nrStar™ Human tRNA Repertoire PCR Array

Cat#: AS-NR-001

Instruction Manual Version 1.0

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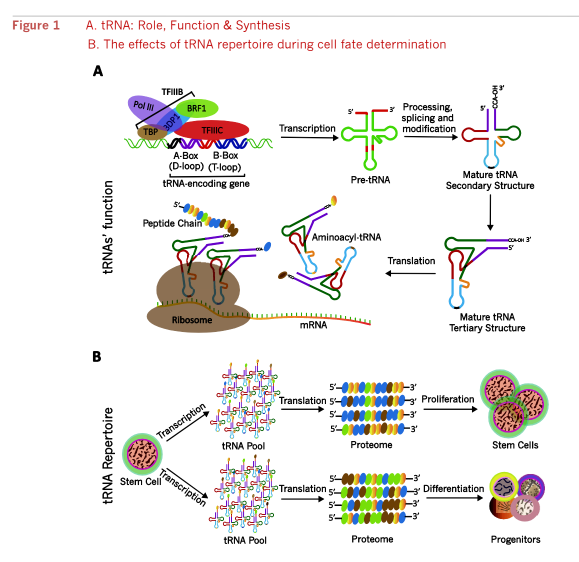
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I. Introduction

A. Overview

Transfer RNAs (tRNAs) are ubiquitous and the most abundant of all small non-coding RNA molecules. As a fundamental component in translation, tRNAs serve as the physical link between the mRNA coding and protein sequences (**Figure 1**). Genomes exhibit substantial variations in their preference for specific codons across their coding sequences. The source of this bias, though still debated, likely reflects selection for translational efficiency and accuracy[1-3]. Alterations of tRNA repertoire affect cell-fate choices during cell development (**Fig. 1B**), and dysregulated tRNAs expression are associated with many disease including cancer. nrStar™ Human tRNA Repertoire PCR Array profiles 66 PCR-distinguishable nuclear tRNA isodecoders and all mitochondrial tRNA species in the human tRNA repertoire by optimized SYBR Green qPCR assays in a PCR panel. The panel covers all anti-codons compiled in GtRNAdb and tRNAdb database, making it a powerful tool for analyzing the profile of tRNA repertoire. As tRNAs are heavily modified posttranscriptionally, which badly affect tRNA cDNA synthesis efficiency, scientists at Arraystar have developed rtStar™ tRNA-optimized First-Strand cDNA Synthesis Kit to efficiently remove the tRNA modifications and greatly improve the cDNA synthesis quality. With the powerful combination of this kit and the PCR array, researchers can obtain a new level of accuracy on the tRNA pool alterations and gain further insight to interpret the proteome and tRNA-derived fragments.



B. tRNA repertoire and cell state

A wide variety of biological processes like cell proliferation[4], differentiation[4, 5] and apoptosis[6] are always accompanied with tRNA level variation (**Fig. 1B**). It has been found that codon usage is different between genes serving cell-autonomous functions and genes involved in multicellularity. Proliferation-induced and differentiation -induced tRNAs often carry anti-codons that correspond to the codons enriched among the cell-autonomous and the multi- cellularity genes, respectively. Because mRNAs of cell-autonomous genes are induced in proliferation and cancer in particular, the concomitant induction of their codon-enriched tRNAs suggests coordination between transcription and translation[4]. According to another study, overexpression of tRNAi (Met) significantly alters the global tRNAs expression profile and results in increased cell metabolic activity and cell proliferation[5]. In addition, tRNA regulates apoptotic sensitivity at the level of cytochrome c mediated apoptosome formation[6]. Microinjection of tRNA can inhibit cytochrome c-induced apoptosis[7]. In sum, alteration of tRNA levels can change the cell state by various mechanisms.

C. tRNA repertoire and diseases

Multiple lines of evidence have associated the disruption of tRNA levels with many diseases. For example, certain dysregulated tRNAs can induce tumorigenesis and cancer progression[5]. Studying tRNA repertoire has become increasingly important in human diseases.

