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Dual-Labeled Expression-Tiling Microarray Protocol for Empirical Annotation of Genome Sequences

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Introduction

The utilization of microarrays for empirical annotation of a genome and differential gene expression analysis focuses on protein-encoding RNA of Prokaryote and Eukaryote systems. Currently, our workflow measures gene expression, while simultaneously providing empirical evidence for gene annotation, from test and reference samples of Eukaryote systems by dual-color microarray hybridization. The workflow is designed to detect steady-state gene expression levels from limited samples. Briefly, RNA is obtained from the biological source of interest and processed to stable labeled targets for microarray hybridization, from which a high-resolution image is recorded and signal intensities are extracted and documented for analysis. First, Total RNA is extracted from the biological source with the use of organic solvents and purified with a silica-based centrifugation column, coupled with on-column Dnase I treatment. Next, T7-based RNA amplification technique allows for linear amplification of polyA+ RNA from the limited sample. Then, double-stranded cDNA amplification technique converts the amplified RNA into a suitable intermediate for random primer labeling. After that, NimbleGen Systems Hybridization technique allows for uniform hybridization of the labeled targets to produce high interarray correlations between replicate hybridizations. Subsequently, array scanning excites the fluorescent dyes to record a high-resolution image file, from which NimbleScan Software extracts quantifiable data for analysis. At the completion of the procedure, data is presented in a format suitable for normalization and comparative analysis.

Precautions

Laboratory safety. It is assumed that users have a sound knowledge of molecular biology techniques and safe laboratory practices. Before undertaking a new protocol or using unfamiliar reagents, users should review relevant Material Safety Data Sheets to identify potential hazards and recommended precautions. For background in general molecular biology please see *Molecular Cloning A Laboratory Manual*, J. Sambrook and D. W. Russell. Cold Spring Harbor Laboratory Press.

Materials

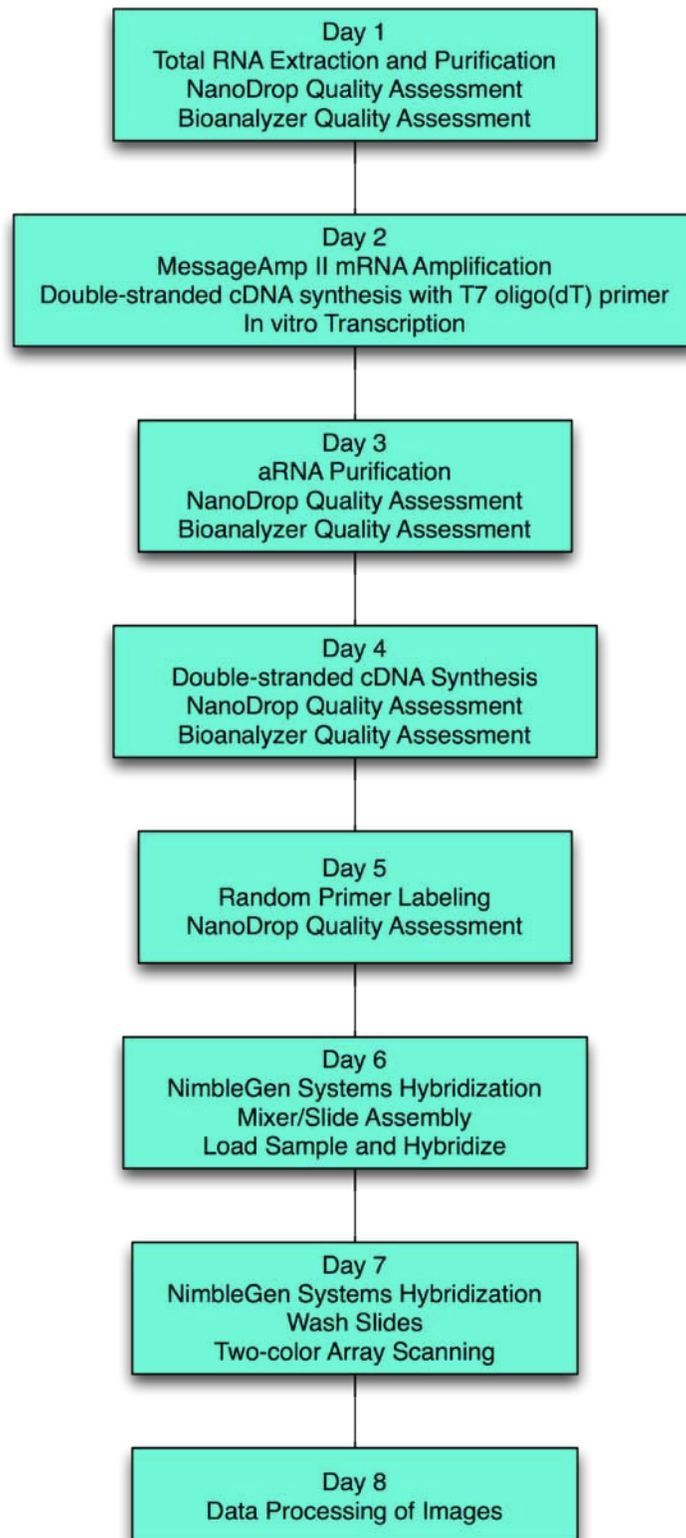
Application	Item Name	Company	Catalog Number	Unit Size
RNA Isolation	Chloroform	EMD Chemicals	CX1055-6	500 ml
RNA Isolation	TRIzol Reagent	Invitrogen	15596-026	100 ml
RNA Isolation	Rnase-free Dnase set (50)	Qiagen	79254	50 rxn
RNA Isolation	RNeasy Mini Kit	Qiagen	74104	50 rxn
RNA Isolation	Disposable pestles; 1.5mL, Plastic	VWR	KT749521-1590	100 Units
RNA amplification	MessageAmp II aRNA Amplification Kit (Part one & two)	Ambion	AM1751	20 rxn
Double-stranded cDNA Synthesis	100mM dNTP set, 4x25µmol	Invitrogen	10297-018	1000 µl
Double-stranded cDNA Synthesis	5X Second Strand Buffer	Invitrogen	10812-014	500 µl
Double-stranded cDNA Synthesis	ChargeSwitch PCR Clean-Up Kit	Invitrogen	CS12000	100 rxn
Double-stranded cDNA Synthesis	DNA Polymerase I (10U/µl)	Invitrogen	18010-025	1000 Units
Double-stranded cDNA Synthesis	E. coli DNA Ligase (10U/µl)	Invitrogen	18052-019	100 Units
Double-stranded cDNA Synthesis	Rnase H (2U/µl – 5U/µl)	Invitrogen	18021-071	120 Units
Double-stranded cDNA Synthesis	RNASEOUT (40U/µl)	Invitrogen	10777-019	5000 Units
Double-stranded cDNA Synthesis	RT-PCR Grade Water	Invitrogen	AM9935	1.5 ml x 10
Double-stranded cDNA Synthesis	SuperScript II (200U/µl)	Invitrogen	18064-014	10000 Units
Double-stranded cDNA Synthesis	T4 DNA Polymerase (5U/µl)	Invitrogen	18005-025	250 Units
Double-stranded cDNA Synthesis	Random Primer	Promega	C1181	20 µg
Double-stranded cDNA Synthesis	Rnase A Solution, 4 mg/ml	Promega	A7973	1000 µl
Double-stranded cDNA Synthesis	0.5M EDTA, pH 8.0	Sigma-Aldrich	E7889-100mL	100 ml
Double-stranded cDNA Synthesis	7.5M Ammonium Acetate	Sigma-Aldrich	A2706-1L	1 liter
Double-stranded cDNA Synthesis	Ethanol, Absolute, 200 proof	Sigma-Aldrich	E7023-500mL	500 ml
DNA labeling	NimbleGen Dual-Color Labeling Kit	Roche NimbleGen	05223547001	20 rxn
DNA labeling	β-Mercaptoethanol	Sigma-Aldrich	M3148-25mL	25 ml
DNA labeling	Isopropanol	Sigma-Aldrich	I9516-500mL	500 ml
Hybridization	NimbleGen Hybridization Kit, LS	Roche NimbleGen	05583934001	40 slides
Hybridization	NimbleGen Wash Buffer Kit	Roche NimbleGen	05584507001	10 uses
Hybridization	NimbleGen Array Processing Accessories	Roche NimbleGen	05223539001	unlimited
Hybridization	Microman Capillary Pistons, CP100, nonsterile	Gilson	F148412	192 tips

Hybridization	Water, Reagent grade, ACS, nonsterile, Type I	VWR	RC91505	20 L
Hybridization	Kimwipe	Cole-Parmer	EW-33670-04	280 x 60 boxes
Hybridization	Forceps	Multiple vendors	---	---
QA	Agilent RNA 6000 Nano Kit	Agilent	5067-1511	25 chips
Supplies	Rnase Zap	Ambion	AM9780	250 mL
Supplies	UltraPure Dnase/Rnase-Free Distilled water	Invitrogen	10977-015	500 mL
Supplies	PCR strip w/ cap (8 tubes/strip)	VWR	20170-004	125 strips
Supplies	Premium Research Microcentrifuge tubes (1.5 ml)	MIDSCI	AVSS1700	1,000 Units
Supplies	ART 10	Molecular BioProducts	2139	960 Tips
Supplies	ART 20P	Molecular BioProducts	2149P	960 Tips
Supplies	ART 200	Molecular BioProducts	2069	960 Tips
Supplies	ART 1000E	Molecular BioProducts	7029E	800 Tips

Equipment

Application	Item Name	Company	Catalog Number	Capacity / Version
Sample Processing	Mastercycler® gradient	Eppendorf	5331 000.010	0.2ml x 96
Sample Processing	Microcentrifuge 5424	Eppendorf	22620401	1.5ml x 24
Sample Processing	Microcentrifuge 5415 R	Eppendorf	5425 725.000	1.5ml x 24
Sample Processing	Vacuum Centrifuge	Multiple vendors	---	1.5ml x 24
Sample Processing	Microcentrifuge	Multiple vendors	---	1.5ml x 8& PCR strip
Sample Processing	Heat block	Multiple vendors	---	1.5ml x 24
Sample Processing	Water bath	Multiple vendors	---	20 L
Sample Processing	MagnaRack Magnetic Rack	Invitrogen	CS15000	1.5ml x 12
Hybridization	Microman M100 Pipette	Glison	F148504	---
Hybridization	Hybridization System 4 (110V)	Roche NimbleGen	5223652001	4 slides
Hybridization	Precision Mixer Alignment Tool (PMAT)	Roche NimbleGen	Included with hyb system	1 slide
Hybridization	Disassembly tool	Roche NimbleGen	Included with hyb systems	1 slide
Hybridization	NimbleGen MS 200 Microarray Scanner	Roche NimbleGen	05 394 341 001	48 slides
Hybridization	Microarray High Speed Centrifuge	ArrayIT	MHC110V	1 Slide
QA	NanoDrop 1000	ThermoScientific	SID-10135606	1 sample
QA	Bioanalyzer 2100	Agilent	G2938C	1 chip
QA	Chip Priming Station	Agilent	5065-4401	1 chip
QA	IKA Vortex Chip Adapter	Agilent	5065-9966	1 chip
Software	NimbleGen MS 200 Data Collection Software	Roche NimbleGen	---	1.1
Software	NimbleScan	Roche NimbleGen	05 933 315 001-	2.6
Software	NanoDrop 1000 Software	ThermoScientific	---	2.5
Software	Bioanalyzer Software	Agilent	G2941AA	B.02.05

Workflow



Thermocycler Programs

MessageAmp II aRNA Amplification

- 1) 70°C for 10 minutes, lid temperature 105°C
- 2) 42°C for 60 minutes x 2, lid temperature 50°C
- 3) 16°C for 60 minutes x 2, heated-lid off
- 4) 37°C for 60 minutes x 6, lid temperature 105°C; 4°C Hold

Double-stranded cDNA Synthesis

- 1) 70°C for 10 minutes, lid temperature 105°C
- 2) 42°C for 60 minutes, lid temperature 50°C
- 3) 16°C for 60 minutes x 2, heated-lid off, then "Pause", followed by 16°C for 5 minutes, heated-lid off, and finally 4°C Hold
- 4) 37°C for 10 minutes, lid temperature 105°C

Random Primer Labeling

- 1) 98°C for 10 minutes, lid temperature 105°C
- 2) 37°C for 60 minutes x 2, lid temperature 105°C

Quality Assessment (QA)

- 1) 70°C for 2 minutes, lid temperature 105°C

References

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Acknowledgements

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Portions of the microarray processing and analysis protocols provided in this document are courtesy of Roche NimbleGen, Inc. NimbleGen microarrays are intended for life science research only and are not for use in diagnostic procedures. The protocol was developed by the authors of this document and it is not standard for processing NimbleGen Catalog Gene Expression microarrays.

We thank Tsetska Takova (Director, Global Marketing, Arrays and Reagents at Roche NimbleGen, Inc.) and Kary Staples (Manager, Global Marketing Communications, at Roche NimbleGen, Inc) for help in creating this training manual, and Jennifer Steinbachs (CGB) for help with formatting.

