Global analysis of microRNA target gene expression reveals that miRNA targets are lower expressed in mature mouse and Drosophila tissues than in the embryos

Zhenbao Yu*, Zhaofeng Jian¹, Shi-Hsiang Shen, Enrico Purisima¹ and Edwin Wang¹,*

Mammalian Cell Genetics Group and ¹Computational Chemistry & Biology Group, Biotechnology Research Institute, National Research Council Canada, Montreal, Quebec, H4P 2R2, Canada

ABSTRACT

MicroRNAs (miRNAs) are non-coding small RNAs of ~22 nt that regulate the gene expression by base pairing with target mRNAs, leading to mRNA cleavage or translational repression. It is currently estimated that miRNAs account for ~1% of predicted genes in higher eukaryotic genomes and that up to 30% of genes might be regulated by miRNAs. However, only very few miRNAs have been functionally characterized and the general functions of miRNAs are not globally studied. In this study, we systematically analyzed the expression patterns of miRNA targets using several public microarray profiles. We found that the expression levels of miRNA targets are lower in all mouse and Drosophila tissues than in the embryos. We also found miRNAs more preferentially target ubiquitously expressed genes than tissue-specifically expressed genes. These results support the current suggestion that miRNAs are likely to be largely involved in embryo development and maintaining of tissue identity.

INTRODUCTION

MicroRNAs (miRNAs), encoded in the chromosomal DNA and transcribed as longer stem–loop precursors, termed pri-miRNAs, are non-coding small (21–23 nt) RNAs that regulate the expression of target mRNAs [reviewed in (1–4)]. Upon transcription, pri-miRNA is converted to mature miRNA duplex through sequential processing by RNase III family of endonucleases Drosha and Dicer (3,4). One strand of the processed duplex is incorporated into a silencing complex and guided to target sequences by base pairing [reviewed in (5,6)]. This results in the cleavage of target mRNAs or repression of their productive translation (5,6). In the past few years, several hundred miRNAs were identified in animals and plants. It is currently estimated that miRNAs account for ~1% of predicted genes in higher eukaryotic genomes (7).

Despite the large number of identified miRNAs, only a handful of them have been functionally characterized. For example, lin-4 and let-7 regulate the timing of larval development in Caenorhabditis elegans (8,9). Lsy-6 and miR-273 act sequentially to control the left/right asymmetric gene expression in C.elegans chemosensory neurons (10). Bantam promotes cell proliferation and inhibits apoptosis in Drosophila (11). MiR-14 suppresses cell death and regulates fat metabolism (12). MiR-181 potentiates B-cell differentiation (13). These findings, together with the complicated expression patterns and large number of predicted targets, imply that miRNAs may regulate a broad range of physiological and developmental processes.

Identification of the targets of each miRNA is crucial for understanding the biological function of miRNAs. Accumulating empirical evidence has revealed the importance of the 5-terminal segment of miRNAs with 6–8 nt in length, called ‘seed’ region, for miRNA function (14–17). For example, systematical single nucleotide mutation studies demonstrated that base pairing of miRNAs to their targets with 7 nt at the 5-terminus of miRNAs from position 2 to position 8 is essential and sometimes sufficient for miRNAs to knockdown their target expression (14). Based on these discoveries, several computational methods have been developed to search for miRNA targets (18–27). Most of these methods have been biologically validated and proved to be very efficient and accurate. The accuracy of these methods has also been proved by large scale gene expression profile studies (28,29). In one study, Lim et al. (28) reported that transfections of miR-1 and miR-124 into HeLa cells, respectively caused down-regulation of large numbers of target mRNAs and majority (76 and 88%, respectively) of down-regulated mRNAs showed a segment with 6 nt complementary to the 5′-terminus of the transfected miRNAs.
(the ‘seed’ sequence). In another study, Krutzfeldt et al. (29) demonstrated that knockdown of miRNA-122 by intravenous administration of miRNA ‘antagomirs’ led to up-regulation and down-regulation of a large number of genes in liver. They found that the 3′-untranslated regions (3'-UTRs) of up-regulated genes are strongly enriched in miRNA-122 ‘seed’-match motifs, whereas down-regulated genes are depleted in these motifs (29).

These methods have yielded a large number of candidate targets in both plants and animals. The estimated human miRNA targets can account for up to one-third of human genes (23). The diversity and abundance of miRNA targets reflect that miRNAs and their targets appear to form a complex regulatory network. For example, a single miRNA can regulate hundreds of mRNAs and a single mRNA can be targeted by several different miRNAs.