Cancer

After cataloging the tRNA repertoire, Gingold et al demonstrated there are different tRNA pools between cancer cells and differentiated cells[4]. tRNAs that are upregulated in differentiated/arrested cells are repressed in proliferating cells. Conversely, tRNAs whose levels are high in proliferating cells become low in differentiated/arrested cells. Cancer cells appear to adjust their tRNA pools to selectively bolster translation of mRNAs that are required for tumor progression. When comparing tRNAs expression in breast tumors versus normal breast tissues, Pavon-Eternod et al found that nuclear- and mitochondrial-encoded tRNAs exhibit distinct expression patterns, indicating the potentials of using tRNAs as biomarkers for breast cancer[8]. Recently, Goodarzi et al also confirmed that specific tRNAs are upregulated in human breast cancer cells as they gain metastatic activity, among which tRNA^{Glu-UUC} and tRNA^{Arg-CCG} are implicated as promoters of breast cancer metastasis. Further studies conclusively showed that tRNA^{Glu-UUC} promotes metastatic progression by directly enhancing EXOSC2 and GRIPAP1 expression [9]. These and other cases have conclusively demonstrated dysregulated tRNA repertoire can promote tumorigenesis and cancer progression[5, 8-15].

Huntington's disease

Huntington's disease (HD), a dominantly inherited neurodegenerative disorder caused by the expansion of a CAG-encoded polyglutamine (polyQ) repeat in huntingtin, displays a highly heterogeneous etiopathology and disease onset. Analyses of HD-affected brain tissues revealed traces of polyalanine (polyA) or polyserine (polyS) proteins within the polyQ aggregates. These species probably result from a shift in the GIn-encoding CAG frame to an Ala-encoding ·1 GCA frame or a Ser-encoding +1 AGC frame. But what is the role of translational frame-shifting in the pathogenesis of polyQ diseases? Girstmair et al found that depletion of tRNA^{GIn-CUG} pairing to the CAG codon was the main cause of ·1 frameshifting. In addition, frameshifted proteins form morphologically distinct aggregates in vivo dependent on the Q:A ratio. These results suggested that frameshifting within expanded CAG stretches may contribute to the heterogeneity in the course and onset of HD at both cellular and individual level[16].

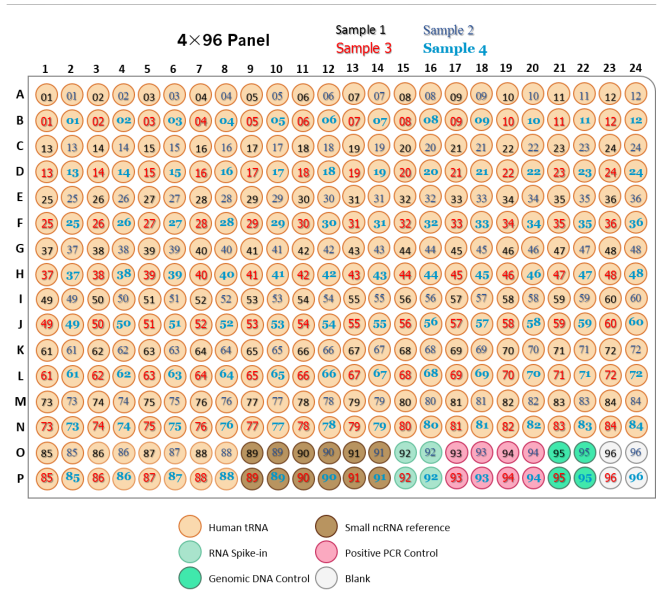
Virus Infection

Viruses are wholly dependent on the host translation machinery to synthesize their proteins. Consequently, viral

codon usage is thought to be under selective pressure to adapt to the host cell tRNA pool. Since host codon usage generally reflects the host tRNA pool, viral translation should be more efficient when viral codon usage is similar to that of the host genes. In many cases, however, viral codon usage seems poorly adapted to that of its host. After profiling the tRNA repertoire, Pavon-Eternod et al found that influenza A and vaccinia viruses could manipulate tRNA populations to favor translation of their own viral genes[17]. In another research, the codon usage of HIV-1 early genes is similar to that of highly expressed host genes, but codon usage of HIV-1 late genes was better adapted to the altered tRNA pool late in the viral infection[18]. This is a striking example of the virus modulating the tRNA pool to optimize its translation efficiency..