Total RNA Extraction and Purification

Introduction

The utilization of microarrays for gene expression analysis requires the extraction of RNA from a biological source of interest, either from a Prokaryote or Eukaryote source. We describe this procedure in detail along with RNA quality assessment measures. Briefly, Total RNA is extracted from the biological source with the use of organic solvents. The extracted Total RNA is purified with a silica-based centrifugation column. The addition of on-column based Dnase I digestion facilitates the removal of genomic DNA contamination. At the completion of the procedure, use of a spectrophotometer and microfluidic capillary electrophoresis assesses the quality of the purified Total RNA.

Precautions

Laboratory safety. It is assumed that users have a sound knowledge of molecular biology techniques and safe laboratory practices. Before undertaking a new protocol or using unfamiliar reagents users should review relevant Material Safety Data Sheets to identify potential hazards and recommended precautions. For background in general molecular biology please see *Molecular Cloning A Laboratory Manual*, J. Sambrook and D. W. Russell. Cold Spring Harbor Laboratory Press.

Caution. TRIzol Reagent, a phenol-based solution, is toxic when in contact with skin or if swallowed. After contact with skin, wash the area immediately with plenty of detergent and water. This solution should be handled with caution.

Prevent excessive Rnase exposure. As with all steps involved in RNA isolation, it is essential to avoid latex glove. Use only powder-free nitrile gloves. It is likewise important to guard against sources of dust and nucleases. We recommend using Dnase- and Rnase-free plastics, including barrier pipette tips. In addition, decontaminate both workspaces and pipettes with RNase Zap, according to manufacturer's instructions.

Instrument Setup:

1. Chill a refrigerated centrifuge to 4°C.
2. Equilibrate TRIzol Reagent to room temperature for at least 1-hour prior to use.

Sample Preparation Setup:

1. For each sample,
 - 1.1. Labeled 1 x 1.5ml Rnase-free microcentrifuge tube
 - 1.2. Labeled 1 x 1.5ml RNeasy collection tube
 - 1.3. Labeled 1 x RNeasy mini-spin column, plus an additional 2ml collection tube
 - 1.4. Rnase-free Dnase Kit: Add 35µl of RDD Buffer to 5µl of Dnase I stock solution. Mix by gently pipetting. Store at 4°C for up to one day.
2. Rnase-free Dnase Kit: Reconstitute lyophilized Dnase I as directed by manufacturer.
3. RNeasy Mini Kit: Add Ethanol, 100% Soln, to Buffer RPE as directed by manufacturer.
4. All centrifugation steps are at room temperature, unless otherwise noted.
5. Remove the frozen tissue samples from -80°C storage. Keep on dry ice until ready to perform extraction procedure.

Procedure

1. Add 500µl of TRIzol Reagent directly to the tube containing the frozen tissue sample. Immediately homogenize tissue. Grind tissue using disposable blue pestle. After tissue has been completely homogenized, add another 500µl of TRIzol Reagent to the tube containing the homogenized tissue. Incubate mixture at least 5 minutes at room temperature to ensure complete dissociation of nucleoprotein complexes. Repeat step 1 for the remaining frozen tissue samples to be processed. Continue to step 2 after all tissue samples have been homogenized.
2. Add 200µl of Chloroform to each sample. Vigorously shake the mixture for 15 seconds. Do not vortex.
3. Centrifuge the mixture for 15 minutes at 4°C at 11,600 rcf (< 12,000 x g).
4. Transfer upper, aqueous phase (approx. 400µl) to labeled Rnase-free 1.5ml Rnase-free microcentrifuge tube (supplied by user).
5. Precipitate RNA by adding 0.5 volume (approx. 200µl) of Absolute Ethanol, 100% Soln.
6. Gently pipette 4 times to mix, and flick tube 3 times. Transfer precipitated RNA from the previous step to the labeled RNeasy mini-spin column. Proceed immediately. Degradation and acidification of RNA can occur with prolonged expose at this point.

Maximum loading volume	700µl
Maximum binding capacity	100µg

7. Centrifuge the column for 30 seconds at 10,000 rcf. Discard flow through. Replace column into same 2mL collection tube.
8. On-Column Dnase Treatment: Wash the column with 350µl of Buffer RW1. Centrifuge the column for 30 seconds at 10,000 rcf. Discard flow through. Replace column into same 2mL collection tube. Pipette the Dnase I/RDD Buffer mixture (40µl) directly onto the RNeasy silica-gel membrane of the RNeasy column. Incubate for 10 minutes at room temperature. Wash the column with another 350µl of Buffer RW1. Centrifuge the column for 30 seconds at 10,000 rcf. Discard flow through. Replace column into same 2mL collection tube.
9. Wash the column with 500µl of Buffer RPE. Centrifuge the column for 30 seconds at 10,000 rcf. Discard flow through. Replace column into same 2mL collection tube.
10. Wash the column with another 500µl of Buffer RPE. Centrifuge the column for 30 seconds at 10,000 rcf. Discard flow through. Place column in a NEW 2mL collection tube.
11. Centrifuge for 2 minutes at 10,000 rcf. Transfer the column to the labeled 1.5ml collection tube (supplied in Rneasy kit).
12. To elute, pipette 30µl of Rnase-free water directly onto silica-gel membrane of the column. Incubate for 1 minute at room temperature.
13. Centrifuge the column for 1 minute at 10,000 rcf.
14. The elutate now contains purified Total RNA.

STOP POINT: Samples may be stored at -80°C until quality assessment is completed.

15. Determine concentration with NanoDrop. See section *NanoDrop Protocol for NimbleGen Microarray*.
16. Determine integrity using BioAnalyzer / RNA 6000 Nano kit. See section *Bioanalyzer RNA Protocol for NimbleGen Microarray*.

STOP POINT: Samples may be stored at -80°C until ready for processing.

MessageAmp II aRNA Amplification

Introduction

The utilization of microarrays for transcription analysis requires large amounts of RNA from a biological source of interest. T7-based RNA Amplification technique, originally developed in the laboratory of James Eberwine (Russell N. Van Gelder 1990), allows for linear amplification of polyA+RNA from limited samples to produce enough material for microarray hybridizations. Procedures have been adapted to be compatible with Prokaryote or Eukaryote systems. The protocol described here was adapted from Ambion's MessageAmp II Amplification kit for amplification of polyA+RNA from Eukaryote systems. We describe this procedure in detail along with RNA quality assessment measures. Briefly, Total RNA is reverse transcribed with a T7 Oligo dT primer to synthesis first-strand cDNA. T7 promoter site is completed by second strand cDNA synthesis. The RNA Polymerase driven in vitro transcription converts the double-stranded cDNA with T7 RNA Polymerase promoter site into the final product, amplified-antisense RNA (aRNA). At the completion of the procedure, use of a spectrophotometer and microfluidic capillary electrophoresis assesses the quality of the purified aRNA.

Precautions

Laboratory safety. It is assumed that users have a sound knowledge of molecular biology techniques and safe laboratory practices. Before undertaking a new protocol or using unfamiliar reagents users should review relevant Material Safety Data Sheets to identify potential hazards and recommended precautions. For background in general molecular biology please see Molecular Cloning A Laboratory Manual, J. Sambrook and D. W. Russell. Cold Spring Harbor Laboratory Press.

Caution. Ambion's aRNA binding Buffer, containing beta-mercaptoethanol, is toxic when in contact with skin or if swallowed. After contact with skin, wash the area immediately with copious amounts of water. This solution should be handled with caution.

Prevent excessive Rnase exposure. As per all steps involved in RNA amplification, it is essential to avoid latex glove. Use only powder-free nitrile gloves. It is likewise important to guard against sources of dust and nucleases. We recommend using Dnase- and Rnase-free plastics, including barrier pipette tips. In addition, decontaminate both workspaces and pipettes with RNase Zap, according to manufacturer's instructions.

Instrument Setup:

Reverse Transcription to Synthesize First Strand cDNA

Instrument Setup

1. Program MasterCycler/Thermocycler
 - 1.1. 70°C for 10 minutes, lid temperature 105°C
 - 1.2. 42°C for 60 minutes x 2, lid temperature 50°C

Sample Preparation Setup

1. Thaw the following reagents from MessageAmp II aRNA Amplification Kit (part two):
 - 1.1. ON ICE: T7 Oligo dT Primer, dNTP mix
 - 1.2. ROOM TEMP: 10X First Strand Buffer
2. Prepare
 - 2.1. Labeled 1 x 1.5ml microcentrifuge tube for First Strand Master Mix
 - 2.2. Labeled 0.2ml thin-walled PCR tube for each sample
3. All centrifugation steps are at room temperature, unless otherwise noted.
4. Remove the Total RNA samples from -80°C storage. Keep on dry ice until ready to proceed with RNA amplification.

Procedure

1. Prepare first strand RNA/primer mixture in a 0.2ml PCR tube.

Component	Amount
Total RNA	0.25µg – 1.5µg
Primer	1.0µl
Nuclease-Free water	To final volume
Total Volume	12.0µl

2. Incubate RNA/Primer mixture for 10 minutes at 70°C. Immediately, remove the RNA/primer mixture from the thermocycler and chill for 2 minutes in an ice-bath. Centrifuge briefly. Keep on ice.

MasterCycler	70°C 10min
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3. At room temperature, prepare first strand master mix in a 1.5ml microcentrifuge tube labeled First Strand Master Mix.

Component provided in kit part 2	Amount plus 0.05% for error
10X First Strand Buffer	2.10µl
dNTP Mix	4.20µl
RNase Inhibitor	1.05µl
ArrayScript	1.05µl
Total Volume	8.40µl

4. Gently pipette the First Strand Master Mix 4 times and flick 3 times to mix. Centrifuge briefly. Keep at room temperature.
5. Transfer RNA/primer mixture to room temperature. Add 8µl first strand master mix to each tube (20µl total volume). Gently pipette the reaction 4 times and flick 3 times to mix. Centrifuge briefly.
6. Incubate for 2 hours at 42°C, lid temperature 50°C.

MasterCycler	42°C, 2hr
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7. Remove the reaction from the thermocycler. Centrifuge briefly. Place reaction on ice. Immediately proceed to second strand cDNA synthesis.

Second Strand cDNA Synthesis

Instrument Setup

1. Program MasterCycler/Thermocycler
 - 1.1. 16°C for 60 minutes x 2, heated-lid off

Sample Preparation Setup

1. Thaw the following reagents from MessageAmp II aRNA Amplification Kit (part two):
 - 1.1. ON ICE: Nuclease-free water, 10X Second Strand Buffer, dNTP mix
2. Prepare
 - 2.1. Labeled 1x 1.5ml microcentrifuge tube for Second Strand Master Mix
3. All centrifugation steps are at room temperature, unless otherwise noted.

Procedure

1. Prepare second strand master mix on ice in a 1.5ml microcentrifuge tube labeled Second Strand Master Mix.

Component provided in kit part 2	Amount plus 0.05% for error
Nuclease Free water	66.15µl
10X Second Strand Buffer	10.50µl
dNTP Mix	4.20µl
DNA Polymerase	2.10µl
RNase H	1.05µl
Total	84.0µl

2. Gently pipette the Second Strand Master Mix 4 times and flick 3 times to mix. Centrifuge briefly. Keep on ice.
3. Add 80µl second strand master mix to the reaction (100µl total volume). Gently pipette the reaction 4 times and flick 3 times to mix. Centrifuge briefly.
4. Incubate for 2 hours at 16°C, heated lid off.

MasterCycler	16°C, 2hr
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5. Remove the reaction from thermocycler. Centrifuge briefly. Place the reaction on ice.

STOP POINT: Reaction may be stored at -20°C overnight; however, it is better to complete the cDNA purification before stopping.

cDNA Purification

Instrument Setup

1. Heat block, 53°C

Sample Preparation Setup

1. MessageAmpII aRNA Amplification Kit (part one):
 - 1.1. cDNA Binding Buffer
 - 1.2. Wash Buffer (ethanol added, as directed by manufacturer)
 - 1.3. Nuclease-free water
 - 1.4. Labeled 1 x 2ml cDNA filter cartridge with cDNA collection tube per sample
 - 1.5. Labeled 1 x 2ml cDNA collection tube per sample
2. If a precipitate is visible in the cDNA Binding Buffer, warm solution to 37°C for up to 10 minutes. Cool to room temperature before use.
3. Prepare
 - 3.1. Labeled 1 x 1.5ml microcentrifuge tube per reaction for precipitation
4. All centrifugation steps are at room temperature, unless otherwise noted.

