Methodology Report

A Multisampling Reporter System for Monitoring MicroRNA Activity in the Same Population of Cells

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MicroRNAs (miRNAs) downregulate gene expression by binding to the partially complementary sites in the 3′ untranslated region (UTR) of target mRNAs. Several methods, such as Northern blot analysis, quantitative real-time RT-PCR, microarray, and the luciferase reporter system, are commonly used to quantify the relative level or activity of miRNAs. The disadvantage of these methods is the requirement for cell lysis, which means that several sets of wells/dishes of cells must be prepared to monitor changes in miRNA activity in time-course studies. In this study, we developed a multisampling reporter system in which two secretable bioluminescence-generating enzymes are employed, one as a reporter and the other as an internal control. The reporters consist of a pair of vectors containing the Metridia luciferase gene, one with and one without a duplicated miRNA targeting sequence at their 3′UTR, while the other vector coding for the secreted alkaline phosphatase gene is used as an internal control. This method allows miRNA activity to be monitored within the same population of cells over time by withdrawing aliquots of the culture medium. The practicability and benefits of this system are addressed in this report.

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1. Introduction

MicroRNAs (miRNAs) are small noncoding RNAs found in plants and animals which downregulate gene expression by binding to complementary sites on the 3′ untranslated region (UTR) of target mRNAs. As a result, the targeted miRNAs are degraded if the miRNA and target sequence are a perfect match or their translation is inhibited [1]. Over 800 different miRNAs are found in the human genome (miRBase: http://microrna.sanger.ac.uk/; Sources of predicted miRNA target: http://microrna.org), and each miRNA may have hundreds of target mRNAs. For this reason, miRNAs have profound effects on global gene expression and are known to play roles in various cellular processes, such as development, differentiation, proliferation, and apoptosis [2–4]. A number of miRNAs have been linked to human cancer [1, 5].

Several methods are used to detect the relative level or activity of miRNAs, including Northern blot analysis [6], quantitative real-time RT-PCR [7], microarray [6, 8], and the firefly luciferase reporter system [7, 9]. To circumvent the tediousness and accuracy-compromising drawbacks of the lysis of the test cells that is required in all the above methods, in this study, we developed a multisampling reporter system that makes it possible to monitor changes in miRNA activity over time within the same population of cells by sampling the culture medium, thus avoiding the need for a lysis step. Briefly, the system contains two distinct vectors that encode different secretable bioluminescence-generating enzymes. The reporters consist of a pair of vectors containing the Metridia luciferase (MLuc) gene with or without a duplicated miRNA targeting sequence at their 3′UTR, while the other vector coding for the secreted alkaline phosphatase (SEAP) gene is used as an internal control. This method allows miRNA activity to be monitored within the same population of cells over time by withdrawing aliquots of the culture medium. The practicability and benefits of this system are addressed in this report.
luminescence [12]. Taking advantage of these features, we can sample small aliquots of medium over time and keep the monitored cells intact. In this study, this system was used to monitor the activity of miR-15/miR-16, which may downregulate the expression of BCL-2, an antiapoptotic protein [13, 14], by binding to their target sequence in the 3’ UTR of the Bcl-2 gene [15, 16].

2. Materials and Methods

2.1. Plasmid Construction. The MLuc gene from the pMetLuc-Control plasmid (Clotech Inc., Palo Alto, CA, USA) was amplified by PCR using the forward primer 5’-CGAAGGTATGACATCAAGGTTGGTTG-3’ and the reverse primer 5’-TATGTATCTAGAGTCGGCGG-3’ (HindIII and XbaI restriction sites are indicated in bold letters). The amplicon was cut by HindIII and XbaI. pG3-Control (Promega Inc., Madison, WI, USA) was cut using the same restriction enzymes to remove the firefly luciferase gene, and the MLuc amplicon was inserted. The recombinant vector was designated pMLuc. Two complementary oligonucleotides 5’TCTAGAAATATCCAA TTCTGGTGCTGATCCGTGGACCATATCCAAATTCTCTGT GCTGATTCAAGA3’ and 5’AGATCTTATAGGTTAGG ACAGCAGATAAGGAACGTTTTTAGGGTGAGGACGA CGAGATAGATC3’ containing two miR15/miR-16 target sequences (the 3’ UTR target site in human Bcl-2 mRNA), indicated in bold letters, were annealed. The double-stranded oligonucleotide was inserted into the XbaI site of pMLuc which is located downstream of the stop codon of the MLuc gene. The recombinant vector obtained was named pMLuc-UTR. pSEAP2-Control (Clotech Inc.) containing the SEAP gene was used as an internal control.

