Melt curve analysis for interpretation of MicroRNAs’ gene expression data in foods

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Abstract
This article illustrates the importance of melt curve analysis (MCA) in interpretation of mild nutrigenomic micro(mi)RNA expression data, by measuring the magnitude of the expression of key miRNA molecules in stool of healthy human adults as molecular markers, following the intake of Pomegranate juice (PGJ), functional fermented sobya (FS), rich in potential probiotic lactobacilli, or their combination. We elaborate on these data and present a detailed review on use of melt curves for analyzing nutrigenomic miRNA expression data, which initially appear to show no significant expressions, but are actually more subtle than this simplistic view, necessitating the understanding of the role of MCA for a comprehensive understanding of what the collective expression and MCA data collectively imply.

Introduction
Gene expression and its control by miRNAs

Cell’s gene expression profile determines its function, phenotype and cells’ response to external stimuli, and thus help elucidate various cellular functions, biochemical pathways and regulatory mechanisms [1]. Several gene expression profiling methods at the RNA level have emerged during past years and have been successfully applied to cancer research. Profiling by microarrays allows for the parallel quantification of thousands of genes from multiple samples simultaneously, using a single RNA preparation, and has become valuable because microarrays provide a comprehensive approach to characterize cancer molecularly, as seen in studies on colon cancer [1].

Control of gene expression has been studied by miRNA molecules, a small non-coding RNA molecule (18–24 nt long), involved in transcriptional and post-transcriptional regulation of gene expression by inhibiting gene translation. MiRNAs silence gene expression through inhibiting mRNA translation to protein, or by enhancing the degradation of mRNA. Since first reported in 1993 [2], the number of identified miRNAs in June 2014, version 14.0, the latest miRBase release (v20) [3] contains 24,521 miRNA loci from 206 species, processed to produce 30,424 mature miRNA products. MiRNAs are processed by RNA polymerase II to form a precursor step which is a long primary transcript. Pri-miR is converted to miRNA by sequential cutting with two enzymes belonging to a class of RNA III endonucleases, Drosha and Dicer. Drosha converts the long primary transcripts to ~70 nt long primary miRNAs (pri-miR), which migrate to the cytoplasm by Exportin 5, and converted to mature miRNA (~22 net) by Dicer [4]. Each miRNA may control multiple genes, and one or more miRNAs regulate a large proportion of human protein-coding genes, whereas each single gene may be regulated by multiple miRNAs [5]. MiRNAs inhibit gene expression through interaction with 3-untranslated regions (3 UTRs) of target mRNAs carrying complementary sequences [4,5].

Effect of antioxidant polyphenols --abundant in Mediterranean diets-- on gene expression unraveled by the availability of molecular biology techniques, reveals our adaptation to environmental changes [6]. Efforts to study the human transcriptome have collectively been applied to tissue, blood, and urine (i.e., normally sterile materials), as well as stool (a non-sterile medium). Extraction protocols that employ commercial reagents to obtain high-yield, reverse-transcribable (RT) PCR, SNPs

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RNA from human stool in studies performed on colon cancer have been reported [7,8].

Micro(mi)RNAs as biomarkers, and their roles in disease processes

A biomarker is believed to be a characteristic indicator of normal biological processes, pathogenic processes, or pharmacological responses to therapeutic interventions. In contrast, clinical endpoints are considered as variables representing a study subject’s health from his/her perspective [9]. A variety of biomarkers exist today as surrogates to access clinical outcomes in diseases, predict the health of individuals, or improve drug development. An ideal biomarker should be safe and easily measured, is cost effective to follow up, is modifiable with treatment, and is consistent across genders and various ethnic groups. Because we never have a complete understanding of all processes affecting individual’s health, biomarkers need to be constantly reevaluated for their relationship between surrogate endpoints and true clinical endpoints [10]. miRNAs have been used herein as biomarkers for assessing the effect of intake of PP-rich or fermented foods on the expression of 88 miRNA genes known to influence cancer.

