNA target to each Exon or Gene Arrays respectively.

14. What is the hybridization condition?

As described in the *GeneChip® Whole Transcript Sense Target Labeling Assay Manual*, a final concentration of 7% DMSO is included in the hybridization cocktail for hybridizing the WT sense target to ST arrays.

15. Can I hybridize the DNA target to the HG-U133 arrays?

The WT Sense Target Labeling Assay is optimized to produce targets specifically for hybridization to ST array type of design. The target is in the sense orientation and the GeneChip® Human Genome U133 Plus 2.0 Array is designed to be compatible with anti-sense targets. Therefore, it is not recommended to mix and match the assays and the array types.

16. Can I use this protocol for prokaryotic arrays?

This has not been tested at the moment; therefore, it is not recommended to use the protocol for any application other than on ST arrays.

17. How does this protocol perform on partially degraded samples?

Utilizing the WT assay for partially degraded samples may be an attractive strategy for profiling these samples. However, it has not been tested thus far in development; therefore, it is recommended that only high-quality total RNA samples should be used.

Array Hybridization, Washing, Staining, and Scanning

18. Why is there no pre-hybridization step for the arrays using the targets from the WT Assay? The pre-hybridization step was required for the 3' target in the *GeneChip® Expression Analysis Technical Manual*.

No pre-hybridization step is necessary for the WT targets. There are many differences between the WT targets and the 3' targets in terms of the nature of the molecules (DNA vs. RNA), as well as labeling molecule and hybridization cocktail makeup. It has been found that pre-hybridization is not necessary for the WT targets.

19. What Fluidics Protocol do I use for the GeneChip® ST Arrays?

New Fluidics Protocols have been developed for this assay, FS450_0001 for Exon Arrays and FS450_0007 for Gene Arrays. In addition to tubes containing SAPE and anti-streptavidin biotinylated antibody, there is a tube containing 1X Array Holding Buffer, which is added to the cartridge following the wash/stain procedure. Please refer to the *GeneChip® Whole Transcript Sense Target Labeling Assay Manual*, Chapter 5, for more details.

20. How long does it take to scan an array?

It takes approximately 35 minutes to scan each Exon Array and approximately 10 minutes to scan each Gene Array.

21. What are the internal grid lines on the array image for?

There is a new gridding algorithm in GCOS 1.3, specifically designed for image analysis of arrays with 5 μm or smaller feature size. This algorithm utilizes 169 subgrids (13x13) to address probe cells on the array. This method is superior to global gridding for arrays with feature pitch less than 11 μm .

22. How do I check for the correct gridding of the new array?

If any of the subgrids fail to align, GCOS will fail to generate a .cel file. The failed subgrids will be visualized with red borders and an “x” in the center, rather than white borders for successfully aligned subgrids.

Gel-Shift Assay

The efficiency of the labeling procedure can be assessed using the following procedure. This quality control protocol prevents hybridizing poorly labeled target onto the probe array. The addition of biotin residues is monitored in a gel-shift assay, where the fragments are incubated with avidin prior to electrophoresis. The nucleic acids are then detected by staining, as shown in the gel photograph [Figure B.1](#). The procedure takes approximately 90 minutes to complete.



NOTE: The absence of a shift pattern indicates poor biotin labeling. The problem should be addressed before proceeding to the hybridization step.

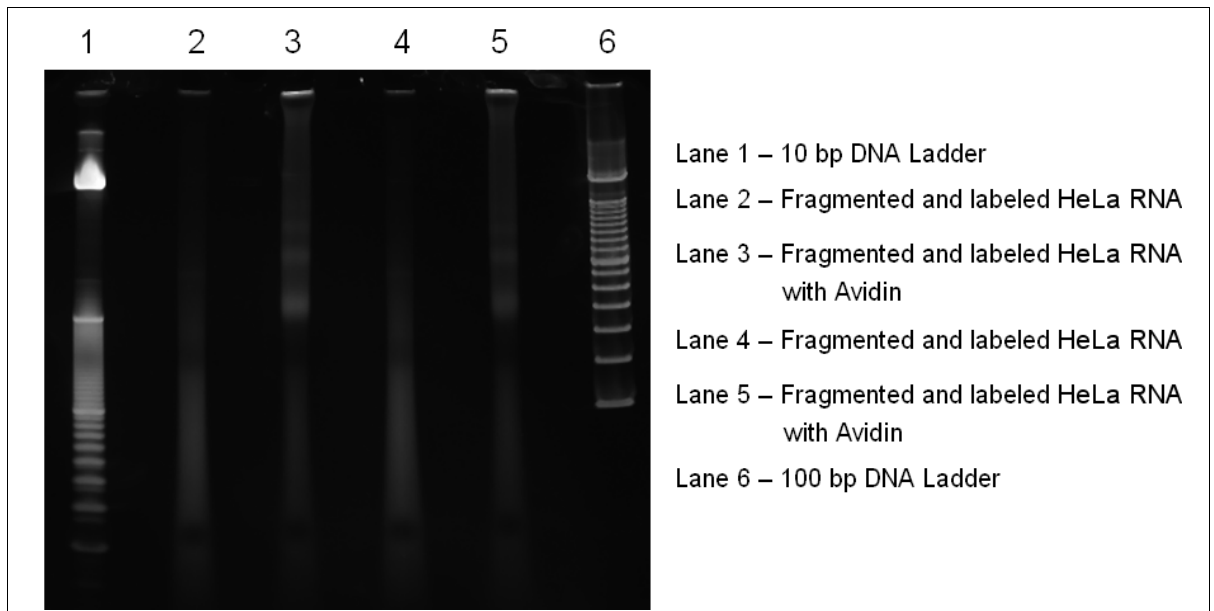


Figure B.1 Gel-Shift

1. Prepare a NeutrAvidin solution of 2 mg/mL in PBS.
2. Place a 4% to 20% TBE gel into the gel holder and load system with 1X TBE Buffer.
3. For each sample to be tested, remove two 1 μ L aliquots of fragmented and biotinylated sample to fresh tubes. Heat the aliquots of samples at 70°C 2 minutes.
4. Add 5 μ L of 2 mg/mL NeutrAvidin to one of the two tubes for each sample tested.
5. Mix and incubate at room temperature for 5 minutes.
6. Add loading dye to all samples to a final concentration of 1X loading dye.
7. Prepare 10 bp and 100 bp DNA ladders
(1 μ L ladder + 7 μ L water + 2 μ L loading dye for each lane).
8. Carefully load samples and two ladders on gel. Each well can hold a maximum of 20 μ L.
9. Run the gel at 150 volts until the front dye (red) almost reaches the bottom. The electrophoresis takes approximately 1 hour.
10. While the gel is running, prepare at least 100 mL of a 1X solution of SYBR Gold for staining.



NOTE: SYBR Gold is light sensitive. Therefore, use caution and shield the staining solution from light. Prepare a new batch of stain at least once a week.

11. After the gel is complete, break open cartridge and stain the gel in 1X SYBR Gold for 10 minutes.
12. Place the gel on the UV light box and produce an image following standard procedure. Be sure to use the appropriate filter for SYBR Gold.