D. Product summary

nrStar™ Human tRNA Repertoire PCR Array profiles 66 PCR-distinguishable nuclear tRNA isodecoders and all human mitochondrial tRNA species, covering all anti-codons compiled in GtRNAdb and tRNAdb databases.





Kit Contents

Table 1	List of	human	t RNAs	and	the	Controls
	LISC OI	nunun	1111/13	and	the	001111015

01	Ala-AGC	25	His-GTG	49	Ser-TGA	73	mt-Glu-TTC
02	Ala-CGC	26	IIe-AAT	50	Sup-CTA	74	mt-Gly-TCC
03	Ala-GGC	27	IIe-TAT	51	Sup-TTA	75	mt-His-GTG
04	Ala-TGC	28	Ini-CAT	52	Thr-AGT-1	76	mt-Ile-GAT
05	Arg·ACG	29	Leu-AAG	53	Thr-AGT-2	77	mt-Leu-TAA
06	Arg-CCG	30	Leu-CAA	54	Thr-CGT	78	mt-Leu-TAG
07	Arg-CCT	31	Leu-CAG	55	Thr-TGT-1	79	mt·Lys·TTT
08	Arg-TCG	32	Leu-TAA	56	Thr-TGT-2	80	mt-Met-CAT
09	Arg-TCT	33	Leu-TAG	57	Trp-CCA	81	mt-Phe-GAA
10	Asn-ATT	34	Lys-CTT-1	58	Tyr-ATA	82	mt-Pro-TGG
11	Asn-GTT	35	Lys-CTT-2	59	Tyr-GTA-1	83	mt-Ser-GCT
12	Asp-ATC	36	Lys-TTT	60	Tyr-GTA-2	84	mt-Ser-TGA
13	Asp-GTC	37	Met-CAT	61	Val-AAC	85	mt-Thr-TGT
14	Cys-GCA	38	Phe-GAA	62	Val-CAC-1	86	mt-Trp-TCA
15	GIn-CTG-1	39	Pro-AGG	63	Val-CAC-2	87	mt-Tyr-GTA
16	GIn-CTG-2	40	Pro-CGG	64	Val-CAC-3	88	mt-Val-TAC
17	GIn-TTG-1	41	Pro-GGG	65	Val-TAC-1	89	RNU6-2
18	GIn-TTG-2	42	Pro-TGG	66	Val-TAC-2	90	SNORD43
19	Glu-CTC	43	Sec-TCA	67	mt-Ala-TGC	91	SNORD95
20	Glu-TTC	44	Ser-ACT	68	mt-Arg-TCG	92	RNA Spike-in
21	Gly-CCC-1	45	Ser-AGA	69	mt-Asn-GTT	93	PPC
22	Gly-CCC-2	46	Ser-CGA	70	mt-Asp-GTC	94	PPC
23	Gly-GCC	47	Ser-GCT	71	mt-Cys-GCA	95	GDC
24	Gly-TCC	48	Ser-GGA	72	mt·GIn·TTG	96	Blank

Description of the control assays

nrStar[™] Human tRNA Repertoire PCR Array includes a series of external and internal controls for effective correction and normalization of sample and qPCR variabilities. In addition, Positive PCR Control and Genomic DNA Control are included to monitor the experiment process and the quality of RNA sample. These controls are described below.

- Ref (small ncRNA Control Reference; Internal Controls): Three stably expressed small ncRNA genes RNU6-2 (Ref 1), SNORD43 (Ref 2), and SNORD95 (Ref 3) are included in the array as the quantification references for tRNA. nrStar[™] PCR system provides multiple reference genes selected among commonly used reference genes by using a stringent bioinformatic algorithm, which offers the flexibility of choosing the most valid reference gene(s) for qPCR normalization for your sample types.
- **RNA Spike-In** (External Control): One External RNA Spike-In Mix is added in the RNA sample prior to the first strand cDNA synthesis. The RNA Spike-In control assay indicates the overall success and the efficiency of the reaction beginning from the cDNA synthesis to the final qPCR. Any problem(s) in these steps will result in a

failed or compromised RNA Spike-In outcome. RNA spike-in assay results for samples are compared and outliers or failed reactions may be identified and excluded from further data analysis.