Procedure

1. Preheat Nuclease-free water to 53°C for at least 10 minutes.
2. Assemble cDNA Filter Cartridges and 2ml cDNA collection tubes.
3. Transfer cDNA reaction to labeled 1.5ml microcentrifuge tube for precipitation.
4. If necessary, adjust cDNA reaction volume to 100µl with Nuclease-free water.
5. Add 250µl of cDNA Binding buffer to cDNA reaction.
6. Pipette the precipitation mixture 4 times to mix, and transfer to the cDNA Filter Cartridge. Centrifuge for 1 minute at 10,000 rcf. Discard the flow-through. Replace the cDNA Filter Cartridge in the same collection tube.
7. Wash the column with 500µl Wash Buffer. Centrifuge for 1 minute at 10,000 rcf. Discard the flow-through. Replace the cDNA Filter Cartridge in the same collection tube.
8. Centrifuge for 1 minute at 10,000 rcf to remove trace amounts of wash buffer. Transfer cDNA Filter Cartridge to labeled 2ml cDNA collection tube for elution.
9. Add 18µl of Nuclease-free Water (preheated to 53°C) to the center of the filter in the cDNA Filter Cartridge.
10. Incubate for 2 min at 53°C. Centrifuge for 1.5 minutes at 10,000 rcf.
11. The eluate now contains purified double-stranded cDNA with T7 promoter site.

STOP POINT: Samples may be stored at -20°C overnight; however, it is better to proceed to In Vitro Transcription before stopping.

In Vitro Transcription

Instrument Setup

1. Program MasterCycler/Thermocycler

- 1.1. 37°C for 60 minutes x 6, lid temperature 105°C; 4°C Hold

Sample Preparation Setup

1. Thaw the following reagents from MessageAmp II aRNA Amplification Kit (part two):

- 1.1. ON ICE: T7 ATP Soln, T7 CTP Soln, T7 GTP Soln, T7 UTP Soln
- 1.2. ROOM TEMP: 10X Reaction Buffer

2. Prepare

- 2.1. Labeled 1 x 1.5ml microcentrifuge tube for In Vitro Transcription Master Mix
- 2.2. 0.2ml thin-walled PCR tube for each sample

3. All centrifugation steps are at room temperature, unless otherwise noted.

Procedure

1. Transfer the 16µl ds cDNA with T7 promoter site from previous section to 0.2ml PCR tube. Centrifuge briefly. Place on ice.
2. Assemble the in vitro transcription master mix in 1.5ml microcentrifuge tube labeled In Vitro Transcription Master Mix at room temp in the order shown:

Component provided in kit part 2	Amount plus 0.05% for error
T7 ATP Soln (75mM)	4.2µl
T7 CTP Soln (75mM)	4.2µl
T7 GTP Soln (75mM)	4.2µl
T7 UTP Soln (75mM)	4.2µl
T7 10X Reaction Buffer	4.2µl
T7 Enzyme Mix	4.2µl
Total	25.2µl

3. Transfer ds cDNA with T7 promoter site to room temperature. Add 24µl of in vitro transcription master mix to each reaction (40µl total volume). Gently pipette the reaction 4 times and flick 3 times to mix. Centrifuge briefly.

4. Incubate for 6 hours at 37°C, followed by 4°C hold overnight.

MasterCycler	37°C 6hr; 4°C HOLD
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5. Remove from thermocycler. Centrifuge briefly. Add 60µl Nuclease-free water to the reaction (100µl total volume). Flick the reaction 3 times to mix. Centrifuge briefly.

STOP POINT: Reaction may be stored at -80°C overnight until purification; however, it is better to proceed to aRNA purification before stopping.

aRNA Purification

Instrument Setup

1. Heat block, 55°C

Sample Preparation Setup

2. MessageAmpII aRNA Amplification Kit (part one):
 - 2.1. aRNA Binding Buffer
 - 2.2. Wash Buffer (ethanol added, as directed by manufacturer)
 - 2.3. Nuclease-free water
 - 2.4. Labeled 1 x 2ml aRNA filter cartridge with collection tube per sample
 - 2.5. Labeled 1 x 2ml aRNA collection tube per sample
3. Prepare
 - 3.1. Labeled 1 x 1.5ml microcentrifuge tube per reaction for precipitation
 - 3.2. Absolute Ethanol, 100% Soln
4. All centrifugation steps are at room temperature, unless otherwise noted.

Procedure

1. Preheat Nuclease-free water to 55°C.
2. Assemble aRNA Filter Cartridges and aRNA collection tubes.
3. Transfer the in vitro transcription reaction to labeled 1.5ml microcentrifuge tube for precipitation.
4. Add 350µl aRNA Binding Buffer to reaction.
5. Add 250µl Absolute Ethanol, 100% Soln, to reaction. Proceed immediately to the next step.
6. Pipette the precipitation mixture 4 times to mix, and transfer to aRNA Filter Cartridge. Centrifuge for 1 minute at 10,000 rcf. Discard the flow-through. Replace the aRNA Filter Cartridge into the same aRNA Collection tube.
7. Wash column with 650µl Wash Buffer. Centrifuge for 1 minute at 10,000 rcf. Discard the flow-through. Replace the aRNA Filter Cartridge in the same aRNA Collection tube.
8. Centrifuge for 1 minute at 10,000 rcf to remove trace amounts of wash buffer.
9. Transfer aRNA Filter Cartridges to labeled 2ml aRNA Collection tube for elution.
10. Add 100µl Nuclease-Free Water (preheated to 55°C) to the center of the filter of the aRNA Filter Cartridge.
11. Incubate for 2 minutes at 55°C, and then centrifuge for 1.5 minutes at 10,000 rcf.
12. The eluate now contains purified aRNA.
STOP POINT: Samples may be stored at -80°C until quality assessment.
13. Determine concentration with NanoDrop. See section *NanoDrop Protocol for NimbleGen Microarray*.
14. Determine integrity using BioAnalyzer / RNA 6000 Nano kit. See section *Bioanalyzer RNA Protocol for NimbleGen Microarray*.
STOP POINT: Samples may be stored at -80°C until proceeding to next step.

Double-stranded cDNA Synthesis

Introduction

Double-stranded cDNA Synthesis technique allows for amplification of RNA to produce enough material suitable for random primer labeling. The protocol described here was adapted from Invitrogen's SuperScript Double-Stranded cDNA Synthesis Kit for amplification of aRNA derived from Eukaryote systems. We describe this procedure in detail along with RNA quality assessment measures. Briefly, aRNA is primed with random hexamer primer in a reverse transcription reaction to synthesize first strand cDNA. Second strand cDNA synthesis completes the strand to produce double-stranded cDNA. In the presence of T4 DNA Polymerase, the double-stranded cDNA is modified to generate the final product, blunt end double-stranded cDNA (ds cDNA). At the completion of the procedure, use of a spectrophotometer and microfluidic capillary electrophoresis assesses the quality of the purified ds cDNA.

Precautions

Laboratory safety. It is assumed that users have a sound knowledge of molecular biology techniques and safe laboratory practices. Before undertaking a new protocol or using unfamiliar reagents users should review relevant Material Safety Data Sheets to identify potential hazards and recommended precautions. For background in general molecular biology please see *Molecular Cloning A Laboratory Manual*, J. Sambrook and D. W. Russell. Cold Spring Harbor Laboratory Press.

Prevent excessive RNase exposure. As per all steps involved in RNA amplification, it is essential to avoid latex glove. Use only powder-free nitrile gloves. It is likewise important to guard against sources of dust and nucleases. We recommend using Dnase- and RNase-free plastics, including barrier pipette tips. In addition, decontaminate both workspaces and pipettes with RNase Zap, according to manufacturer's instructions.

Reverse Transcription to Synthesize First Strand cDNA

Instrument Setup

1. Program MasterCycler/Thermocycler
 - 1.1. 70°C for 10 minutes, lid temperature 105°C
 - 1.2. 42°C for 60 minutes, lid temperature 50°C

Reagent Setup

1. Prepare 10mM dNTP mix.
 - 1.1. Combine 100µl of each dNTP (dATP, dCTP, dGTP, and dTTP) and bring to a final volume of 1000µl with Nuclease-free water.
 - 1.2. Store at -20°C for up to 6 months.

Sample Preparation Setup

1. Thaw the following reagents from Invitrogen, unless otherwise noted:
 - 1.1. ON ICE: Random Hexamer Primer (Promega), 0.1M DTT Soln, 10mM dNTP mix
 - 1.2. ROOM TEMP: 5X First Strand Buffer
2. Prepare
 - 2.1. 1 x 1.5ml microcentrifuge tube labeled First Strand Master Mix
 - 2.2. 0.2ml thin-walled PCR tube for each sample
3. All centrifugation steps are at room temperature, unless otherwise noted.

- Remove the aRNA samples from -80°C storage. Keep on dry ice until ready to proceed with double-stranded cDNA synthesis.

Procedure

- Prepare aRNA/primer mixture in 0.2mL PCR tube.

Component	Amount
aRNA	10.0µg
Random Primer	1.0µl
Nuclease-Free water	To final volume
Total Volume	11.0µl

- Incubate aRNA/primer mixture for 10 minutes at 70°C. Immediately, remove the RNA/primer mixture from the thermocycler, centrifuge briefly, and chill for 5 minutes in an ice-bath.

MasterCycler	70°C 10min
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- At room temperature, prepare first strand master mix in a 1.5ml microcentrifuge tube labeled First Strand Master Mix.

Component	Amount plus 0.05% for error
5X First Strand Buffer	4.20µl
0.1M DTT Solution	2.10µl
dNTP Mix (10mM)	1.05µl
RNaseOUT (40U/µl)	0.525µl
SuperScript II (200U/µl)	2.10µl
Total Volume	9.975µl

- Transfer aRNA/primer mixture to room temperature. Add 9.5µl First Strand Master Mix to aRNA/primer mixture (20.5µl total volume). Gently pipette the reaction 4 times and flick 3 times to mix. Centrifuge briefly.
- Incubate for 60 minutes at 42°C, lid temperature 50°C.

MasterCycler	42°C, 60min
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- Remove the reaction from the thermocycler. Centrifuge briefly. Place the reaction on ice. Proceed immediately to second strand cDNA synthesis.

Second Strand cDNA Synthesis

Instrument Setup

1. Program MasterCycler/Thermocycler

- 1.1. 16°C for 60 minutes x 2, heated-lid off, then “Pause”, followed by 16°C for 5 minutes, heated-lid off, and finally 4°C Hold

Sample Preparation Setup

1. Thaw the following reagents from Invitrogen, unless otherwise noted:
 - 1.1. ON ICE: Nuclease-free water, 5X Second Strand Buffer, dNTP mix
2. Prepare
 - 2.1. Labeled 1 x 1.5ml microcentrifuge tube for Second Strand Master Mix
3. All centrifugation steps are at room temperature, unless otherwise noted.

Procedure

1. Prepare second strand master mix on ice in a 1.5ml microcentrifuge tube labeled Second Strand Master Mix.

Component	Amount plus 0.05% for error
Nuclease Free water	95.55µl
5X Second Strand Buffer	31.50µl
dNTP Mix (10mM)	3.15µl
E.coli DNA Ligase (10U/µl)	1.05µl
E.coli DNA Polymerase (10U/µl)	4.20µl
E.coli Rnase H (2U/µl)	1.05µl
Total	136.5µl

2. Add 130µl Second Strand Master Mix to reaction (150.5µl total volume). Gently pipette the reaction 4 times and flick 3 times to mix. Centrifuge briefly.
3. Incubate or 2 hours at 16°C, heated lid off.

MasterCycler	16°C, 2hr
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4. At the pause in the cycle, add 2µl E.coli T4 DNA Polymerase to reaction (152.5µl total volume). Gently pipette 4 times to mix.

Component	Amount
E.coli T4 DNA Polymerase (5U/µl)	2.0µl
Total	152.5µl

5. Resume thermocycler program. Incubate for 5 minutes at 16°C, heated lid off.

MasterCycler	16°C, 5min
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6. Remove the reaction from the thermocycler. Centrifuge briefly. Add 10 μ l of 0.5M EDTA, pH 8.0. Flick tube 3 times to mix. Centrifuge briefly.

Component	Amount
0.5M EDTA, pH 8.0	10.0 μ l
Total	162.5 μ l

STOP POINT: Samples may be stored at -20°C overnight, however it is better to complete the cDNA purification before stopping.

Rnase A Treatment and Clean-Up

Instrument Setup

1. Program MasterCycler/Thermocycler
 - 1.1. 37°C for 10 minutes, lid temperature 105°C

Sample Preparation Setup

1. Prepare
 - 1.1. Labeled 2 x 1.5ml microcentrifuge tube per reaction
2. ChargeSwitch PCR Clean-Up Kit
3. MagnaRack Magnetic Rack
4. All centrifugation steps are at room temperature, unless otherwise noted.

Procedure

1. Add 1 μ l of 4mg/ml RNase A Solution to the reaction. Gently pipette the reaction 4 times, and flick 3 times to mix. Centrifuge briefly.
 - 1.1. Important: Use caution when working with RNase A. Use RNase ZAP to clean work area surfaces, if necessary.

Component	Amount
Rnase A (4mg/ml)	1.0 μ l
Total	163.5 μ l

2. Incubate for 10 minutes at 37°C, lid temperature 105°C.

MasterCycler	37°C, 10min
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3. During the 10 minute incubation,
 - 3.1. add 163 μ l ChargeSwitch Purification Buffer (N5) to the first labeled 1.5ml microcentrifuge tube, and then set aside.

Component	Amount
Purification Buffer (N5)	163.0 μ l

4. Remove the reaction from the thermocycler. Centrifuge briefly. Transfer the reaction to the tube containing 163 μ l ChargeSwitch Purification Buffer (N5). Aspirate to mix.