Disease modulation by nutrients

Cardiovascular diseases due to hypercholesterolemia are considered a risk factor for Chronic Heart Disease (CHD), and chronic degenerative diseases--caused wholly or partially by dietary patterns--represent the most serious threat to public health [11,12]. Moreover, nearly one-third of all cancer deaths are due to poor nutrition, lack of physical activity, and obesity; and these risk factors account for nearly 80% of large intestine, breast, and prostate cancers. Chronic inflammation is considered a common factor that contributes to the development and progression of these illnesses, which are caused by and/or modified by diet [13].

Pomegranate juice (PGJ) and derived products are considered the richest sources of polyphenolic compounds, with positive implication on TC, LDL-C and TG plasma lipid profile [14]. Moreover, anthocyanin and ellagittannins pigments, mainly punicagins, inhibit the activities of enzymes 3-hydroxy-3-methylglutaryl-CoA reductase and 3-hydroxyacyltransferase, important in cholesterol metabolism [15]. Probiotic bacteria also contribute to lowering plasma hypercholesterolemia due to the above mechanism, caused by the probiotic bile salt hydrolyase (BSH) activity. This probiotic enzyme hydrolyses conjugates both glycodeoxycholic and taurodeoxycholic acids to hydrolysis products, inhibiting cholesterol absorption and decreasing reabsorption of bile acid [16].

Colonic microbiota is a central site for the metabolism of dietary PP and colonization of probiotic bacteria. A dietary intervention study with probiotic strains from three Lactobacillus species (L. acidophilus, L. casei and L. rhamnosus) given to healthy adults, showed that bacterial consumption caused the differential expression of from hundreds to thousands of genes in vivo in the human mucosa [17]. The interaction of PP with the gut microbiota influences the expression of some human genes (i.e., nutritional transcriptomics), which mediates mechanisms underlying their beneficial effects [17]. Similar in vivo mucosal transcriptome findings have been reported when adults were given the probiotic L. plantarum, illustrating how probiotics modulate human cellular pathways, and show remarkable similarity to responses obtained for certain bioactive molecules and drugs [18].

Materials and methods

Participants

Study subjects were 25 healthy adults, 20 to 34 years old; exclusion were: absence of metabolic diseases, no use of medication for the last 6 weeks, and no signs of allergy or hypersensitivity to food or ingested material. Compliance with the supplementation in all subjects was satisfactory, as assessed daily, and, all subjects continued their habitual diets throughout the study. The research protocol was approved by the institution review board, and all subjects gave written consent prior to their participation in the study.

Design of the study

Figure 1 shows the design of the nutrigenomic randomized study. Estimated dietary intake was assessed by 3 repeated food records, one week before they were enrolled in the trial. The average portion size consumed, as well as composition data values from nutrient composition of the food were combined to assess average daily energy and nutrient intakes by the “nutrisurvey” software program. The characteristics of the voluntary subjects who were enrolled in the study, the mean daily energy intake, as well as selected macro nutrients are presented in Table 1.

Supplements

Pomegranate was obtained in bulk from the Obour Public Market, Cairo, Egypt. Pomegranate fruits were peeled and the juice was extracted using a laboratory pilot press (Braun, Germany). The juice was distributed in aliquots of 100 or 250 grams air tight, light-proof polyethylene bottles, and frozen at −20°C, where pomegranate polyphenols remained stable. Sour sobya, a fermented rice porridge containing per gram 3 × 10^7 cfu diverse lactic acid bacteria (LAB) and 1 × 10^5 cfu Saccharomyces cerevisiae, with added ingredients such as milk, sugar and grated coconut, was purchased twice a week from the retail market, and saved in the refrigerator. Sobya is fermented rice. Table 2 illustrates the proximate initial and final mean urinary polyphenols,