- **PPC** (Positive PCR control): 2 replicates of one artificial DNAs and the PCR primer pairs to indicate the qPCR amplification efficiency. A Ct value greater than 25 is an indication of low qPCR amplification efficiency. More importantly, the PPC are used as inter-plate calibrator (IPC) between PCR plate runs to give the same Ct value for the calibrator, thereby reducing run-to-run variance. Inter-plate calibration (IPC) can easily be performed with the data analysis software available on our website (<u>www.arraystar.com</u>).
- **GDC** (Genomic DNA Control): The control assay consists of PCR primers for an untranscribed genomic region. Non-RT sample or RNA sample are added during the qPCR Process. The Ct values should be greater than 35. A positive GDC signal indicates the array result may have been compromised with genomic DNA contamination.

Shipping and Storage

nrStar[™] PCR Arrays are shipped at ambient temperature, on ice, or on dry ice depending on the destination and accompanying products. Store at –20°C upon receipt. The contents are stable for at least 6 months.

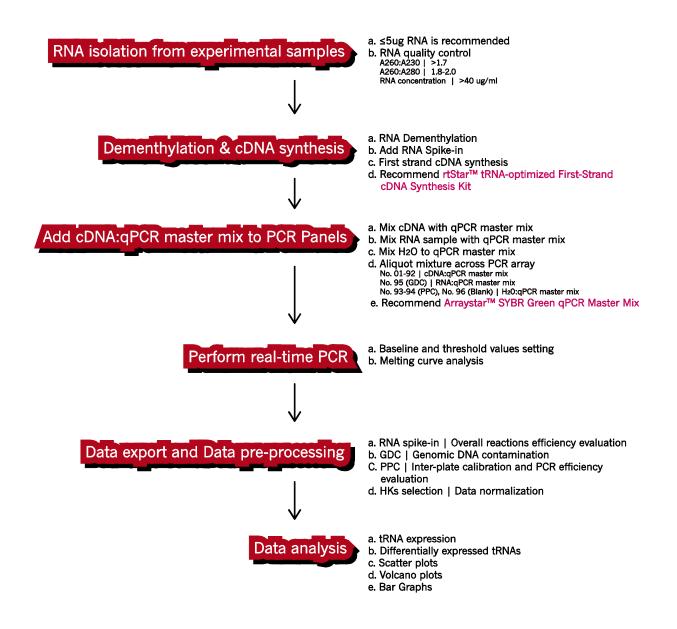
Additional Required Equipment

- Thermal cycler
- Real time qPCR instrument, compatible with 384-well format

Additional Required Reagents

- rtStar™ tRNA.optimized First.Strand cDNA Synthesis Kit (Cat# AS.FS.004)
- Arraystar SYBR® Green qPCR Master Mix(ROX+) (AS·MR·006·5)
- Nuclease-free PCR-grade water

E. Protocol overview



II. Protocol

IMPORTANT: Please read through this entire protocol before your first experiment. Prepare a workspace and use labwares free of DNA, RNA, and nuclease contamination. Wear gloves while performing this protocol.

A. RNA sample preparation and quality control

For best results from the PCR array, RNA samples must meet the QC standards of integrity and purity from protein, organics, genomic DNA contamination and excessive RNA degradation. You can measure the RNA concentration and purity by UV spectrophotometry. You can also check 18S and 28S ribosomal RNA as an indicator of RNA integrity by denaturing gel electrophoresis or by using an Agilent BioAnalyzer.

- A260:A230 ratio greater than 1.7.
- A260:A280 ratio between 1.8 and 2.0.
- Total RNA concentration greater than 40 ng/ μI

Eliminating genomic DNA contamination is essential for accurate gene expression profiling by qPCR, which is particularly important for genes at low expression levels. Due to the presence of pseudogenes, even cross-intron primers are not a guarantee of avoiding amplification from contaminating genomic DNA. The Genomic DNA Control (GDC) in the PCR Array specifically detects potential genomic DNA contamination. A Ct for GDC less than 35 indicates the presence of genomic DNA contamination that may compromise the qPCR results.