2.2. Pre-miRNA. Pre-miR-16 and the negative control pre-miR (NC miR) were purchased from Ambion (Austin, TX, USA).

2.3. Cell Culture and Transfection. HEK293T and HepG2 cells (purchased from the Bioresource Collection and Research Center, Taiwan) were cultured in DMEM containing 10% fetal bovine serum (both from Gibco BRL, Gaithersburg, MD, USA) at 37°C in a humidified CO2 incubator. For transfection, the cells were grown in 6-well plates to 70% confluence. Transfection with plasmids and pre-miRNA was performed using a previously described protocol [17].

2.4. Assays of MLuc and SEAP Activity. Culture medium was collected and centrifuged, then 20 μL aliquots of the cell-free medium diluted in 180 μL of 0.9% NaCl were assayed for MLuc and SEAP activity using kits purchased from Clotech. For the MLuc assay, the 10× MLuc substrate stock was diluted to 1× in reaction buffer then diluted to 0.5× in double-distilled H2O; then 5 μL of the substrate/buffer solution and 50 μL of the diluted sample were mixed and loaded in 96-well black plates to detect luminescence. For the SEAP assay, 30 μL of 1× dilution buffer and 10 μL of the diluted sample were mixed and incubated at 65°C for 30 minutes and 4°C for 2 minutes in 96-well black plates, then 40 μL of SEAP substrate solution was added and the samples incubated at room temperature for 30 minutes before detecting the luminescence.

2.5. Quantification of miR-16 by qRT-PCR. HepG2 cells were grown in 6 cm Petri dishes to 70% confluence then were treated for different periods with 100 μM epigallocatechin gallate (EGCG). Total RNA was extracted from the cells using Trizol reagent (Invitrogen). qRT-PCR analysis for miRNAs was performed using TaqMan MicroRNA assays kit (Applied Biosystem, Foster City, CA, USA). Fifty nanograms of total RNA were subjected to reverse transcription (RT) using miR-16 and U6 RT primers (has-miR-16 UAGCCAGCACGU AAAUAUUGCC; U6 CGCAAGUGAUGCACAC GCAAAU-UUGUGAACAUAAUUUU) purchased from Applied Biosystem. RT was performed on a PCR machine (16°C for 30 minutes, 42°C for 30 minutes, and 85°C for 5 minutes). Real-time PCR was performed using an ABI PRISM 7900 (Applied Biosystem) and specific primer sets for miR-16 and U6 RNAs from Applied Biosystem. The conditions for the real-time PCR were 95°C for 10 minutes and 40 cycles of 95°C for 30 seconds and 60°C for 1 minute. The ΔCt was calculated by subtracting the Ct for U6 small nuclear RNA from that for miR-16. The ΔΔCt was calculated by subtracting the ΔCt for the untreated control from that for the EGCG-treated sample. The fold change in miR-16 levels was calculated as log 2−ΔΔCt.