5. Add 20µl ChargeSwitch Magnetic Beads to the Purification Buffer (N5)/reaction mixture. Aspirate to mix without forming bubbles. Incubate at room temperature for 1 minute.
6. Place the sample on the MagnaRack for 1 minute until the beads form a pellet.
7. Without removing the tube from the MagnaRack, carefully aspirate the supernatant without distributing the bead pellet.
8. Remove tube from the MagnaRack. Add 460µl Wash Buffer (W12). Aspirate to resuspend the bead pellet without forming bubbles.
9. Place the sample on the MagnaRack for 1 minute until the beads form a pellet.
10. Without removing the tube from the MagnaRack, carefully aspirate the supernatant without distributing the bead pellet.
11. Repeat Wash Procedure once more: Remove tube from the MagnaRack. Add 460µl Wash Buffer (W12). Aspirate to resuspend the bead pellet without forming bubbles. Place the sample on the MagnaRack for 1 minute until the beads form a pellet. Without removing the tube from the MagnaRack, carefully aspirate the supernatant without distributing the bead pellet.
12. Remove tube from the MagnaRack. Add 20µl Elution Buffer (E5). Aspirate to resuspend the bead pellet without forming bubbles. Incubate at room temperature for 1 minute.
13. Place the sample on the MagnaRack for 1 minute until the beads form a pellet.
14. Without removing the tube from the MagnaRack, carefully transfer the supernatant containing the purified DNA to the second 1.5ml microcentrifuge tube without distributing the bead pellet.

STOP POINT: Samples may be stored at -20°C until quality assessment.

15. Determine concentration with NanoDrop. See section *NanoDrop Protocol for NimbleGen Microarray*.
16. Determine integrity using BioAnalyzer / RNA 6000 Nano kit. See section *Bioanalyzer RNA Protocol for NimbleGen Microarray*.

STOP POINT: Samples may be stored at -20°C until proceeding to next step.

Random Primer Labeling

Introduction

Random Primer Labeling technique allows for amplification of ds DNA to produce enough material for microarray hybridization. Procedures have been adapted to be compatible with genomic DNA and ds cDNA. The protocol described here was adapted from Roche NimbleGen's Dual-Color DNA Labeling Kit for amplification of ds cDNA derived from amplified RNA of Eukaryote systems. We describe this procedure in detail along with DNA quality assessment measures. Briefly, ds cDNA is primed with CY-labeled random nonamer and the addition of Klenow (exo-) enzyme synthesizes 5' Cy-labeled cDNA. At the completion of the procedure, use of a spectrophotometer assesses the quality of the purified 5' Cy-labeled cDNA.

Precautions

Laboratory safety. It is assumed that users have a sound knowledge of molecular biology techniques and safe laboratory practices. Before undertaking a new protocol or using unfamiliar reagents users should review relevant Material Safety Data Sheets to identify potential hazards and recommended precautions. For background in general molecular biology please see *Molecular Cloning A Laboratory Manual*, J. Sambrook and D. W. Russell. Cold Spring Harbor Laboratory Press.

Caution. Random Primer Buffer, containing beta-mercaptoethanol after preparation, is toxic in contact with skin and if swallowed. After contact with skin, wash the area immediately with copious amounts of water. This solution should be handled with caution.

Prevent excessive light exposure. Cy-labeled Random Primer reagent and subsequent Cy-labeled cDNA is light sensitive and should not be exposed to any unnecessary light. When working with the labeled components turn off all lights and close the blinds to the windows.

Prevent excessive ozone and humidity exposure. Minimize ozone exposure and avoid humidity levels above 40% as this leads to oxidation of the cyanine dyes. Use of desiccant materials or a dehumidifier should be considered.

Sample Preparation Setup

1. Thaw the following reagents from Roche NimbleGen Dual Color Labeling Kit, unless otherwise noted:
 - 1.1. ON ICE: Random Primer Buffer
2. Prepare
 - 2.1. 50 x 0.2ml thin-walled PCR tube
3. All centrifugation steps are at room temperature, unless otherwise noted.

Procedure

1. Prepare Random Primer Buffer.

Component	Amount
Random Primer Buffer	998.25 μ l
β -Mercaptoethanol	1.75 μ l
Total Volume	1000.0 μ l

2. Centrifuge briefly Cy3- and Cy5-Random Nonamer. Dilute each Cy-primer with 462 μ l of Random Primer Buffer with β -Mercaptoethanol.

Component	Amount
Cy-Random Nonamer	11 O.D.
Random Primer Buffer with β -Mercaptoethanol	462.0 μ l
Total Volume	462.0 μ l

3. Aliquot 42 μ l individual reaction volumes in 0.2 ml thin-walled PCR tubes. Store at -20°C, protect from light. Yields approximately 11 individual reactions for each dye.

STOP POINT: Diluted Cy-Random Nonamer may be stored for up to 4 months at -20°C until proceeding to next step. Avoid freeze thaw cycles.

Assemble labeling reaction

Instrument Setup

1. Program MasterCycler/Thermocycler
 - 1.1. 98°C for 10 minutes, lid temperature 105°C
 - 1.2. 37°C for 60 minutes x 2, lid temperature 105°C

Sample Preparation Setup

1. Thaw the following reagents from Roche NimbleGen's Dual Color Labeling Kit, unless otherwise noted:
 - 1.1. ON ICE: Diluted Cy-Random Nonamers, 10mM dNTP mix, Nuclease-free water
 - 1.2. Room temperature: Stop Solution and 5M NaCl solution (heat and vortex if precipitate is visible)
2. Prepare
 - 2.1. 0.2ml thin-walled PCR tube for each sample

- 2.2. Labeled 1 x 1.5ml microcentrifuge tube for Klenow Master Mix
 - 2.3. Labeled 1 x 1.5ml microcentrifuge tube per sample
 - 2.4. Labeled 1 x 1.5ml centrifuge tube per hybridization
 - 2.5. Ethanol, 80% Soln
3. All centrifugation steps are at room temperature, unless otherwise noted.

Procedure

15. Prepare the test and reference samples each in triplicate in separate 0.2ml thin-walled PCR tubes. Dye assignment can be switched, if desired.

Component	Test x 3	Reference x 3
ds cDNA	1.0µg	1.0µg
Diluted Cy3-Random Nonamers	42µl	---
Diluted Cy5-Random Nonamers	---	42µl
Nuclease-free water	To volume (80µl)	To volume (80µl)
Total	80.0µl	80.0µl

16. Incubate cDNA/primer mixture for 10 minutes at 98°C.

MasterCycler	98°C, 10min
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17. Remove the reaction from the thermocycler. Immediately transfer to an ice-water bath and chill for 10 minutes. Centrifuge briefly. Return to ice bath.

18. Prepare the Klenow master mix on ice in 1.5ml microcentrifuge tube labeled Klenow Master Mix.

Component	Amount plus 0.05% for error
10mM dNTP Mix	10.5µl
Nuclease Free water	8.4µl
Klenow Fragment (3'-5' exo-) 50U/µl	2.1µl
Total	21.0µl

19. Add 20µl Klenow master mix to reaction (100µl total volume). Gently pipette the reaction 10 times to mix. Centrifuge briefly.

Component	Amount
dNTP/Klenow Mix	20.0µl
Total	100.0µl

20. Incubate for 2 hours at 37°C, lid temperature 105°C.

MasterCycler	37°C, 2hr
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21. During incubation, add 44µl of Isopropanol to labeled 1.5ml microcentrifuge tube, and then set aside.

22. Remove the reaction from the thermocycler. Centrifuge briefly.
23. Add 10.0µl of Stop Solution (0.5M EDTA, pH 8.0) to the reaction.

Component	Amount
Stop Solution	10.0µl
Total	110.0µl

24. Add 11.5µl of 5M NaCl to the reaction. Vortex to mix, and centrifuge briefly.

Component	Amount
5M NaCl	11.5µl
Total	121.5µl

25. Transfer the reaction to labeled 1.5ml microcentrifuge tube containing 110µl of Isopropanol for precipitation. Vortex well.

Component	Amount
Isopropanol	110.0µl
Total	231.5µl

26. Incubate for 10 minutes at room temperature, protected from light. Centrifuge for 10 minutes at 12,000 rcf. Decant supernatant.
27. Wash pellet with 500µl of 80% Ethanol Solution (v/v). Centrifuge for 2 minutes at 12,000 rcf. Decant supernatant.
28. SpeedVac the pellet on low heat for 5 minutes to remove traces of wash buffer.

STOP POINT: Labeled pellet may be stored at -20°C for up to 5 days, protected from light, until proceeding to next step.

29. Centrifuge briefly before opening. Rehydrate pellet with 25µl of Nuclease-free water. Incubate at room temperature, protected from light, for 5 minutes. Gently vortex and centrifuge briefly. Combine replicates into a single pool.
30. Determine concentration with NanoDrop. See section *NanoDrop Protocol for NimbleGen Microarray*.
31. Based on the concentration, calculate the volume of the test sample and reference sample required per hybridization for the array set based on the following table. Combine both test and reference samples into labeled 1.5ml microcentrifuge tube for hybridization.

Sample Requirement	2.1M Array	2.1M 2-Array Set	2.1M 3-Array Set	2.1M 4-Array Set
Test Sample	18.0 µg	36.0 µg	54.0 µg	72.0 µg
Reference Sample	18.0 µg	36.0 µg	54.0 µg	72.0 µg

32. SpeedVac on low heat, protected from light, until all the Nuclease-free water is removed, and the pooled labeled cDNA for hybridization is reduced to a pellet.

STOP POINT: Labeled cDNA pellet may be stored at -20°C , protected from light, until proceeding to next step.

NimbleGen Systems Hybridization

Introduction

NimbleGen Systems Hybridization technique allows for uniform hybridization of labeled DNA to produce high interarray correlations between replicate hybridizations. The protocol described here was adapted from Roche NimbleGen User's Guide for Expression Analysis for Cy-labeled cDNA derived from Eukaryote systems. We describe this procedure in detail for dual-color competitive hybridization of 2.1M Array Set format. Briefly, Cy-labeled cDNA is hybridized to high-density, long oligonucleotide microarray using custom microarray slide mixer and specialized hybridization system. At the completion of the procedure, the hybridized microarray is ready for data collection.

Precautions

Laboratory safety. It is assumed that users have a sound knowledge of molecular biology techniques and safe laboratory practices. Before undertaking a new protocol or using unfamiliar reagents users should review relevant Material Safety Data Sheets to identify potential hazards and recommended precautions. For background in general molecular biology please see Molecular Cloning A Laboratory Manual, J. Sambrook and D. W. Russell. Cold Spring Harbor Laboratory Press.

Caution. Hybridization Component A, containing formamide, is toxic in contact with skin and if swallowed. After contact with skin, wash the area immediately with copious amounts of water. This solution should be handled with caution.

Prevent excessive light exposure. Cy-labeled cDNA is light sensitive and should not be exposed to any unnecessary light. When working with this component turn off all lights and close the blinds to the windows.

Prevent excessive ozone and humidity exposure. Minimize ozone exposure and increased humidity levels as this leads to oxidation of the cyanine dyes. Use of desiccant materials or a dehumidifier should be considered.

Instrument Setup

1. Heat block, 95°C
2. Heat block, 42°C
3. NimbleGen Hybridization System, 42°C

Sample Preparation Setup

1. Thaw on ice from the NimbleGen Hybridization Kit,:
 - 1.1. 2X Hybridization Buffer
 - 1.2. Hybridization Component A
 - 1.3. Alignment Oligo Soln
2. Prepare
 - 2.1. Labeled 1 x 1.5ml microcentrifuge tube for Hybridization Master Mix
3. All centrifugation steps are at room temperature, unless otherwise noted.

Procedure

1. Set the Hybridization System to 42°C. With the cover closed, allow at least 3 hours for the temperature to stabilize. Set one heat block to 95°C, and a second heat block to 42°C, allow at least 3 hours for the temperature to stabilize.

1.1. For single array platforms, resuspend the dried labeled pellet with Nuclease-free water.

Component	2.1M Array	2.1M 2-Array Set	2.1M 3-Array Set	2.1M 4-Array Set
Nuclease-free water	12.3µl	24.6µl	36.9µl	49.2µl

1. Using components from the NimbleGen Hybridization Kit, prepare the Hybridization master mix according to the following table. The amounts listed in the table provide a sufficient volume (1.25x volumes) to hybridize all arrays of a single slide. When processing multiple slides, adjust the amounts for the number and type of array.

Component	2.1M Array	2.1M 2-Array Set	2.1M 3-Array Set	2.1M 4-Array Set
2X Hybridization Buffer	29.5µl	59.0µl	88.5µl	118.0µl
Hybridization Component A	11.8µl	23.6µl	35.4µl	47.2µl
Alignment Oligo	1.2µl	2.4µl	3.6µl	4.8µl
Total	42.5µl	85.0µl	127.5µl	170.0µl

2. Add the appropriate amount of hybridization master mix to each rehydrated labeled cDNA according to the following table.