B. First-strand cDNA synthesis

Use the same amount of total RNA in this reaction for every sample. High quality cDNA synthesis is vital for the following qPCR performance. Because of the disincentive effects of tRNA modification on cDNA synthesis, we highly recommend using rtStar[™] tRNA-optimized First-Strand cDNA Synthesis Kit (Cat# AS-FS-004) for tRNA first strand cDNA synthesis, which is specifically optimized for and fully compatible with the nrStar[™] tRNA repertoire PCR Array. It can efficiently remove the modifications and greatly improve cDNA synthesis quality so that obtain more accurate tRNA expression data.

RNA demethylation

1. Prepare reagents

Gently thaw the Demethylation Reaction buffer $(5\times)$ and Nuclease-free water, and immediately place on ice. Mix by vortexing. Immediately before use, remove the Demethylase from the freezer, mix by flicking the tubes and place on ice. Spin down all reagents.

2. Combine reagents

Combine reagents orderly according to the following table. If performing RNA demethylation on multiple RNA samples, it is recommended to prepare an demethylation master mix of the Demethylation Reaction Buffer (5×), Demethylase and Nuclease-free Water (in the proportion indicated in Table). Considering the pipetting losses, 10% excel of all reagents is recommended.

Nuclease-Free Water	Variable
Demethylation Reaction buffer (5×)	40 µl
Demethylase	5 µl
RNase Inhibitor	1 µl
Input RNA	≤5 µg
Total volume	200 µl

3. Perform & stop RNA demethylation reaction

Incubate the above mix at 37°C for 100 min. Then orderly add 160 μ l Nuclease-free water and 40 μ l 5 \times Stop Buffer to terminate the reaction.

- 4. RNA precipitation
 - a. Add 400 µl phenol: chloroform to the sample. Mix well by inverting. Incubate at room temperature for 10 min. Centrifuge at 12,000 rpm for 10min.
 - b. Carefully pipette off top layer to RNase-free tube and discard bottom to phenol waste.
 - c. Add 400 µl chloroform to sample, mix well then microfuge at 12,000 rpm for 10 min.
 - d. Carefully pipette off top layer to RNase-free tube and discard bottom to phenol waste.
 - e. Add 1ml isopropanol to the aqueous phase. Mix well by inverting. Incubate at room temperature for 10 min. Centrifuge at 12,000 rpm for 10 min.
 - f. Remove the supernatant from the tube, leaving only the RNA pellet.
 - g. Add 1ml 75% ethanol (in DEPC-treated water). Mix well by inverting.
 - h. Centrifuge the tube at 7,500 rpm for 5min at 4°C. Discard the wash.
 - i. Vacuum or air dry the RNA pellet for 5–10min.
 - j. Resuspend the RNA pellet in 12 µl Nuclease-free Water.
 - k. Incubate in a water bath or heat block set at 55–60°C for 10–15 min.

First strand cDNA synthesis

NOTE: The recommended amount of starting material can vary from 10 pg to 5 μ g of total RNA according to the expression of interested RNA.

5. Prepare reagents

Gently thaw all of the kit components except for Reverse Transcriptase, and immediately place on ice. Mix by vortexing. Spin down all reagents.

NOTE: The first time to use this kit, please reconstitute the RNA spike-in by adding 200 μ l Nuclease-free water to the tube. Mix by vortexing and spin down. Leave on ice for 20 ~ 30 min to fully dissolve the RNA spike-in. Vortex again, then spin down.

6. Combine Annealing Mix according to Table

If performing first-strand cDNA synthesis on multiple RNA samples, it is recommended to prepare an

Annealing Mix of the Primer, dNTP Mix and RNA Spike-in (in the proportion indicated in Table). 10% excess volume for pipetting losses is recommended.

Random Primers	1 µl
dNTP Mix	1.5 μl
RNA Spike-in	1 µl
Template Total RNA	10.5 μl
Total volume	14.0 μl

- 7. Incubate in a thermal cycler at 65°C for 5 min, and then immediately place on ice for at least 1 min. Collect the contents of the tube by brief centrifugation.
- 8. Combine cDNA Synthesis Mix

cDNA Synthesis Mix is recommended to prepare for multiple RNA samples. It includes the components in the following table. 10% excess volume for pipetting losses is recommended.