3. Results and Discussion

To monitor the activity of miRNA in vivo without destroying the test cells, we devised a secretable reporter system consisting of three vectors that allows accurate and continuous measurement of the test miRNA. All 3 vectors (pMLuc, pMLuc-UTR, and pSEAP2-Control) were designed to use the same SV40 promoter to regulate the expression of the reporters or the internal control. pSEAP2-Control serves the dual role of (1) normalizing the transfection efficiency and (2) normalizing the activity of the SV40 promoter, which may differ in the context of various treatments. We first examined the signal-to-noise (background) ratio of the system and the stability of the two enzymes in culture medium to freeze thawing to determine whether samples from a time-course study could be frozen then assayed simultaneously. To examine these parameters, HEK293T cells were transfected with pMLuc or pSEAP2-Control, then 1 mL of culture medium was withdrawn 24 hours after transfection and centrifuged. The cell-free medium was then incubated at 37°C in a CO2 incubator and 20 μL aliquots were withdrawn at different time points and assayed for MLuc or SEAP activity. The results, shown in Figure I(a), indicate that the background signals from DMEM and conditioned medium from nontransfected cells were <0.1% of that from conditioned medium from pMLuc- or pSEAP2-Control-transfected cells. The secreted enzymes were stable.
in culture medium at conditions of 37°C and 5% CO2 for several hours (Figure 1(b)). A single freeze-thaw cycle reduced MLuc activity by 20%, but activity was not further reduced by freezing for 7 days (Figure 1(c)). Freeze-thaw and freezing for 7 days had no effect on SEAP activity (Figure 1(c)). This shows that we can collect and freeze all the samples from a single experiment and test them later without compromising the accuracy of the results. Thus, MLuc and SEAP are practicable for a multisampling reporter assay.

To test whether our system faithfully reflected the cellular activity of miR-15/miR-16, HEK293T cells were cotransfected with pSEAP2-Control, pMLuc or pMLuc-UTR, and pre-miR-16 or NC miR. The culture medium was replaced with fresh medium at 16, 24, and 36 hours after transfection and aliquots of medium were taken 2 hours after each medium change and assayed for MLuc and SEAP. The MLuc/SEAP activity ratios after various treatments are shown in Figure 2(a). The results showed that the ratios for the pMLuc-UTR + pre-miR-16 group (black bars) were 2- to 3-fold lower than those for the pMLuc-UTR + NC miR group (gray bars), indicating that pre-miR-16 suppressed the expression of MLuc. In addition, the ratios for the pMLuc-UTR + NC miR group (gray bars) were 1.5-fold lower than those for the pMLuc + NC miR group (white bars). This is because endogenous miR-15/miR-16 can target the MLuc-UTR mRNA generated from pMLuc-UTR. Cotransfection with pMLuc, which does not contain miR-15/miR-16 target sites, resulted in a higher ratio. These results indicate that our reporting system can be used to detect miR-15/miR-16 activity without disrupting the cells. The fourth group pMLuc + pre-miR-16 (hatched bars) showed higher ratios than those of the pMLuc + NC miR group (white bars), but a similar trend. This may be due to some global effect of miR-16 on the treated cells. In addition, by monitoring

![Figure 1: Characterization of the multisampling reporter system. HEK293T cells were grown in 6-well plates to 70% confluency, then some were left untreated and others were transfected with 0.5 μg of pMLuc or pSEAP2-Control for 24 hours. (a) An aliquot of the medium was assayed for MLuc (left) or SEAP (right). (b) The culture medium was collected and incubated at 37°C in a 5% CO2 humidified chamber and aliquots were withdrawn at different time points and assayed for MLuc (left) or SEAP (right). (c) Medium samples were frozen at −20°C in separate Eppendorf tubes, and thawed at the indicated times, and assayed. The data presented are the mean ± SD of triplicates.](image-url)
Figure 2: Reporting ability of the multisampling reporter system. HEK293T cells were grown in 6-well plates to 70% confluency. (a) The cells were cotransfected with 0.5 μg of pSEAP2-Control, 0.5 μg of pMLuc or pMLuc-UTR, and 30 nM pre-miR-16 or NC miR. The medium was replaced with fresh medium at different time points after transfection, and an aliquot of medium was withdrawn 2 hours later and assayed for MLuc and SEAP. (b) The cells were cotransfected with 0.5 μg of pMLuc-UTR and pSEAP2-Control, together with different concentration of pre-miR-16 (0–35 nM) and NC miR (35–0 nM), then, 24 hours later, an aliquot of medium was withdrawn from each culture medium and assayed. The data presented are the mean ± SD of triplicates.