Component	2.1M Array	2.1M 2-Array Set	2.1M 3-Array Set	2.1M 4-Array Set
Pooled labeled cDNA in Nuclease-free water	12.3µl	24.6µl	36.9µl	49.2µl
Hybridization Master Mix	31.7µl	63.4µl	95.1µl	126.8µl
Total Volume	44.0µl	88.0µl	132.0µl	176.0µl

3. Vortex well for 15 seconds, and centrifuge briefly.
4. Incubate the hybridization mixture for 5 minutes at 95°C, protected from light.
5. Incubate the hybridization mixture for 5 minutes at 42°C, protected from light, and maintain at 42°C until ready for sample loading. Vortex well for 15 seconds, and centrifuge prior to sample loading.

Prepare Mixer/Slide assembly for hybridization

Instrument Setup

1. NimbleGen Precision Mixer Alignment Tool (PMAT)
2. NimbleGen Mixer Brayer
3. Forceps
4. NimbleGen Hybridization System, 42°C

Sample Preparation Setup

1. NimbleGen Array Mixer
2. NimbleChip Microarray Slide

Procedure

1. Remove the appropriate mixer from its package. Handle the mixer by the Tab with the NimbleGen barcode. (Avoid touching the body of the mixer because this will transfer unwanted fingerprints and other residues, which may compromise array hybridization.)

Array Format	2.1M Array
Mixer	HX1 Mixer

2. Position the Precision Mixer Alignment Tool (PMAT) so the hinge is on the left, and then open it.
3. Snap the mixer onto the two alignment pins on the lid of the PMAT, with the tab end of the mixer toward the inside hinge and the mixer's adhesive gasket facing outward.
4. Carefully unpack the slide, and set aside its blue cassette (for use at a later time). While pushing back the PMAT's plastic spring with your thumb, place the slide in the base of the PMAT so that the barcode is on the right, farthest from the PMAT hinge, and the corner of the slide sits against the plastic spring. The NimbleGen logo and barcode number should be readable. Remove your thumb and make sure the spring is pushing the corner of the slide so that the entire slide is touching the edge of the PMAT closest to you. In addition, be sure that the slide is lying flat against the PMAT.
5. Using forceps, remove the protective backing from the mixer's adhesive and close the PMAT so that the mixer's adhesive makes contact with the slide.
6. Lift the lid by grasping the long edges of the PMAT while simultaneously applying pressure with a finger through the window in the lid of the PMAT to free the Mixer/Slide assembly from the pins of the PMAT. Be careful to avoid the arrays when applying pressure through the window.
7. Remove the Mixer/Slide assembly from the PMAT. Place the Mixer/Slide assembly on a 42°C heating block for 1 minute to facilitate adhesion of the mixer to the slide.
8. Rub the Mixer Brayer over the mixer with moderate evenly distributed pressure to adhere the mixer and remove any bubbles between the adhesive and slide's surface. For HX1 mixers, start in the center of the array and rub outwards. The adhesive will become clear when fully adhered to the slide's surface.
9. Place the Mixer/Slide assembly in the hybridization bay of the Hybridization System.

Load and hybridize samples to array

Instrument Setup

1. Microman Capillary Piston Pipette, CP100
2. Microman Capillary Piston Pipette Tips, CP100

Sample Preparation Setup

1. NimbleChip Microarray Mixer/Slide Assembly
2. Mixer Seals
3. Forceps
4. KimWipes

Procedure

1. At this point, the hybridization reaction and Mixer/Slide assembly are ready for sample loading.
2. Using the appropriate Gilson Microman pipette and tip, draw up the designated loading volume; inspect the pipette tip for bubbles. Discharge and reload the pipette tip if there is a bubble. It is recommended to have residual hybridization reaction in the tube to avoid introducing bubbles into the array. When ready, slowly dispense the appropriate loading volume into the fill port until the hybridization reaction starts to leak out of the array's corresponding vent port. Using a KimWipe, carefully to remove residual hybridization reaction surrounding the fill port and vent port. Adhered the mixer seal over the fill port and vent port. Close the hybridization system's bay clamp.

Component	2.1M Array
Loading Volume	41.0µl
Pipette & Tip	CP100

3. Be sure all slides have been processed: hybridization reactions loaded, mixer seals applied, and hybridization bay clamps closed. Turn on the mixing panel of the Hybridization System, with the mix mode set to Mode B.
4. Confirm the Hybridization System recognizes the slide in each occupied bay (indicator light becomes green when hybridization bay is in-use).
5. Hybridize for 16 – 20 hours at 42°C, mix Mode B.

Wash Hybridized Slides

Instrument Setup

1. Water bath, 48°C
2. Shallow dish (P1000 tip box lid, 500ml capacity)
3. NimbleGen Array Processing Containers
4. NimbleGen Disassembly Tool
5. ArrayIT High Speed Microarray Centrifuge

Sample Preparation Setup

1. 1X NimbleGen Wash Buffer I, 42°C & RT
2. 1X NimbleGen Wash Buffer II
3. 1X NimbleGen Wash Buffer III
4. All centrifugation steps are at room temperature, unless otherwise noted.

Procedure

1. **IMPORTANT** for ALL Washing Steps:
 - 1.1. Wash one slide at a time since the ArrayIT High Speed Microarray Centrifuge accommodates only one slide at a time. Restart washing protocol when each microarray slide is dry.
 - 1.2. Handle the slide from the barcode edge with a firm grip; do not use forceps. Barcode edge is at the far end of the Mixer Disassembly Tool; so the slide will have to be turned around (180 degrees) before inserting into wash containers. When transferring the slides into the next wash buffer, minimize the amount of wash buffer carryover by tapping the edge of the slide gently on a Kimwipe.
2. Setup ArrayIT High Speed Microarray Centrifuge with slide carriage.
3. Prepare NimbleGen Hybridization Wash Buffers,
 - 3.1. Pre-warm overnight 300ml of 1X Wash Buffer I in a water bath set to 48°C.
 - 3.2. Prepare 1X NimbleGen Wash Buffers in NimbleGen Array Processing containers.
 - 3.2.1. For the centrifuge tube with blue cap, final volume is 30ml.

SMALL SCALE (1 slide)				
Component	Wash I	Wash I	Wash II	Wash III
VWR water, Type I	270 ml	27 ml	27 ml	27 ml
10X NimbleGen Wash Buffer	30 ml	3 ml	3 ml	3 ml
1M DTT Soln	30 µl	3 µl	3 µl	3 µl
Total	300 ml	30 ml	30 ml	30 ml
Temperature	42°C	RT	RT	RT

4. Pour Wash Buffer I (48°C) into Shallow dish (P1000 tip box lid, 500ml capacity) with Disassembly Tool. Measure the temperature and wait until it reaches 42°C, then proceed immediately with slide washing. Heat Wash Buffer I if temperature is below 42°C. Microwave may be used to adjust wash buffer temperature.

5. Remove the Mixer/Slide assembly from the Hybridization System and immediately load it into the Disassembly Tool. Submerge the loaded Disassembly Tool into the Shallow dish containing warm Wash I (42°C). Do not allow the slide to cool.
6. With the Mixer/Slide assembly submerged, carefully peel the mixer off the slide. Discard the mixer. While submerged in Wash Buffer I, carefully remove the slide from the Disassembly Tool. Gently agitate the slide for 10 seconds, and quickly transfer the slide to Wash I, RT.
 - 6.2. It is important to agitate the slide in the warm Wash Buffer I (42°C). This will facilitate the removal of hybridization reaction from the surface of the slide. Failure to do so leads to poor washing results.
7. Wash Buffer I: Incubate for 2 minutes at RT with vigorous, constant agitation.

Wash Buffer I, RT	2 min
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8. Transfer slide to Wash Buffer II.
9. Wash Buffer II: Incubate for 1 minute at RT with vigorous, constant agitation.

Wash Buffer II, RT	1 min
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10. Transfer slide to Wash Buffer III.
11. Wash Buffer III: Incubate for 15 seconds at RT with vigorous, constant agitation.

Wash Buffer III, RT	15 sec
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12. Remove the slide from Wash III. Load it, active side up, into the ArrayIT High Speed Microarray Centrifuge. Centrifuge the slide for 1 minute.
13. Remove the slide from the carriage, and return the dry hybridized slide to its cassette.
14. Proceed immediately to scanning.

Two-color Array Scanning

Introduction

Array scanning technique allows for imaging of fluorescently labeled DNA to produce biologically relevant data from microarray hybridizations. Lasers excite the fluorescent dyes while photomultiplier tubes record the illuminated pixels as a high-resolution image file. The protocol described here was adapted from Roche NimbleGen Array User's Guide for CGH and CNV Arrays. The microarray is scanned using the NimbleGen MS 200 Microarray Scanner with resolution down to 2 μ m. Photomultiplier tube (PMT) Gain is automatically adjusted by the Data Collection software to achieve consistent, reproducible experimental results. After image acquisition, single-tiff images are ready to load into NimbleScan 2.6 Software to grid images for data extraction.

Precautions

Laboratory safety. It is assumed that users have a sound knowledge of molecular biology techniques and safe laboratory practices. Before undertaking a new protocol or using unfamiliar reagents users should review relevant Material Safety Data Sheets to identify potential hazards and recommended precautions. For background in general molecular biology please see Molecular Cloning A Laboratory Manual, J. Sambrook and D. W. Russell. Cold Spring Harbor Laboratory Press.

Prevent excessive light exposure. Hybridized Cy-labeled cDNA is light sensitive and should not be exposed to any unnecessary light. When working with this reagent turn off all lights and close the blinds to the windows.

Prevent excessive ozone and humidity exposure. Minimize ozone exposure and increased humidity levels as this leads to oxidation of the cyanine dyes. Use of desiccant materials or a dehumidifier should be considered.

Prevent loss of data. After image is saved, it is recommended to copy the files to a secondary independent data storage device or location.

Instrument Setup

1. NimbleGen MS 200 Microarray Scanner
2. NimbleGen MS 200 Data Collection Software

Procedure

1. Start the control unit and log into your user account as msOperator or other account as set up by your system administrators.
Account: msOperator
Password: 1-msOperator
2. Turn on the scanner using the power switch on the left side.
3. Insert slides into the Slide Magazine as described in Figure 1. Numbered slots provide spacing for inserting slides.

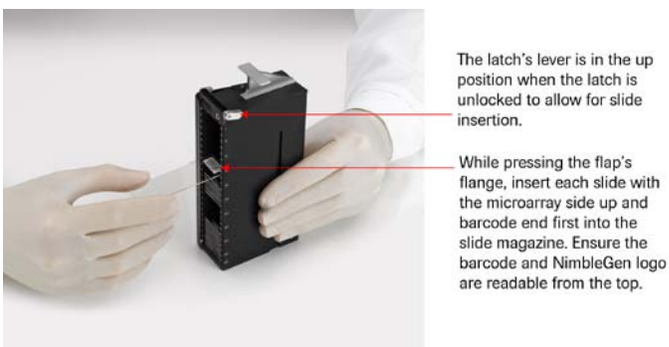


Figure 1: Inserting Slides into the Slide Magazine

4. Press the insert/eject magazine button on the scanner to open the stacker cover. Insert the slide magazine with loaded slides, aligning the slot on the slide magazine's side to join with the rail profile inside the scanner. The lowering of the slide magazine is interrupted by a mechanical hold point. Apply gentle pressure to complete insertion. Press the insert/eject magazine button to close the stacker cover (Figure 2). The initialization process starts, checking the slide magazine to determine which slots are occupied.



Figure 2: Inserting the Slide Magazine into the Scanner

5. Double-click the NimbleGen MS200 icon to launch the MS 200 Data Collection Software. Make sure that the software has completely loaded before continuing.
6. Click the **Green Laser** and **Red Laser** buttons in the Laser Control (Figure 3) to switch on the lasers. Allow lasers to warm for 10 minutes.

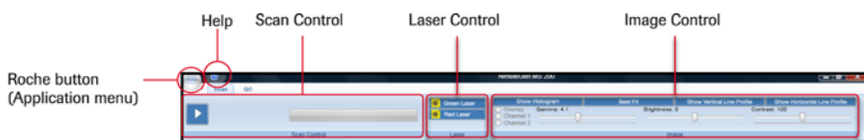


Figure 3: Top of Data Collection Workspace, showing Scan Control, Laser Control, and Image Control

7. Review the Magazine Control (Figure 4) in the Data Collection Software. Ensure that a green box appears in the *Slide Present* field for each slide loaded into the slide magazine.