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**Table 1. Composition of the supplements**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Unit</th>
<th>Control (Portion served)</th>
<th>FS (Portion served)</th>
<th>PGJ (Portion served)</th>
<th>FS+PGJ (Portion served)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Portion Size</td>
<td>g</td>
<td>–</td>
<td>170</td>
<td>250</td>
<td>150+10</td>
</tr>
<tr>
<td>Total Solids</td>
<td>g</td>
<td>–</td>
<td>40.01</td>
<td>17.75</td>
<td>48</td>
</tr>
<tr>
<td>Carbohydrate</td>
<td>g</td>
<td>–</td>
<td>51.1</td>
<td>32.75</td>
<td>59</td>
</tr>
<tr>
<td>Dietary Fiber</td>
<td>g</td>
<td>–</td>
<td>54</td>
<td>0.25</td>
<td>48.6</td>
</tr>
<tr>
<td>Energy</td>
<td>kcal</td>
<td>–</td>
<td>263</td>
<td>135</td>
<td>290</td>
</tr>
<tr>
<td>Lactobacillus</td>
<td>diverse cfu/ serving size</td>
<td>–</td>
<td>5.1 x 10^6</td>
<td>4.5 x 10^6</td>
<td></td>
</tr>
<tr>
<td>Yeast</td>
<td>cfu/serving size</td>
<td>–</td>
<td>2.77 x 10^8</td>
<td>2.44 x 10^8</td>
<td></td>
</tr>
<tr>
<td>Total PP</td>
<td>mg/portion g</td>
<td>–</td>
<td>519.1±8.75</td>
<td>207.65±3.5</td>
<td></td>
</tr>
<tr>
<td>Antioxidant activity</td>
<td>(AEAC)**</td>
<td>–</td>
<td>7.74±1.33</td>
<td>11.35±2.2</td>
<td>11.37±2.2</td>
</tr>
</tbody>
</table>

*mg galic acid equivalent (GAE), FS: Functional fermented sobya; PGJ: Pomegranate juice **mmol ascorbic acid equivalent antioxidant activity [5]

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plasma and urinary antioxidative activity, urinary thiobarbituric acid reactive species (TBARS), and erythrocytic glutathione-S-transferase (GST).

**Stool collection and storage**

Stool was obtained from the 25 healthy adults, twice at day 0 and three weeks after the dietary intervention. All stools were collected with sterile, disposable wood spatulas in clean containers, after stools were freshly passed, and then placed for storage into Nalgene screw top vials (Thermo Fisher Scientific, Inc., Palo Alto, CA, USA), each containing 2 ml of the preservative RNA later (Applied Biosystems/Ambion, Austin, TX, USA), which prevents the fragmentation of the fragile mRNA molecule [7], and vials were stored at ~ 70 °C until samples were ready for further analysis. Total small RNA, containing miRNAs, was extracted from all frozen samples at once, when ready, and there was no need to separate mRNA containing small miRNAs from total RNA, as small total RNA was suitable to make ss miRNA c-DNA.

**Extraction of total small RNA**

A procedure used for extracting small total RNA from stool was carried out using a guanidiunium-based buffer, which comes with the RNeasy isolation kit, Qiagen, Valencia, CA, USA, as we have previously detailed [7]. DNase digestion was not carried out, as our earlier work demonstrated no difference in RNA yield or effect on RT-PCR after DNase digestion [7]. The time to purify aqueous RNA from all of the 25 frozen stool samples was ~ three hours. Small RNA concentrations were measured spectrophotometrically at λ 260 nm, 280 nm and 230nm, using a NanoDrop spectrophotometer (Thermo-Fischer Scientific). The integrity of total RNA was determined by an Agilent 2100 Bioanalyzer (Agilent Technologies, Inc., Palo Alto, CA, USA) utilizing the RNA 6000Nano LabChip®. RNA integrity number (RIN) was computed for each sample using instrument’s software [7].

**Preparation of ss-cDNA for molecular analysis**

The Rt² miRNA First Strand Kit® from SBAbiosciences Corporation (Frederick, MD, USA) was employed for making a copy of ss-DNA in a 10.0 µl reverse transcription (RT) reaction, for each RNA samples in a sterile PCR tube, containing 100 ng total RNA, 1.0 µl miRNA RT primer & ERC mix, 2.0 /µl 5X miRNA RT buffer, 1.0 µl miRNA RT enzyme mix, 1.0 µl nucleotide mix and Rnase-free H₂O to a final volume of 10.0 µl. The same amount of total RNA was used for each sample. Contents were gently mixed with a pipettor, followed by brief centrifugation. All tubes were then incubated for 2 hours at 37°C, followed by heating at 95°C for 5 minutes to degrade the RNA and inactivate the RT. All tubes were chilled on ice for 5 minutes, and 90 µl of Rnase-free H₂O was added to each tube. Finished miRNA First Strand cDNA synthesis reactions were then stored overnight at -20°C [7].