Systematical analysis of gene expression profiles has been proved to be valuable for studies on diverse biological processes (30–33). Based on its biochemical function, the biological functions of a miRNA should depend on the combination of its action to each of all its targets for their expression. Accordingly, Farh et al. (34), Stark et al. (35) and Sood et al. (36) have recently found a clear correlation of the expression of miRNAs and their targets through analysis of mRNA target expression profiles and in situ hybridization. In this study, we undertook a global analysis of the expression of miRNA targets in human, mouse and Drosophila using several public gene expression datasets (37–39). To our surprise, we found that the average expression levels of the total targets of all miRNAs are significantly different in distinct tissues compared to the expression of the total genes. For example, we found that the expression levels of miRNA targets are significantly lower in all mouse mature tissues and Drosophila later life stages than in the embryos. We also found that miRNA targets are more ubiquitously expressed.

MATERIALS AND METHODS

Stand-alone Java programs or Perl scripts were used where necessary to facilitate the following analyses.

Datasets used in this study

The datasets used in this study include two complete lists of human miRNA targets published by Lewis et al. (23) and Krek et al. (21), one complete list of human and mouse miRNA targets published by John et al. (19), two complete lists of Drosophila miRNA targets published by Enright et al. (18) and Stark et al. (26), a microarray expression dataset for more than 10 000 human genes across 52 tissues and cell lines (38), a microarray expression dataset for nearly 40 000 known and predicted mRNAs in 55 mouse tissues (39) and a microarray expression data for nearly one-third of all Drosophila genes during the whole life-cycle (37).

All miRNA target datasets were downloaded from the most recently updated websites. The datasets published by Lewis et al. (23) and Krek et al. (21) contain human miRNA targets but not mouse miRNA target. To get mouse miRNA targets, we mapped the human miRNA targets to the orthologous genes in mouse genome. To do so, we first converted the human gene IDs from the datasets published by Lewis et al. (23) and Krek et al. (21) to NCBI GeneIDs using ID Converter (http://idconverter.bioinf.cipf.es/). To obtain the corresponding mouse orthologous gene pairs, we queried the homoloGene data file downloaded from HomoloGene (ftp://ftp.ncbi.nih.gov/pub/HomoloGene/) using the human genes.

For the human gene microarray dataset (38), we removed some samples which derived from the same or similar tissues in order to avoid over-representation of these samples for the analysis. For example, several brain samples were present in the dataset, we removed most of them. Totally 41 human tissues were analyzed.

As the mouse microarray gene chips were designed to detect 40 000 known and predicted mouse mRNAs (39), it is expected that many predicted miRNAs should not be expressed. Totally 21 622 genes could be detected in at least one tissue in the microarray gene chips. The rest (almost half) of them were not detected in any tissues. Therefore, only 21 622 genes were analyzed in this study.

Ranking of miRNA target genes

To study the correlation of the expression of miRNAs and their targets, we analyzed the microarray expression data containing ~10 000 genes published by Johnson et al. (38) across 41 human tissues by ranking each gene over all tissues according to its expression level in the respective tissue as described previously (28). A lower rank number represents less expression level. For example, if a gene is expressed less in a defined tissue than in any other tissues, the rank number of this gene in this tissue is 1. Similarly, rank number 41 means that the expression level of a gene in this tissue is lower than that in any other tissues. We then mapped miRNA targets to the genes in the microarray dataset. The miRNA targets we used in this analysis are from the datasets published by Lewis et al. (22) and John et al. (19), respectively. We only chose the miRNAs, which have more than 50 targets found in the microarray dataset for further analysis. On average, 180 targets per miRNA could be found in the microarray dataset. For each miRNA, in each tissue, we obtained a rank number of each of its targets.

To facilitate a more global view, we grouped the genes into two sets, one with lower half of rank numbers and the other with higher half. When there is an odd number of a rank, the middlemost rank is excluded. In our specific example with 41 tissue samples, one set consists of ranks 1–20 and the other from 22–41, excluding rank 21. For any miRNA in any tissue, we counted the total number of its targets within lower-rank set and divided it by the total number of its targets within higher-rank set to yield the calculated rank ratio, RR. For example, for a miRNA in the 41-tissue set, RR = N_Rank 1–20/N_Rank 22–41. The RR value is an indicator of the preferential tissue expression of a given miRNA’s target genes. An RR value greater than one means that the majority of expressed targets of a miRNA in this tissue have a lower expression level than the median level of expression of the miRNA’s targets across all the tissues. If the RR value of a miRNA is greater in a particular tissue than that in any others, the expression level of the targets of this miRNA in this tissue is very likely to be the lowest among the 41 human tissues. We also did the same analysis for total genes presented in
the microarray dataset. The RR value of target genes for each miRNA in a tissue was normalized by the RR value of total genes in the same tissue and then plotted as a function of tissues and miRNAs, respectively. The RR value provides a global descriptor of the tissue distribution of a miRNA’s target genes rather than the expression levels of individual genes. It does not provide gene-specific information but allows the extraction of global trends of a group of genes (miRNA target genes) amid the noisy data for individual genes. This method was also used to analyze a microarray dataset containing 55 mouse samples. In this case, \[ RR = \frac{N_{\text{Rank 1–20}}}{N_{\text{Rank 22–41}}} \]

Comparison of the expression level of miRNA targets in the embryos of mouse and Drosophila with that in their mature tissues

To explore the difference of the expression level of the total targets of all available miRNAs in mouse mature tissues from that in 12.5-day mouse embryo, we compared the absolute expression value of each gene between tissues and the embryo to determine if a gene has a higher or lower expression level in a tissue than in embryo. We then calculated the ratio of lower-expressed targets to the higher-expressed targets in each tissue (termed \( R_{\text{mirna}} \)).