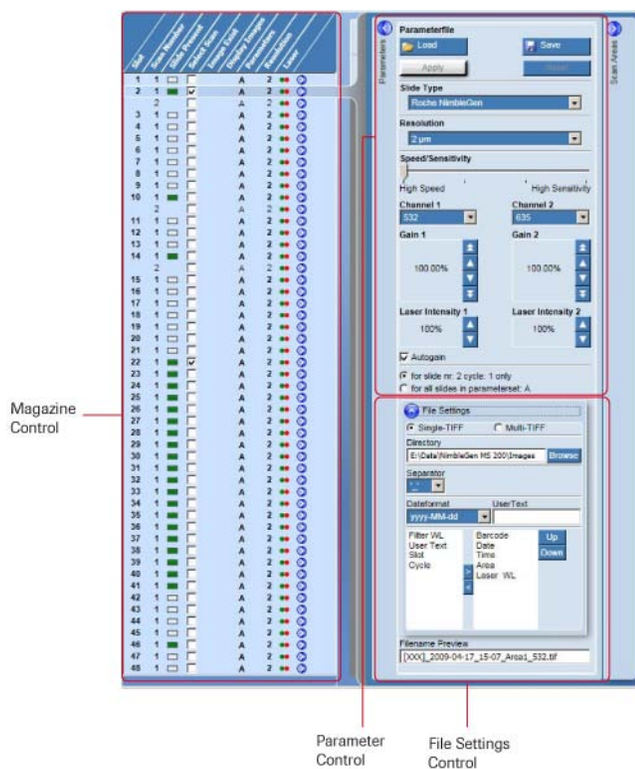


Figure 4: Magazine Control, Parameter Control, and File Setting Control

8. Use the Parameter Control (Figure 4) to set the parameters to use when scanning:
 - a. To open the Parameter Control if not displayed, go to the row of a slide to scan in the Magazine Control and click its **Open Parameter Control** button (🔍) to display the Parameter Control.
 - b. Do not adjust the following default parameters:
 - Slide type
 - Channel 1 and Channel 2
 - Laser Intensity 1 and Laser Intensity 2
 - for slide nr: 2 cycle: 1 only
 - c. Adjust the following default parameters if necessary:
 - Resolution
 - Speed/Sensitivity
 - Autogain
 - d. To save any changes to the settings, click **Apply**.
 - e. (Optional) Click **Save** to save settings to a parameters file, which allows the settings to be applied to other slides.

(Optional) To process the slide multiple times using the same or different parameters, create up to 12 cycles. To create a cycle, close the Parameter Control by clicking the **Close Parameter Control** button (🔍). Go to the slide row in the Magazine Control, right-click, and select **Add Cycle**. Open the Parameter Control. Then specify and save parameters for the cycle as described above (8b - 8e).
9. Use the File Settings Control (Figure 4) to specify image file settings:

- a. To open the File Settings Control if not displayed, go to the row of a slide to scan in the Magazine Control and click its **Open Parameter Control** button (🔍) (Figure 4). In the Parameter Control, click the **Open/Close File Settings** button (🔍) to open the File Settings Control.
- b. Choose the *Single-TIFF* option button to generate one image file in Tagged Image File Format (TIFF, .tif) per channel.
- c. If necessary, change the path to which the images files will be saved. The default path is E:\Data\NimbleGen MS 200. To change the path, click **Browse** to open a dialog box to specify a location in the directory and click **OK** to confirm.
- d. Use the annotation list to add or change annotations to include in file names. Click an annotation in the left list box and then click the right arrow button (➡) to add to the right list box.

If you will be using NimbleScan software for data analysis, specify and order the annotations as follows:

<Barcode>_<User Text>_<Laser WL>.tif

where “WL” means wavelength. To change the order of the annotations, select the annotation and click the **Up** or **Down** button. For “User Text,” make sure to type the text in the *User Text* field.

The *Filename Preview* text box of the Parameter Control displays the entire naming convention of the image file (.tif).

- e. Click **Apply** to confirm settings.
 - f. (Optional) To save the file settings to a file, click **Save** in the Parameters Control (Figure 4).
10. Use the Area Definition Control (Figure 5; denoted as *Scan Areas* in the software interface) to set scan, barcode, and autogain areas.
- a. To open the Area Definition Control if not displayed, in the Parameter Control, click the **Open/Close Area Definition Control** button (🔍) (shown on the right side of Figure 4 above the *Scan Areas* label in the software interface). Figure 5 shows the components of the Area Definition Control.

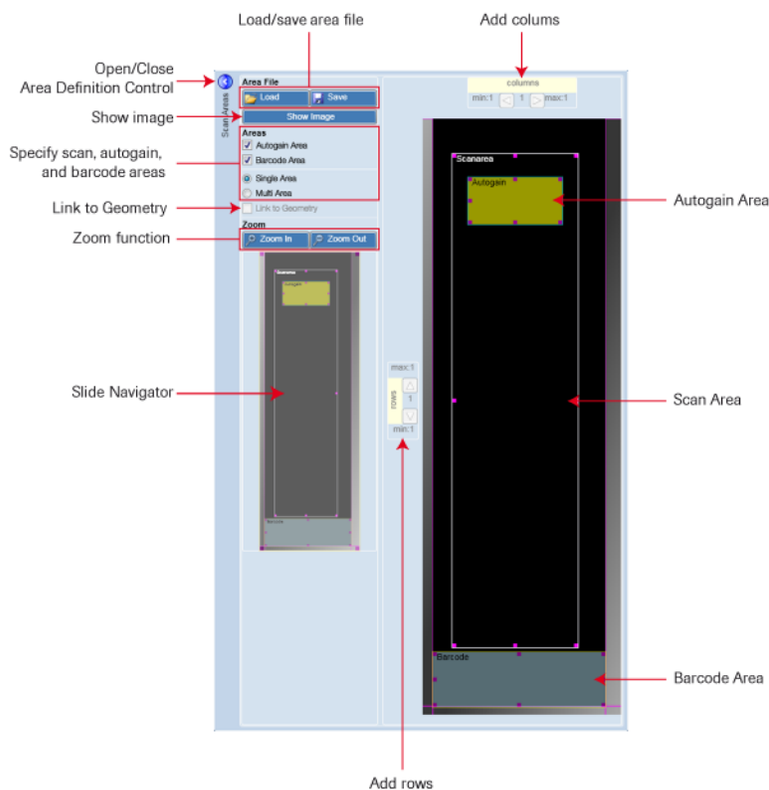


Figure 5: Components of the Area Definition Control

- b.** Ensure the following are selected:
 - *Barcode area* checkbox
 - *Autogain area* checkbox, if the Autogain checkbox was selected in the Parameter Control (Figure 4)
 - *Single Area* option - this is the preferred option when scanning single and multiplex NimbleGen arrays. For multiplex arrays, you will use NimbleScan software's burst functionality to create individual image files for multiplex arrays.
- c.** Review and if necessary adjust the scan area:
 - For NimbleGen 2.1M, 3x720K, and 12x135K arrays, use the default selection for the scan area.
 - (Optional) For NimbleGen 385K and 4x72K arrays, reduce the scan area, which will reduce the scan time. To define the scan area, download and use area files available at www.nimblegen.com/scanner/ under *Download*. These files identify scan areas that are specific for NimbleGen 385K or 4x72K arrays. For instructions on how to use area files or how to manually adjust the scan area, refer to the *NimbleGen MS 200 Microarray Scanner Operator's Manual*.
 - If desired, click **Save** to save settings to an area file for future use.
- d.** Review and if necessary adjust the autogain area.

To adjust the autogain area, position the mouse pointer inside the respective rectangle and click to enable the move cursor. Drag the rectangle to the desired array location and about 10 mm from the edge of the default scan area. Size the rectangle by dragging the side and corner handles (3 mm x 3 mm to 22 mm x 22 mm). Click **Apply** to confirm your settings.

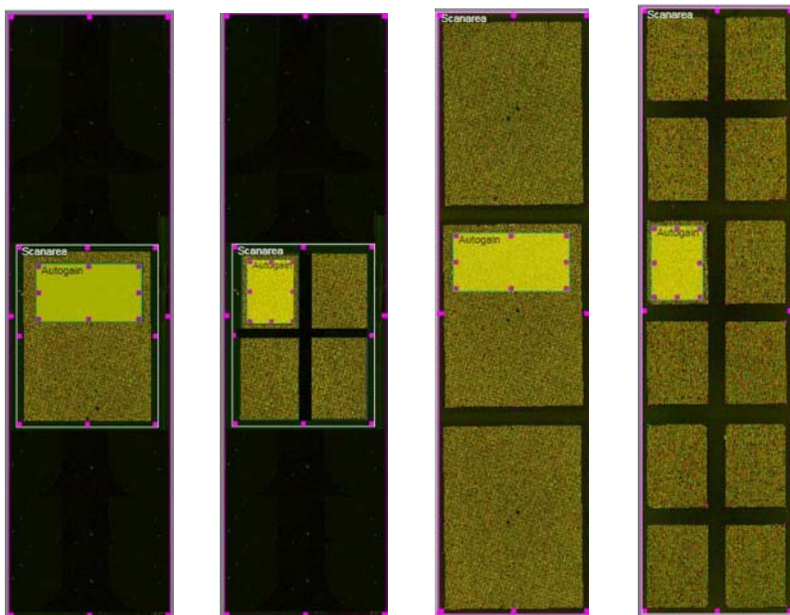


Figure 6: Example of Autogain Areas for NimbleGen Arrays

11. Repeat the instructions above for each slide to process during the experiment.

If you saved parameter file(s) and area file(s), click **Load** in the Parameter Control or Area Definition Control, respectively, to select a file and then click **Apply** to confirm your selection.

12. Click the **Start/Stop Scan** button () in the Scan Control (Figure 7).

13. When prompted, specify the folder and file naming to save the session file.

The scan process is then initiated. The Parameter Control and Area Definition Control close. The Image View, Slide View, and Spot View open (Figure 7), and the Scan Control shows the current task that is being performed above the progress bar.

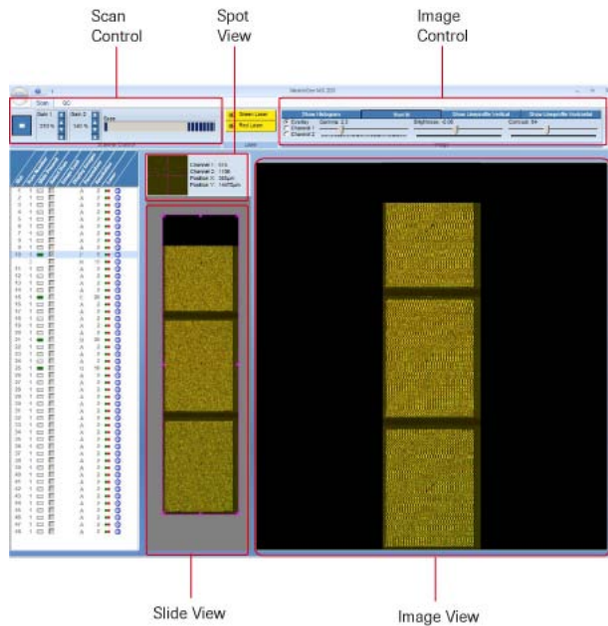


Figure 7: Example of a Run Time Window

14. (Optional) To manually adjust the PMT gain while scanning at 5 μm or higher resolution:

- a. Use the *Gain 1* and *Gain 2* spin boxes that appear in the Scan Control (Figure 7) to adjust the gain.



- b. Click **Show Histogram** in the Image Control (Figure 7) to view the histogram. Use the zoom feature (magenta box in the Slide View, **Error! Reference source not found.**Figure 9), to select where to view the histogram.



The best data are achieved when the red and green curves are superimposed or as close as possible to one another. If the red curve is above the green curve, lower the red curve (Channel 2). The curves should level out near $3 \log_{10}$.

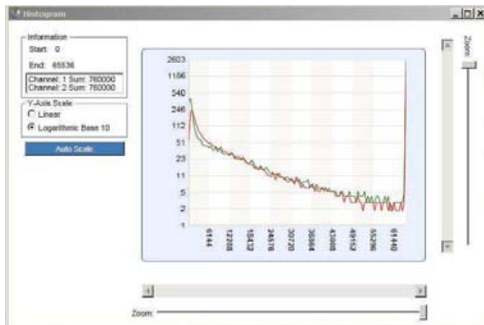


Figure 8: Histogram Window

- c. Record the final gain settings to use to scan the slide again.

15. After the scanning experiment is completed, the light in the upper-left corner of the insert/eject

magazine button on the scanner becomes green. The scanner unlocks its stacker cover, and you can remove the slide magazine.

16. Once the scan of a slide is completed, a blue **Displays Images** button (🖼️) appears in the *Display Images* field of the Magazine Control (Figure 4). Move the mouse pointer over the button to display the directory location and file name of the acquired image. Click the blue **Displays Images** button (🖼️) to display the image (both images) in the Image View.
17. To view a small region of the array and to select where to view the histogram (Figure 9), reduce the size of the magenta box that outlines the slide image in the Slide View so that the magenta box surrounds the area of interest. The area of interest will appear in the Image View. Click **Show/Hide Histogram** to display the histogram.