RT Reaction buffer (5×)	4.0 μl
0.1 M DTT	1.0 μl
RNase Inhibitor	0.6 µl
Reverse Transcriptase	0.4 μl
Total volume	6.0 µl

- 9. Add cDNA Synthesis Mix to the tube from STEP 7. Vortex the sample briefly to mix, and collect by brief centrifugation. Incubate at 25°C for 5–10 min, followed by 50 min at 50°C
- 10. Terminate the reactions at 85°C for 5 min. Chill on ice.
- 11. OPTIONAL. To check the synthesized cDNA quality, reconstitute the RNA Spike-in qPCR Primer Mix in 200 μl nuclease-free water. Use 2 μl RNA Spike-in qPCR Primer Mix with 2 μl cDNA, 5 μl SYBR Green Master Mix, and 1 μl nuclease-free water. Run the PCR program described in "Running Real-Time PCR Detection" below. A Ct value < 30 for the RNA spike-in indicates a successful tRNA cDNA synthesis.</p>

NOTE: The cDNA synthesis product can proceed directly to PCR or can be stored at .20°C.

C. Perform qPCR for the PCR array

NOTE: The fellow operations are designed for one sample. If repetitive experiments are planned, the volume of the reagent should be accordingly increased. To make it easier to understand, we take Sample1 in Figure 2 for example.

 Dilute the cDNA in Nuclease-free water. If 1.5 µg input RNA is used with rtStar[™] tRNA-optimized First-Strand cDNA Synthesis Kit (Cat#AS-FS-004), the dilution factor is 1:20. Mix well and spin down. The diluted cDNA is used as the qPCR template in the wells for tRNA Repertoire Transcript assays, Internal Control References, and Spike-in External Controls.

- 2. For GDC Controls, combine 1.5 μl NRT (no RT) sample or 1.5 μl RNA sample, 7.5 μl SYBR Green Master Mix, and 6 μl Nuclease-free water. Mix well and spin down.
- 3. For Blank Controls, combine 20 μ L SYBR Green Master Mix and 20 μ L Nuclease-free water. Mix well and spin down.
- 4. Use Arraystar SYBR Green Real-Time Quantitative PCR Master Mix to prepare the qPCR Master Mix. There are total of 96 wells of PCR reaction. Some extra amount is included for consumption by the liquid dispensing operation. Prepare the cocktail according to the following table.

SYBR Green Master Mix	500 µl
diluted cDNA template	400 µl
ddH ₂ O	100 μl
total volume	1000 μl

5. Loading the PCR Array plate.

NOTE: In order to better understand the fellow operations, we take **Sample1** (in **Figure 2**) for example. If repetitive experiments are planned, it is important to note that the reagents should load to the related well number corroding to **Figure 2** and **Table 1**.

- a. CAREFULLY remove the plate seal from the PCR Array;
- Add 10 μl of the cocktail from STEP 4 to each PCR Array plate well (except No.93-No.96; i.e. well 017, well 019, well 021, well 023);
- c. Add10 µl GDC Mixture aliquot from STEP 2 into the No.95 (well O21) to detect genomic DNA contamination.
- Add 10 µl Blank Mixture aliquot from STEP 3 into the No.93 (well 017), No.94 (well 019) and No.96 (well 023).
- e. CAREFULLY but tightly seal the PCR Array plate with the optical adhesive cover. Be sure that no bubbles appear in any of the wells. To remove bubbles, tap the plate gently on the bench top or centrifuge the plate briefly.
- f. Keep the plate on ice while setting up the PCR program described in "Running Real-Time PCR Detection" below.
- 6. Running Real-Time PCR Detection

Cycles	Temperature	Time
1	95 ℃	10 minutes
10	95 ℃	10 seconds
40	60 °C	1 minutes
Melting curve analysis		

D. Data pre-processing and data analysis

Pre-process the qPCR run data according to the qPCR instrument manufacturer's instructions. If using baseline and threshold method for Ct calculations, such as with ABI 7900HT, the optimal baseline and threshold manual settings applied consistently across all assays on the plate are preferred over the software automatic settings for better reliability and accuracy.

Inspect the melting curve analysis of the post-PCR products to verify the amplification specificity. If the melting curve has multiple peaks or poor peak morphology deviated from that of the known single amplicon product, it may indicate non-specific off-target amplification or primer dimer formation, which will compromise the quantification. In such cases, gel electrophoresis or DNA fragment analysis can be performed to verify whether the product is from a single correct amplicon.