Figure 3: Increase in miR-16 activity after EGCG treatment of HepG2 cells. HepG2 cells were grown in 5 mL of DMEM in 6cm Petri dishes to 70% confluency. (a) The cells were cotransfected for 6 hours with 4 μg of pMLuc or pMLuc-UTR and 4 μg of pSEAP2-Control, then 5 μL of 100 mM of EGCG (dissolved in DMSO) or DMSO was added. The medium was replaced with fresh medium at 24 and 36 hours and an aliquot of medium was withdrawn 2 hours later and assayed for MLuc and SEAP. The fold repression is defined as the MLuc (from pMLuc)/SEAP ratio divided by the MLuc (from pMLuc-UTR)/SEAP ratio. (b) The cells were treated with or without EGCG for 24 or 36 hours, then total RNA was extracted and subjected to qRT-PCR to detect levels of miR-16 and U6 small nuclear RNA (see Section 2 for details). The data presented are the mean ± SD of triplicates; ∗P < .05.

the same population of cells at 16, 24, and 36 hours, we found that the difference between the pMLuc-UTR + NC miR group ratios (gray bars) and the pMLuc-UTR + pre-miR-16 group ratios (black bars) decreased from 2.9-fold to 2.4-fold, then to 2.2-fold. This suggests that there is a gradual reduction in miR-16 levels in cells with time. To test the sensitivity of our reporting system, HEK293T cells were cotransfected with pMLuc-UTR + pSEAP2-Control, and different concentrations of pre-miR-16 and NC miR (total concentration of the two 35 nM) and aliquots of medium were taken 24 hours after transfection and assayed for MLuc and SEAP activity. As shown in Figure 2(b), dose-dependent suppression of MLuc was seen in response to an increasing amount of pre-miR-16.

HepG2, a liver cancer cell line, is known to exhibit upregulation of miR-16 levels (1.5-fold) following treatment with epigallocatechin gallate (EGCG) [18]. We, therefore, used our reporter system to monitor changes in this endogenous miRNA. HepG2 cells were cotransfected with pSEAP2-Control and either pMLuc-UTR or pMLuc then were cultured in the presence or absence of EGCG. The result is shown in Figure 3, in which “fold repression” is defined as
the MLuc (from pMLuc)/SEAP ratio divided by the MLuc (from pMLuc-UTR)/SEAP ratio. A significant increase in fold repression was seen in 24 hours of EGCG treatment compared to the blank control. A concomitant increase in miR-16 levels was confirmed by quantitative real-time RT-PCR (Figure 3(b)). The role of miRNA in fine tuning of protein synthesis can be elicited from the results of Figures 2(b) and 3. We have used our system to screen compounds which might up- or down-regulate a given miRNA. However, we found that some compounds had a direct effect on the enzymatic activities of MLuc and/or SEAP, leading to changes in the MLuc/SEAP ratio, while miRNA activities remained unchanged. We overcome this potential fault by calculating the fold repression instead of the MLuc/SEAP ratio, as the MLuc and SEAP activities were normalized in the calculation of the fold repression. We, therefore, strongly favor using the fold repression data for drug screening.

4. Conclusions

Our pMLuc/pSEAP2-Control-based reporter facilitates multisampling for monitoring the activity of a given miRNA using culture medium. The extracellular reporters are less affected by intracellular modifications. Time-course studies can be performed using the same population of cells. Samples collected at constant interval after each medium refreshment provide an activity readout of a given miRNA in real time. Moreover, the same population of cells used in detecting miRNA activity can be used for RNA extraction for further determination of miRNA levels. This method offers the benefits of accuracy, convenience, and cost effectiveness. With no need for cell lysis or addition of protease inhibitors, our system is more applicable to robust high-throughput drug screening. Finally, the system is versatile, as it can be easily switched to monitor the activity of any given miRNA by inserting different synthesized miRNA-targeting sequences into the XbaI site of pMLuc.

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