**Use of cancer RT² miRNA PCR array 96-well plate to study miRNAs’ expressions**

We used a SBAbiosciences RT² miRNA qPCR Array Plate System for Human (Qiagen) to analyze miRNA expression using real-time, reverse transcription PCR (RT-qPCR) as a sensitive and reliable quantitative method for miRNA expression analysis. The arrays employ a SYBR Green real-time PCR detection system, which has been optimized to analyze the expression of many mature miRNAs simultaneously. Each 96-well array plate contains a panel of primer sets for 88 relevant miRNA focused pathways (one universal primer and one gene-specific primer for each miRNA sequence), plus four housekeeping genes (Human SNORD 48, 47 and 44, and U6), and two RNA and two PCR quality controls. Duplicate RT Controls (RTC) to test the efficiency of the miRNA RT reaction, with a primer set that detects the template synthesized from the built in miRNA External RNA Control (ERC). There are duplicate RT controls (RTC) to test the efficiency of the miRNA RT process, with a primer set to detect the template synthesized from the kit’s built-in miRNA External RNA Control (ERC). There is also duplicate positive PCR controls (PPC) to test the efficiency of the PCR process, using a per-dispersed artificial DNA sequence and the primer set that detects it. The two sets of duplicate control wells (RTC and PPC) also test for inter-well, intra-plate consistency. The human RT² miRNA PCR Arrays reflect miRNA sequences annotated by the Sanger miRBase Release 14. Figure 2 shows the layout of the MAH-102F array.

**Performing real-time quantitative polymerase chain reaction (qPCR)**

We used RT² SYBR Green qPCR Master Mix (SBA Biosciences) to obtain accurate results from our qPCR arrays. The following components were mixed in a 15-ml tube for 96-well plate format: 1275 µl of 2X RT² SYBR Green PCR Master Mix, 100 µl of diluted first strand reaction, 1175 µl of ddH₂O (total volume 2550 µl, of which 2400 was needed for 96 reactions, each well having 25 µl, with150 µl cocktail remaining.

We employed a Roche LightCycler 480® 96-well block PCR Machine (Roche, Mannheim, Germany) to carry out quantitative real-time miRNA expressions. When ready, we removed the needed miRNA qPCR Arrays, each wrapped in aluminum foil, from their sealed bags, added 25 µl of the same cocktail to each well, adjusted the ramp rate to 1°C/sec. We used 45 cycles in the program, and employed the Second Derivative Maximum method, available with the LightCycler 480® software for data analysis [19]. We first heated the 96 well plate for 10 min at 95°C to activate the HotStart DNA polymerase, then used a three-step cycling program (a 15 seconds heating at 95°C to separate the ds DNA, a 30 seconds annealing step at 60°C to detect and record

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**Table 2. Initial and final mean urinary polyphenols, plasma and urinary antioxidative activity, urinary TBARS and erythrocytic GST**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Unit</th>
<th>Control</th>
<th>Sobya</th>
<th>Pomegranate</th>
<th>Sobya +Pomegranate</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>X±SE</td>
<td>X±SE</td>
<td>X±SE</td>
<td>X±SE</td>
</tr>
<tr>
<td>Urinary polyphenol</td>
<td>GAE/mg creat</td>
<td>10.36±1.8</td>
<td>8.11±2.2</td>
<td>11.84±6.2</td>
<td>9.86±1.8</td>
</tr>
<tr>
<td>Urinary antioxidant activity</td>
<td>AEAC/mg creat*</td>
<td>9.74±2.0</td>
<td>8.13±2.7</td>
<td>3.89±0.9</td>
<td>10.30±2.3</td>
</tr>
<tr>
<td>Urinary TBARS</td>
<td>ug/mg creat</td>
<td>83.04±12.1</td>
<td>75.17±15.3</td>
<td>82.77±27.8</td>
<td>29.97±4.4</td>
</tr>
<tr>
<td>Plasma antioxidant</td>
<td>AEAC/100 ml</td>
<td>6.36±2.81</td>
<td>5.99±2.66</td>
<td>3.70±0.33</td>
<td>4.55±0.27</td>
</tr>
<tr>
<td>E-GST activity</td>
<td>IU/g Hb</td>
<td>5.94±3.3</td>
<td>5.45±4.1</td>
<td>4.26±0.5</td>
<td>7.21±0.8</td>
</tr>
</tbody>
</table>