Export the raw Ct values to Excel. Free, easy-to-use data analysis software is available online, please refer to <u>www.arraystar.com</u> for detailed instruction. The data analysis procedures include:

Data pre-processing

1. Set all Ct values \geq 35 or N/A (not detected) to 35. From this point on, any Ct value equal to 35 is considered a negative result.

2. Examine the Ct values of the Genomic DNA Control (GDC) wells. If the value is greater than 35, no genomic DNA contamination is detected and no action is needed. If the value is less than 35, genomic DNA contamination is evident and the result may be compromised.

3. Before initiating the data analysis, the RNA spike in wells are compared. Outlier samples (Ct >25) may be identified and considered for exclusion in the further data analysis.

4. Inter-plate calibration (IPC) can be performed using the PPC assay replicates either with the data analysis software or manually. For each plate, verify that the replicates have Ct standard deviation ≤ 0.5 . If this is not the case, exclude the outlier if it can be identified. Calculate the average of the replicates for each plate, the overall average (average of IPC values from all plates). The calibration factor for each plate is calculated as the difference between the plate average and the overall average:

calibration factor = IPC(plate n) - IPC(overall)

The Ct value is corrected with the calibration factor as

 $Ct_{RNA} = Ct_{RNA}(Raw value, plate n) - IPC(plate n) + IPC(overall)$

 $Ct_{RNA} = Ct_{RNA}(Raw value, plate n) - calibration factor$

	Plate 1	Plate 2	Plate 3
Ala-TGC	20.26	20.83	20.44
IPC Plate average	16.70	16.70	16.40
IPC overall average	16.60	16.60	16.60

or

Calibration factor	0.10	0.10	-0.20	
Ala-TGC (Calibrated)	20.16	20.73	20.64	

5. Calculate the Δ Ct for each tRNA in the plate.

 $\Delta Ct_{RNA} = Ct_{RNA} - average(Ct_{Refs})$

Where average (Ct _{Refs}) is the average of the Ct values derived from the multiple reference genes. Three most stably expressed small ncRNA Control References were selected from abroad range of samples by our stringent algorithm that evaluates the optimal properties and the number for endogenous controls.

Data analysis

1. Calculate the $\Delta\!\Delta\text{Ct}$ for each tRNA

 $\Delta\Delta Ct = \Delta Ct(sample 1) - \Delta Ct(sample 2)$, between samples

or

 $\Delta\Delta Ct = \Delta Ct(group 1) - \Delta Ct(group 2)$, between groups

2. Calculate the fold changes for each gene from sample 1 to sample 2 or group 1 to group 2 as following:

Fold Change =
$$2^{-\Delta\Delta Ct}$$

NOTE: By convention, if the fold change is greater than 1, the result is reported as a fold up-regulation. If the fold change is less than 1, its negative inverse is reported as a fold down-regulation.

3. When comparing profiling differences between two groups (such as disease versus control) with biological replicates, the statistical significance of the difference can be estimated as p-value by *t*-test. RNAs having fold changes ≥ 2 and *p*-values ≤ 0.05 are selected as the significantly differentially expressed RNAs.

NOTE: Fold change is related to biological effect size. Ranking by fold change is preferred over *p*-value. qPCR as commonly used in confirmation has a limit of quantification of 0.5 Δ Ct, which is equivalent to approximately 2 fold change.

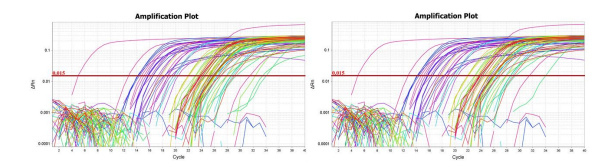
4. Other analyses such as scatter plots, volcano plots, list of differentially expressed genes and bar graph of expression differences for the tRNAs are performed and included in the standard analysis package.