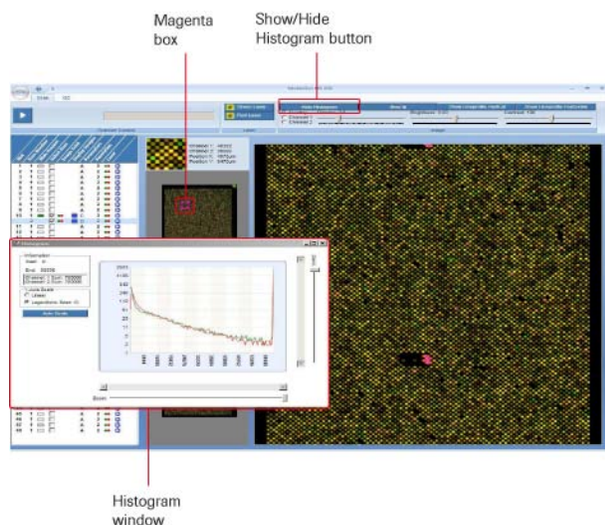


Figure 9: Example Window Showing Magenta Box in the Side View, Show/Hide Histogram Button, and Histogram Window

18. Adjust contrast and brightness using the Image Control (Figure 10) to provide an improved view of features on the image. Adjusting these settings allows faint features to be more easily seen.



Figure 10: Image Control

Data Processing of Images

Introduction

NimbleScan Software allows for automated gridding of microarray image files to produce quantifiable data for analysis. The protocol described here was adapted from Roche NimbleGen User's Guide for Expression analysis. Briefly, a tif image is uploaded into NimbleScan along with NimbleGen design files, which describe both probe identities and locations, to generate a tab-delimited report with probe identities and feature signal intensities. At the completion of the procedure, data is presented in an appropriate format for normalization and comparative analysis.

Precautions

Prevent loss of data. After raw data is saved, it is recommended to copy the files to a secondary independent data storage device or location.

Instrument Setup

1. Launch NimbleScan 2.6 Software.

Procedure

1. If the slide image is a multiplex array (4x72K or 12x135K), follow the steps below to separate each subarray into a separate image file. Otherwise, proceed to step 2.
 - 1.1. Select FILE > BURST MULTIPLEX IMAGE. The burst image dialog box appears. To choose the .tif files, click ADD IMAGES. Select the slide image of the same design. Click ADD TO BATCH. Browse to select the correct multiplex description file (.ncd). Browse to select the desired output file destination in the OUTPUT BURST IMAGES TO field.
 - 1.2. Click BURST. NimbleScan creates one file for each array of the slide, and each array designation for each burst image is appended to the original file name.
 - 1.3. Example: Barcode#A01_wavelength.tif
 - 1.4. Repeat steps 1 for each set of multiplex images of different designs.
2. Select FILE > OPEN. The open an alignment dialog box appears. Select a single array image. Select the design file (.ndf). Select the gene description file, if available. Click OPEN.
3. Once the image has loaded, the data is ready to match signal intensities with probe locations. Select the AUTO ALIGN TOOL (Ctrl + A). Zoom into the top corner of the array (Ctrl + G, then Ctrl + 7). Verify all fiducial controls line up correctly with the grid by shifting to each corner (Ctrl + 9, Ctrl +3, and Ctrl + 1).
4. Click FILE > SAVE to save the gridded image.
5. Confirm Experimental Integrity (STC report) for multiplex array formats only: Select ANALYZE > SAMPLE TRACKING. Click ADD FILES. Browse to select all 532nm gridded images of the multiplex array to be analyzed. Click ADD TO LIST. Click BROWSE to select destination of output file. Save outfile with extension .xls to review the STC report in MicroSoft Excel. Open the file. Ensure that a "PRESENT" call is reported for only the unique STC added to the hybridization reaction. If more than one "PRESENT" call is report, there may be contamination between hybridizations.
6. Create Pair Reports: Select ANALYZE > PAIR. Click ADD IMAGES to choose the files for analysis. Select the design file (.ndf). Browse to select the desired output file destination in

the Choose the destination folder field. Click REPORT. NimbleScan creates one PAIR report (.pair) for each image file.

- 6.1. For 385K and 4x72K array formats, select local alignment.
- 6.2. For 12x135K array formats, deselect local alignment.
7. The PAIR report is the raw data of the array hybridization. Other reports may be created. For more information see, NimbleScan Software User's Guide on <http://www.nimblegen.com> > Products > Software > literature.

NanoDrop Protocol for NimbleGen Microarray

Introduction

The NanoDrop ND-1000 utilizes UV/Vis spectroscopy to measure nucleic acid concentration. The ND-1000 is a micro-scale spectrophotometer that measures RNA or DNA concentration without dilution. With a pre-loaded application, Nucleic Acid, absorbance is measured across a broad wavelength range 190 – 840nm. Values for 260nm absorbance determine nucleic acid concentration, while 230nm and 280nm absorbance are used to calculate purity.

Precautions

Prevent loss of data. After report is generated, it is recommended to save the report and copy the files to a secondary independent data storage device or location.

Instrument Setup

1. Launch Nanodrop software to start application and adjust setting:
 - 2.1. Select assay “Nucleic Acid”.
3. A prompt will appear. Following the instructions from the prompt:
 - 3.1. With a KimWipe, gently buff the top and bottom pedestal 5 times.
 - 3.2. Load 1.5µl of nuclease-free water to pedestal. Gently, lower arm and click “OK”.

Procedure

1. Select “Sample Type” from drop-down menu displayed at the top left of the measurement window.
 - 1.1. For Total RNA and aRNA quantification,
 - 1.1.1. Select Sample type “RNA – 40” for extinction coefficient = 40
 - 1.2. For DNA quantification (i.e. ds cDNA and Cy-labeled cDNA),
 - 1.2.1. Select Sample type “DNA – 50” for extinction coefficient = 50
2. Buff top and bottom of pedestal to remove the nuclease-free water.
3. To BLANK, apply to the pedestal 1.5µl of elution buffer in which the nucleic acid is dissolved. Click BLANK to calibrate the instrument.
 - 3.1. To make sure the blank is calibrated properly, apply 1.5µl of Elution buffer and click Measure. Reading should be close to zero.
4. Instrument is now ready to measure concentration of the nucleic acid sample. Be sure to enter a “Sample Name” for each measurement.
5. To quantify sample, apply 1.5µl of the nucleic acid sample to the pedestal and click Measure.
 - 5.1. For Total RNA and aRNA,
 - 5.1.1. 260/280 ratio of > 2.0
 - 5.1.1.1. If the ratio is lower, this may indicate the presence of protein, phenol or other contaminants that absorb at or near 280 nm.
 - 5.1.2. 260/230 ratio range of 1.8 – 2.2
 - 5.1.2.1. If the ratio is low, this may indicate the presence of co-purified contaminants (e.g. Mg²⁺, salts, or other wash buffers).
 - 5.1.3. Spectrophotometer graph shows smooth curve.

- 5.2. For ds cDNA and Cy-labeled cDNA
 - 5.2.1. 260/280 ratio of > 2.0
 - 5.2.1.1. If the ratio is lower, this may indicate the presence of protein, phenol or other contaminants that absorb at or near 280 nm.
 - 5.2.2. 260/230 ratio range of 1.8 – 2.2
 - 5.2.2.1. If the ratio is low, this may indicate the presence of co-purified contaminants (e.g. Mg²⁺, salts, or other wash buffers).
6. Save Report:
 - 6.1. From Menu Bar at top of window, select REPORTS and click Save Report.
 - 6.1.1. Select Export Table (Option 2) for a tab-delimited text file.
 - 6.1.1.1. This file can be viewed in MS Excel application.
 - 6.1.2. Select Full Report for a NanoDrop formatted file.
 - 6.1.2.1. This file can be viewed in the NanoDrop software only. In addition, the report can be loaded at the start of the next session, if a single continuous record / file is desired.
 - 6.1.2.2. To load previous full report, select REPORTS and click Load Report. Browse for the desired report, select it, and click OK to load it. Once loaded, measurements taken during the new session will add to the report. At the end of the session, Save the Full Report.
 - 6.1.3. Browse for the file destination and enter an appropriate file name. Click OK.

7. Trouble shooting:

- 7.1. Make sure sample pedestal is clean. Use 2.0 μ l of deionized water to wash pedestal and wipe dry with a KimWipe.
- 7.2. Redo the calibration with nuclease-free water step presented at the start of the software setup.
- 7.3. Redo the blank setup. If blank measurement is not done properly, strange results will occur.
- 7.4. Nucleic acid sample may not be homogenized. Gently mixing the nucleic acid sample by finger-flicking the micro-centrifuge tube prior to measuring the concentration is recommended. Re-measure the sample.
- 7.5. Use a 1.5 – 2.0 μ l sample size when measuring. Strange results occur when the liquid sample column is not completely formed during the measurement. While making a measurement, visually confirm the water column is completely formed.
- 7.6. If this does not resolve the issue, the nucleic acid sample may need to be re-purified. Recommend Microcon YM-30 Centrifugal Filter Unit (See Manufacturer's Manual).

Bioanalyzer RNA Protocol for NimbleGen Microarray

Introduction

Agilent Bioanalyzer system and reagent kits are designed to analysis limited samples by microfluidic capillary electrophoresis. By replacing ethidium bromide with fluorescent dye, the system is a safer alternative for nucleic acid analysis. Standardized analysis applications, included with the analysis software, provide reliable data, which can be compared between independent chip runs.

Precautions

Laboratory safety. It is assumed that users have a sound knowledge of molecular biology techniques and safe laboratory practices. Before undertaking a new protocol or using unfamiliar reagents users should review relevant Material Safety Data Sheets to identify potential hazards and recommended precautions. For background in general molecular biology please see Molecular Cloning A Laboratory Manual, J. Sambrook and D. W. Russell. Cold Spring Harbor Laboratory Press.

Caution. RNA 6000 dye concentrate contains DMSO, a molecule that facilitates chemical absorption into cells. All materials should be handled with caution.

Prevent loss of data. After report is generated, it is recommended to save the report and copy the files to a secondary independent data storage device or location.

Instrument Setup

1. Confirm the electrophoresis cartridge is installed in the Bioanalyzer 2100 System.
 - 1.1. Electrophoresis cartridge is 1.
2. Switch ON the Bioanalyzer 2100 System.
 - 2.1. Green Light: ON and ready for measurement
 - 2.2. Green Blinking: Measuring
 - 2.3. Orange Blinking: Busy, not measuring
 - 2.4. Red Light: Not ready
3. Switch ON the computer system operating the Bioanalyzer 2100 System and Bioanalyzer 2100 Expert Software.
4. Launch Bioanalyzer 2100 Expert Software.
5. After the software opens, the Instrument control panel will appear.
6. Make sure the Bioanalyzer 2100 System has been detected.
 - 6.1. From instrument control panel, select the instrument.
7. Click on METHODS, select the method for measurement.
 - 7.1. Eukaryote Total RNA Nano
 - 7.1.1. Evaluates 18S and 28S based Eukaryote RNA
 - 7.2. Prokaryote Total RNA Nano
 - 7.2.1. Evaluates 16S and 23S based Prokaryote RNA
 - 7.3. mRNA Nano
 - 7.3.1. Evaluates distribution of nucleic acid molecule to detect ribosomal RNA contamination for amplified RNA (aRNA) and double-stranded (ds) cDNA

8. Input Chip Information.
 - 8.1. Sample Name
 - 8.2. Sample Comments
 - 8.3. Chip Lot Number
 - 8.4. Kit Lot Number
 - 8.5. Comments
9. Wash the electrode pins and allow drying for 10 minutes.
 - 9.1. Obtain electrode-cleansers from the Bioanalyzer Kit, label one RnaseZap and label the other Nuclease-free water.
 - 9.2. Slowly fill the first with RnaseZap (350 μ l) and the second with nuclease free water (350 μ l).
 - 9.3. Open the lid and place the electrode cleanser with RnaseZap into the Agilent 2100 Bioanalyzer. Close the lid and leave it for 1 minute. Open the lid and remove the electrode cleanser chip from the Agilent 2100 Bioanalyzer.
 - 9.4. Open the lid and place the electrode cleanser with nuclease-free water into the Agilent 2100 Bioanalyzer. Close the lid and leave it for 10 seconds. Open the lid and remove the electrode cleanser chip from the Agilent 2100 Bioanalyzer. Keep the lid open for 10 seconds then close the lid.
10. Setup Chip Priming Station.
 - 10.1. Replace the syringe.
 - 10.1.1. Replace the syringe with each new kit or after 25 chips have been processed.
 - 10.1.1.1. Unscrew the old syringe from the lid of the chip priming station.
 - 10.1.1.2. Release the old syringe from the clip. Discard the old syringe.
 - 10.1.1.3. Remove the plastic cap of the new syringe and insert it into the clip.
 - 10.1.1.4. Slide it into the hole of the luer lock adapter and screw it tightly to the chip priming station.
 - 10.2. Adjust the base plate.
 - 10.2.1. Open the chip priming station by pulling the latch.
 - 10.2.2. Lift the base plate and insert it again in position C.
 - 10.3. Adjust the syringe clip.
 - 10.3.1. Release the lever of the clip and slide it up to the top position.