X±SE: Mean ± Standard Error, *:mmol ascorbic acid equivalent antioxidant capacity/mg creatinine; GAE: gallic acid equivalent, Mean values are significantly different if the P-values are less than 0.05 (P<0.05) [5]
SYBR Green fluorescence at each well during each cycle, and a final heating step for 30 seconds at 72°C. Each plate was visually inspected after the run for signs of evaporation from the wells. Data were analyzed using the 2-ΔΔct method [20]. Resulting threshold cycle values for all wells were exported to a blank Excel sheet for analysis. We also ran a Dissociation (Melt) Curve Program after the cycling program [21] and generated a first derivative dissociation curve for each well in the plate, using the LC (Lightcycler’s) software.

**Statistical and bioinformatics analysis**

Gene expressions were standardized by dividing the SNORD48 value while raw melting temperatures were used. Analysis were done using the software R (version 3.1.3), with the package MASS [22]. One individual had so many missing values that this case was not used in the analysis so that the number of individuals is 24. For each standardized gene and each melting temperature, a one-way ANOVA was used to obtain a p-value. There were four levels of the explanatory variable: Control, Sobya, Pom, and Both. Parallel coordinate plots (parcoord command in R) [23] were used to visualize the data for each gene and each melting temperature. Coordinates were ordered using the magnitude of the p-value. The two sample t-test was used on gene expression to compare control to sobya and control to Both. -values are greater (less significant), and the graphs did not show any meaningful separations.

Stool represents a challenging environment, as it contains many substances that may not be consistently removed in PCR, in addition to the existence of certain inhibitors, which all must be removed for a successful PCR reaction. Our results [27] and others [8,28] have shown that the presence of non-transformed RNA and other substances in stool do not interfere with measuring miRNA expressions, because of the use of suitable PCR primers, and the robustness of the real-time qPCR method [19]. Stool colonocytes contain much more miRNA and mRNA than that available in free circulation, as in plasma [29], all factors that facilitate accurate and quantitative measurements.

**Discussion**

**Suitability of stool as a medium for developing a sensitive molecular biomarker screen**

Stool represents a challenging environment, as it contains many substances that may not be consistently removed in PCR, in addition to the existence of certain inhibitors, which all must be removed for a successful PCR reaction. Our results [27] and others [8,28] have shown that the presence of non-transformed RNA and other substances in stool do not interfere with measuring miRNA expressions, because of the use of suitable PCR primers, and the robustness of the real-time qPCR method [19]. Besides, stool colonocytes contain much more miRNA and mRNA than that available in free circulation, as in plasma [29], all factors that facilitate accurate and quantitative measurements.
presence of inhibiting substances, representing diverse compounds with different properties and mechanisms of action, which induce their effects by direct interaction with DNA that will be amplified, or through interference with the employed thermostable DNA polymerase [30].

Agents that reduce Mg2+ availability or interfere with the binding of Mg2+ to DNA polymerase could inhibit the PCR reaction [31]. Calcium ion is another inorganic inhibiting substance, although most PCR inhibitors are organic compounds (e.g., bile salts, sodium dodecyl sulphate, urea, phenol, ethanol, polysaccharides), as well as proteins (e.g., collagen, hemoglobin, immunoglobin G and protease) [32]. The existence of polysaccharides in stool could decrease the capacity to resuspend precipitated RNA or disrupt the enzymatic reaction by mimicking the structure of nucleic acid. The DNA template of the PCR, as well as primers binding to DNA template can be inhibited by inhibitors in stool, such as additional extraction steps, sephadex G-200 chromatography, heat treatment before the PCR, chloroform extraction, treatment with activated carbon, adding BSA, or dilution of sample, have been suggested [34]. We found the dilution method, in which the extracted ribonucleic acid (RNA) is diluted in the reaction mixture with distilled water or an isotonic buffer, to be the most practical method for preventing PCR inhibition using a commercially available diluent [35].