III. Quality Control and Sample Data

A. nrStar[™] Human tRNA Repertoire PCR Array validation

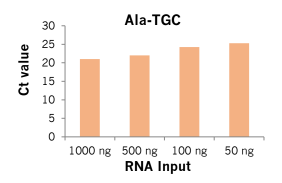
Real-time qPCR Validation

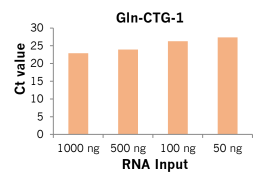
The performance of Human tRNA Repertoire Panel was tested using a cohort of pancreatic carcinoma and para-carcinoma tissues. The extracted RNA samples were converted to cDNA using rtStar™ tRNA-optimized First-Strand cDNA Synthesis Kit (Cat#AS·FS·004). The cDNA were profiled using the PCR array according to aforementioned protocol without modification. The results are charted on the real-time amplification plots for the entire plate for the cell lines.



Sensitivity Test

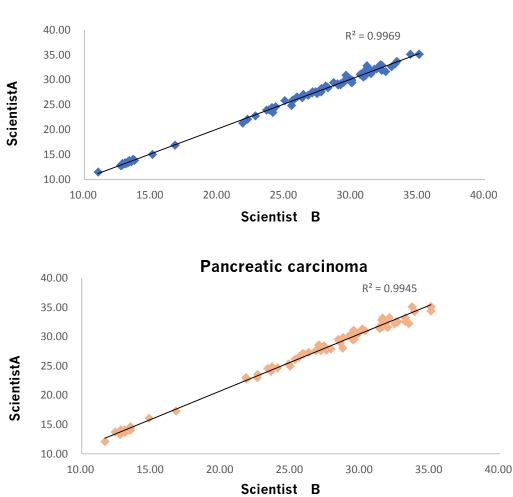
The decreasing input amounts of total RNAs from human glioblastoma cell lines were converted to cDNAs and profiled by the PCR arrays. The Ct values were determined using the software default automatic baseline and Ct settings. Ala-TGC and Gln-CTG-1 transcripts were detected at Ct values of 25.2 and 27.0 respectively, with the input RNA amount as low as 50 ng.





Reproducibility Test

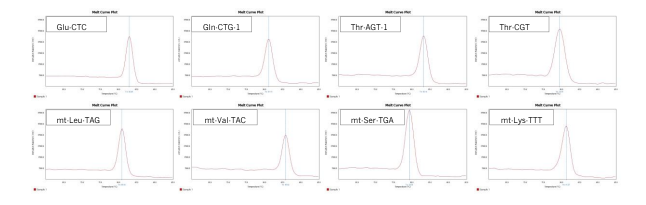
Two independent runs of nrStar[™] Human tRNA Repertoire PCR Array were conducted by two different scientists A and B at two different times using two different cells. The results demonstrate a high degree of reproducibility with correlation R²>0.98.



Para-carcinoma tissue

Specificity Test

The amplification products of transcripts of Glu-CTC, Gln-CTG-1, Thr-AGT-1, Thr-CGT, mt-Leu-TAG, mt-Val-TAC, mt-Ser-TGA and mt-Lys-TTT were analyzed by melting curves, all of which had single sharp melting peaks. The results demonstrate the high amplification specificities for the transcripts with the assays on the array.



B. Sample data: Analysis of human tRNA Repertoire transcripts levels in cell lines

The sample data were generated from RNAs extracted from pancreatic carcinoma and para-carcinoma tissues. The normalization was carried out using the average of the Internal Control Reference genes. The fold change between the pancreatic carcinoma and para-carcinoma tissues are graphed in the bar chart below.



IV. Troubleshooting

Problem	Possible solution
qPCR background too high	Reduce the amount of cDNA used in the SYBR Green Master Mix.
	Inspect if the Internal Controls have valid qPCR signal
No qPCR signals	Set SYBR Green as the Detector's Reporter Dye
	Use more cDNA in the Master Mix
	• Lower the annealing temperature in Protocol STEP C.6 from 60°C to 50°C.
	 Follow the instructions of the qPCR system manufacturer.
Baseline and threshold settings	Contact their technical support as necessary.

V. References

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VI. Technical Support

For additional information, manual download or technical assistance, please visit our website at www.arraystar.com, or contact us at:

Arraystar Inc. 9430 Key West Ave #128 Rockville, MD 20850, USA

Tel: 888-416-6343 Fax: 240-238-9860 Email: support@arraystar.com

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By purchasing and using any part of the nrStar™ Human tRNA Repertoire PCR Array, you agree to accept the following terms and conditions.

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