11. Proceed with Sample Preparation.

Sample Preparation Setup

1. Prepare Nucleic Acid sample(s) to a final concentration of 100ng/ μ l - 400ng/ μ l.
 - 1.1. Total RNA
 - 1.1.1. 150ng per 1.5 μ l of nuclease-free water in a 0.2mL micro-centrifuge tube. Mix by gently pipetting. Centrifuge briefly to collect the contents.
 - 1.2. Amplified RNA (aRNA)
 - 1.2.1. 400ng per 1.5 μ l of nuclease-free water in a 0.2mL micro-centrifuge tube. Mix by gently pipetting. Centrifuge briefly to collect the contents.
 - 1.3. Double-stranded cDNA
 - 1.3.1. 100ng per 1.5 μ l of nuclease-free water in a 0.2mL micro-centrifuge tube. Mix by gently pipetting. Centrifuge briefly to collect the contents.

2. Prepare Nucleic Acid standard.
 - 2.1. RNA 6000 Ladder, Agilent
 - 2.1.1. Transfer 1.3 μ l to a 0.2mL micro-centrifuge tube. Mix by gently pipetting. Centrifuge briefly to collect contents.
3. Heat-denature Nucleic Acid sample(s) and Nucleic Acid standard.
 - 3.1. 70°C for 2 minutes
 - 3.2. Immediately chill in ice bath.
4. Keep Nucleic Acid sample(s) and Nucleic Acid standard on ice until ready to analyze.
 - 4.1. Analyze within 30 minutes. Otherwise return samples to the freezer (-20°C) until ready to process.

Procedure

1. Allow all reagents from Bioanalyzer kit to equilibrate to room temperature protected from light for 30 minutes before use.
 - 1.1. RNA 6000 Nano Marker (Green)
 - 1.2. RNA 6000 Nano dye concentrate (Blue)
 - 1.3. RNA 6000 Gel Matrix
 - 1.3.1. If filtered Gel Matrix is available, equilibrate to room temperature for 30 minutes and skip “prepare gel matrix”.
 - 1.3.2. RNA 6000 Nano Gel Matrix (Red)
2. Prepare gel matrix. (Skip if filtered gel matrix is available.)
 - 2.3. Place 550 μ l of Agilent 6000 Nano gel matrix into the top receptacle of a spin filter (provided).
 - 2.4. Place the spin filter in a microcentrifuge and spin for 10 minutes at 1500 rpm.
 - 2.5. Aliquot 65 μ l filtered gel into 0.5ml Rnase-free tubes (provided). Store unused aliquots at 4°C and use within one month of preparation.
3. Prepare gel-dye mix.
 - 3.3. Vortex RNA 6000 Nano dye concentrate for 10 seconds and centrifuge briefly.
 - 3.4. Add 1 μ l of RNA 6000 Nano dye concentrate to 65 μ l aliquot of filtered gel. Vortex well. Centrifuge for 10 minutes at room temperature at 14000 rpm.
 - 3.5. Use gel-dye mix immediately. Otherwise store at 4°C for up one day. Equilibrate to room temperature for 30 minutes. Centrifuge gel-dye mix for 10 minutes at 14,000 rpm prior to use.
4. Load gel-dye mix.
 - 4.3. Take a new RNA Nano chip out of its sealed bag.
 - 4.4. Place the chip on the priming station.
 - 4.5. Pipette 9.0 μ l of the gel-dye mix from the upper layer of the 0.5ml microcentrifuge tube and dispense into the bottom of the well marked with the dark encircled G of the RNA Nano chip.
 - 4.5.1. When pipetting the gel-dye mix from the 0.5ml microcentrifuge tube, make sure not to draw up particles from the bottom of the gel-dye mix tube. Insert the tip of the pipette to the bottom of the chip well when dispensing. This prevents a large air bubble forming under the gel-dye mix. Placing the pipette at the edge of the well may lead to poor results.

- 4.6. Set the plunger is positioned at 1 ml and then close the chip priming station. Depress the plunger to luer clamp. Hold in place for 30 seconds. Release the plunger clip. Wait 5 seconds. Return the plunger to 1ml position. Open the Priming Station.
5. Load RNA 6000 Nano Marker.
 - 5.3. Pipette 5 μ l of the RNA 6000 Nano marker into the well marked with the ladder symbol and each of the sample wells to be used. For unused sample wells, pipette 6 μ l of the RNA 6000 Nano marker.
 - 5.3.1. Do not leave any wells empty or the chip will not run properly. Unused wells must be filled with 6 μ l of the RNA 6000 Nano marker.
6. Load Ladder and Samples.
 - 6.3. Pipette 1 μ l of the RNA ladder into the well marked with the ladder symbol.
 - 6.4. Pipette 1 μ l of each sample into a separate sample wells.
7. Vortex the chip.
 - 7.3. Place the chip horizontally in the adapter of the IKA vortex mixer and secure the chip to the carriage. Vortex the chip for 1 minutes at 2400 rpm. Remove the chip from the vortex and remove any spilled liquid with a Kimwipe.
8. Insert Chip in the Agilent 2100 Bioanalyzer.
 - 8.3. Open the lid to the Agilent 2100 Bioanalyzer.
 - 8.4. Place the chip into the station.
 - 8.5. Gently close the lid to the Agilent 2100 Bioanalyzer.

Instrument Operation

1. Clicking START to begin the Chip Run.
 - 1.1. Ensure the appropriate assay has been selected.
 - 1.2. Adjust the wells to be measured setting if necessary.
 - 1.3. The runtime is approximately 25 minutes (7 minutes to warm plus 1.5 minutes per well measurement).
2. At the completion of the Chip Run, clean the instrument and dispose of the used chip from the run.
 - 2.1. Open the lid and remove the used chip from the Agilent 2100 Bioanalyzer. Place the electrode cleanser with fresh nuclease-free water into the Agilent 2100 Bioanalyzer. Close the lid and leave it for 10 seconds. Open the lid and remove the electrode cleanser chip from the Agilent 2100 Bioanalyzer. Keep the lid open for 10 seconds then close the lid.

Results / Interpretations

1. Moderate to High Quality Total RNA Profile
 - 1.1. Distinct 18S and 28S peaks
 - 1.2. Low noise between peaks
 - 1.3. Minimal low-molecular weight contamination
 - 1.4. Minimal high-molecular weight contamination.
 - 1.5. High molecular weight contamination may indicate the presence of a contaminant (i.e. DNA).

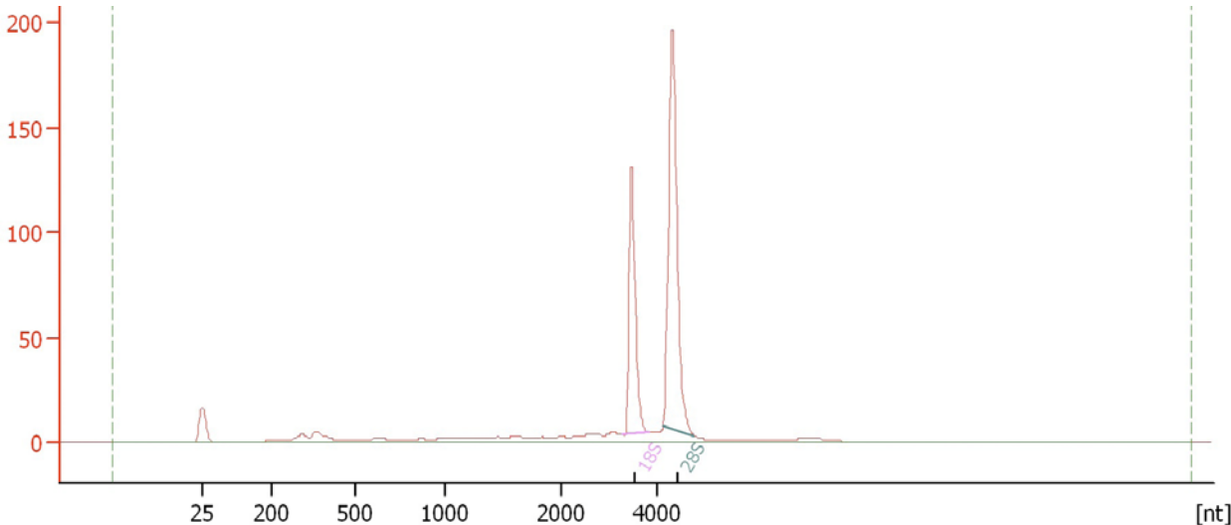


Figure 1 Electropherogram of Total RNA

2. High Quality amplified RNA (aRNA) Profile

- 2.1. One marker peak
- 2.2. Broad distribution (medial-molecular weight molecules)
- 2.3. Minimal low-molecular weight molecules
- 2.4. Minimal ribosomal RNA contamination (< 5%)

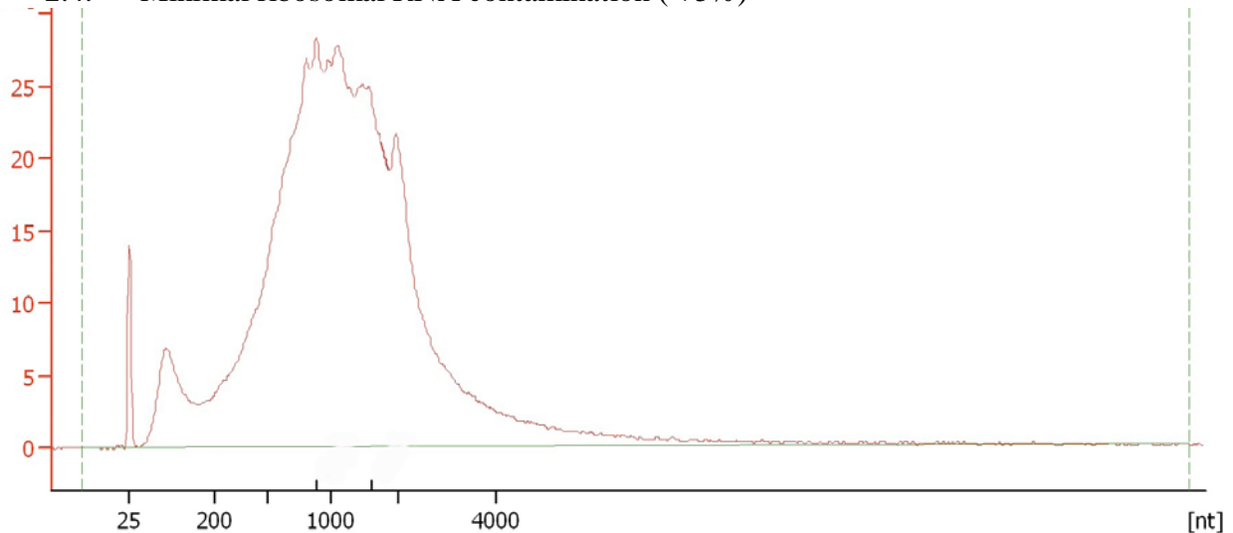


Figure 2 Electropherogram of aRNA

3. High Quality ds cDNA Profile

- 3.1. One marker peak
- 3.2. Broad distribution (medial-molecular weight molecules)
- 3.3. Minimal low-molecular weight molecules
- 3.4. Minimal ribosomal RNA contamination (< 5%)

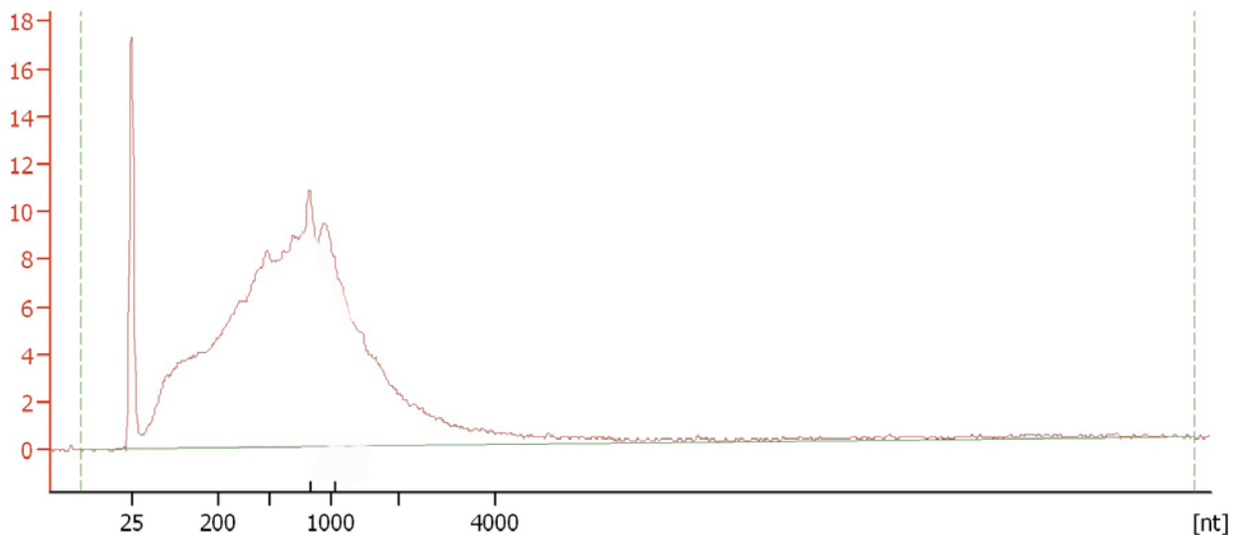


Figure 3 Electropherogram of ds cDNA