Role of biomarker miRNAs in various diseases

MiRNA functions were shown to regulate development [36] and apoptosis [37], and dysregulation of miRNAs has been associated with many diseases such as various cancers [38], heart diseases [39], kidney diseases [40], nervous system diseases [41], alcoholism [42], obesity [43], auditory diseases [44], eye diseases [45], skeletal growth defects [46], as well as key role in host–virus pathogenesis of viral diseases [47]. A negative correlation was found between tissue specificity of interactions and miRNA in a number of diseases, and an association between miRNA conservation and disease, and predefined miRNA groups allow for identification of novel disease biomarkers at the miRNA level [48]. Specific miRNAs are crucial in oncogenesis [49], effective in classifying solid [50] and liquid tumors [51], and function as oncogenes or tumor suppressor genes [52]. MiRNA genes are often located at fragile sites, as well as minimal regions of loss of heterozygosity, or amplification of common breakpoints regions, suggesting their involvement in carcinogenesis [53]. MiRNAs have shown to serve as biomarkers for cancer diagnosis, prognosis and/or response to therapy [54,55]. Profiles of miRNA expression differ more accurately than expression profiles of protein-coding messenger (m)RNA genes [56]. Besides, small miRNAs (~18-22 nt long) are stable molecules than the fragile mRNA [27].

Melting curve analysis (MCA)

MCA is an assessment of dissociation characteristics of dsDNA during heating, leading to rise in absorbance, intensity and hyperchromicity. The temperature at which 50% of DNA is denatured is referred to as melting point, \( T_m \).

Gathered information can be used to infer the presence of single nucleotide polymorphism (SNP), as well as clues to molecule's mode of interaction with DNA, such as intercalator slots in between base pairs through pi stacking and increasing salt concentration, leading to rise in melt temperature, whereas pH can affect DNA's stability, leading to lowering of its melting temperature [57]. Originally, strand dissociation was measured using UV absorbency, but now techniques based on fluorescence measurements using DNA intercalating fluorophores such as SYBR Green I, EvaGreen, or Fluorophore-labelled DNA probes (FRET probes) when they are bound to ds DNA [58] are now common. Specialized thermal cyclers that run the qPCR, such as Roche LightCycler (LC) 480, used in this study, is programmed to produce the melt curve after the amplification cycles are completed. As the temperature increases, dsDNA denatures becoming ss and the dye dissociates, resulting in decrease in fluorescence. The graph of the negative first derivative of the melting-curve (-dF/dT) represents the rate of change of fluorescence in the amplification reaction and allows pin-pointing the temperature of dissociation (50% dissociation) using formed peaks to obviate or complement sequencing efforts [59].
The melting temperature ($T_m$) of each product is defined as the temperature at which the corresponding peak maximum occurs. The MCA confirms the specificity of the chosen primers, as well as reveals the presence of primer-dimers, which usually melt at lower temperatures than the desired product, because of their small size, and their presence severely reduce the amplification efficiency of the target gene as they compete for reaction components during amplification, and ultimately the accuracy of the data. The greatest effect is observed at the lowest concentrations of DNA, which ultimately compromises the dynamic range. Moreover, nonspecific amplifications may result in PCR products that melt at temperatures above or below that of the desired product. Optimizing reaction components ($Mg^{2+}$, detergents, SYBR Green I concentration) and annealing temperatures aid in decreasing nonspecific product formation. Adequate product design, however, is considered to be the best method to avoid nonspecific products’ formation. Including a negative control will determine if there is a coamplified genomic DNA [57,58]. The formula for $T_m$ calculation is shown by the equation:

$$T_m = \frac{-\Delta H^o_{\text{add}}}{\Delta S^o_{\text{add}}} + \frac{R \ln C_T}{\Delta S^o_{\text{add}}} + 273.15,$$

where thermodynamic parameter $\Delta H^o_{\text{add}}$ is Enthalpy changes, $\Delta S^o_{\text{add}}$ parameter is Entropy changes, and $C_T$ is total strand concentration; these free-energy parameters predict $T_m$ of most oligonucleotide duplexes to within $5\degree C$ and permit prediction of DNA, as well as RNA duplex stabilities. It should be noted that $T_m$ depends on the conditions of the experiment, such as oligonucleotide concentration, salts’ concentration, mismatches and single nucleotide polymorphisms (SNPs) [60]. OligoAnalyzer ® Tool [www.idtdna.com/analyzer/Applications/Oligoanalyzer] allows for calculating the $T_m$ of employed nucleotides.

Microscale thermophoresis is a method that determines the stability, length, conformation and modifications of DNA and RNA. It relies on the directed movement of molecules in a temperature gradient that depends on surface characteristics of the molecule, such as size, charge and hydrophobicity. By measuring thermophoresis of nucleic acids over a temperature gradient, one finds clear melting transitions, and can resolve intermediate conformational states (Figure 3). These intermediate states are indicated by an additional peak in the thermophoretic signal preceding most melting transitions (Figure 3B) [59,61-63]. Agarose gel visualization is the gold standard for analyzing PCR products. Alternatively to reduce the number of gels needed to conform the presence of a single amplicon, “uMelt” melting curve prediction software (http://www.dna.uth.edu/umelt/umelt.html) can be used to confirm that a single amplicon is generated by PCR [64]. This program predicts melt curves and their derivatives for qPCR-length amplicons and is suited to test for multiple peaks in a single amplicon product.

...Because SYBR Green I dye has several limitations, including inhibition of PCR, preferential binding to CG-rich sequences and effects on MCA, two intercalating dyes SYTO-13 and SYTO-82 were tried and did not show these negative effects, and SYTO-82 demonstrated a 50-fold lower detection limit [65], as well as best combinations of time-to-threshold ($T_\alpha$) and signal-to-noise ratio (SNR) [66]. To optimize performance of the buffer, a PCR mix supplemented with two additives, 1M 1,2-propanediol and 0.2 M trehalose, were shown to decrease $T_\alpha$, efficiently neutralize PCR inhibitors, and increase the robustness and performance of qPCR with short amplicons [67]. “uAnalyze®” is another web-based tool, similar to uMELT, for analyzing high-resolution melting PCR products’ data, in which recursive nearest neighbor thermodynamic calculations are used to predict a melt curve. Using 14 amplics of CYBB [cytochrome b-245 heavy chain, also known as cytochromae b (558) subunit], the main +/- standard deviation, the difference between experimental and predicted fluorescence at 50% helicity was -0.04 +/- 0.48°C [68].

MCA has been an effective and economical way for identification of virus stains [69], genes [70], bacterial strains [71], insect species [72], temperature validation of PCR cyclers [73], detection of translocations in lymphomas [74] and RNA interference/gene silencing [61]. Thus, the presence of double peaks during MCA, is not always indicative of non-specific amplification, and other methods such as agarose gel electrophoresis, and use of melt curve prediction software [61,68] are also needed in order to determine the purity of an amplicon. For example, Figure 5A shows a single peak for exon 17b of CFTR (Cystic Fibrosis Transmembrane Conductance Regulator) gene, whereas the melt curve for an amplicon from exon 7 of CFTR shows two peaks, which could be interpreted as indicative of two separate amplicons (Figure 5B). However, analysis by agarose gel electrophoresis showed only one peak. To solve this conflict, an understanding of how melt curves are produced is needed. It should be emphasized that intercalating dyes used in qPCR, such as SYBR Green, will fluoresce only when the dye is bound to ds DNA, but not in the presence of a ssDNA, or when the DNA is free in solution. After the amplification cycle in qPCR, the instrument starts at a preset temperature above the primer $T_m$, and as the temperature increases dsDNA denatures becoming ssDNA and the dye therefore dissociates from the ssDNA (Figure 3A). The change in slope of this curve when plotted as a function of temperature to obtain a melt curve for CFTR exon 17b (Figure 4A). However, if we allow for the possibility that DNA may assume an intermediate state that is neither dsDNA or ssDNA, raw date from CFTR exon 7 melt will look Figure 4B. This could happen when there are regions of the amplicon that are more stable (e.g., G/C rich), which do not melt immediately, but maintain their ds configuration until the temperature becomes sufficiently high to melt it, which results in two phases (Figure 4B). Additional sequence factors, such as amplicon misalignment in A/T rich regions, and designs that have secondary structure in the amplicon region, can also produce products that melt in multiple phases.

An advancement of MCA, referred to as High Resolution Melt (HRM), discovered and developed by Idaho Technology and the University of Utah [75, http://www.dna.uth.edu/Hi-Res/TOP_Hi-Res%20Melting.html], which has been useful for mutation detection and SNPs, enabling differentiation of homozygous wildtype, heterozygous and homozygous mutant alleles from the dissociation patterns. HRM has been used to identify variation in nucleic acid sequences, enabled by use of a more advanced software, and is therefore less expensive than probe-based genotyping methods, and allows for identification of variants quickly and accurately [76]. This method has been widely used in molecular diagnosis and for detection of mutations [77-79].

In our study, we found the melt curve analysis to be a useful and an informative method because after the statistical analysis carried on our miRNA expression samples showed no preferential expression of any of the 88 miRNA genes, a melt curve analysis on the same samples found that we could distinguish 7 miRNA (miR-184, miR-203, miR-373, miR-124, miR-96, miR-373 and miR-301a), due to different separation melting profiles (Figure 6). Thus, we believe that it is imperative for investigators to run this kind of analysis on samples that particularly may not show expression differences in their miRNA or miRNA studied genes, such as nutritional samples.
Bioinformatic methods to correlate seed miRNA Data with messenger(m)RNA Data

To provide information about complex regulatory elements, we correlated miRNA results with our available available mRNA data [80], as well as those data available in the open literature using computer model TargetScan [81,82]. The authenticity of functional miRNA/mRNA target pair once identified was validated by fulfilling four basic criteria: a) miRNA/mRNA target interaction can be verified, b) the predicted miRNA and mRNA target genes are co-expressed, c) a given miRNA must have a predictable effect on target protein expression, and d) miRNA-mediated regulation of target gene expression should equate to altered biological function. Bioinformatics showed 21 upregulated mRNA genes encoding different cell regulatory functions, and 12 of these mRNAs were found to be active in the nucleus and related to transcriptional control of gene regulation. For down-regulated miRNAs, four of the mRNAs appeared to be clustered in cell cycle regulation categories (Table 3) [27].

Clinical significance

The clinical significance of the study presented above is that using melting temperatures for analyzing nutrient-gene data is a promising new approach for identifying key regulatory miRNA genes related to metabolites rich in polyphenols, probiotic lactobacilli, or combinations...
of the two metabolites. Melt curve analysis is a powerful novel approach because after the statistical analysis carried on our miRNA samples produced negative gene expression (Cq) results, running melt curve analysis on the same samples identified 7 of the 88 miRNA genes imprinted on the highly sensitive focused PCR arrays (~ 8% of the genes), and using parallel coordinates plots showed noticeable separation of melt curve profiles. Thus, we believe that it is imperative for investigators to run this kind of MCA on nutrition samples that are mild in nature, and many not always show significant differences in the expression of studied miRNA genes. The same analysis can also be envisioned for messenger mRNA amplifications, using mRNA arrays, and then using bioinformatics resources to correlate mRNA with miRNA data.

We are also planning to validate these initial results by carrying out additional miRNA nutrigenomic expression studies, with much more observations using PP, FS and their combinations, and collectively the obtained results would fully demonstrate the sensitivity/specificity of this powerful systemic molecular approach for analyzing nutrient-gene data.

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Disclosure

Authors certify that there has been no conflict of interest in carrying out this research, interpretation of data, or in preparing this manuscript for publication.

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