To determine the statistical significance of the observation, we performed resampling statistical tests. A more detailed explanation of randomization tests was described previously (40). In each test, we randomly picked up the same number of genes as the number of miRNA targets from total genes. We calculated the ratio of lower-expressed genes to higher-expressed genes in this sub-pool and defined it as \( R_{\text{random}} \). We performed the randomly picking test for 5000 tests and counted the number of times (N) when \( R_{\text{random}} > R_{\text{mirna}} \). We defined \( P \)-value as \( N/5000 \). If \( P < 0.05 \), we thought that the number of lower-expressed miRNA targets is significantly more than that of higher-expressed miRNA targets. We rejected the hypothesis if the \( P < 0.05 \).

We used three sets of miRNA targets, respectively, for this analysis. They are the mouse miRNA dataset published by John et al. (19) and the human miRNA datasets published by Lewis et al. (23) and Krek et al. (21), respectively. Human miRNA targets were converted to mouse orthologous for the analysis. We performed correlation analysis of the results obtained from each of the three datasets and calculated the correlation coefficient using \( R \) (http://www.r-project.org/).

A similar method was used to compare the expression of miRNA targets in the different periods of Drosophila life cycle with that in the 23–24 h embryo using a microarray dataset published by Arbeitman et al. (37), except that the ratio of number of miRNA targets whose expression level in a defined period is 2-fold lower than that in 23–24 h embryo to that of miRNA targets whose expression level is two times higher than that in 23–24 h embryo was represented. The statistical analysis for this set was conducted as described above.

Analysis of the tissue-specificity of miRNA target expression

Using the microarray database published by Zhang et al. (39), in which 21,622 mouse genes including 2276 miRNA targets published by John et al. (19) were expressed in at least one tissue, we analyzed the tissue-specificity of miRNA target expression. The mouse genes are classified into 55 groups according to the number of tissues (1–55) in which a gene was expressed. The total numbers of miRNA targets and total genes in each group were counted, respectively. The percentage of miRNA targets to the total genes in each group was calculated and compared to the percentage (10.52%) of the total miRNA targets (2276) to the total genes (21,622) for determining if the miRNA targets are enriched in or excluded from the respective group.

RESULTS

Expression level of miRNA targets is tissue-dependent

Since the function of a miRNA depends on the combination of its actions to all of its targets for their expression, to understand the global role of these numerous miRNAs, we undertook a global analysis of the expression of miRNA targets in human, mouse and Drosophila using several public datasets. We first analyzed the microarray expression data containing \( \sim 10,000 \) genes over 41 human tissues published by Johnson et al. (38). We compared the relative expression level of the total targets of individual miRNAs across the 41 human tissues. Since each miRNA has many targets and the absolute expression levels of these targets are very different, the contribution of each target to the result should be different if we simply use the average expression level of these targets for comparison. To make each target equally contribute to the comparison, we ranked each gene over 41 human tissues according its expression levels in the respective tissues (see Materials and Methods). A lower rank number means a lower expression level. For each miRNA, in each tissue, we counted the total number of its targets (23) at each rank position (Supplementary Table S1). By comparing the distribution of the rank number of the targets between different tissues, we could find the relative expression levels of the total targets of a miRNA in each tissue compared to other tissues. This method could avoid the effect of the bias of the absolute expression levels of the miRNA targets on the analysis. Figure 1A shows a typical result for the distribution of the rank number of miR-128a targets (23) in liver and brain. In liver, the number of miR-128a targets with a lower rank number is obviously more than that of those with a higher rank number. In contrast, in brain, the result is reversed. This suggests that the overall expression level of the targets of miR-128a in liver is lower than that in brain. To obtain a quick overview, we grouped the targets into two sets, one with rank numbers from 1–20 and the other with rank numbers from 22–41 (see inset in Figure 1A). We then calculated the RR value (see Materials and Methods), \( \frac{N_{\text{Rank 1–20}}}{N_{\text{Rank 22–41}}} \). A higher RR value of a miRNA in a defined tissue means that more targets of the miRNA have a lower expression level (lower ranking number) in the tissue and less targets of the miRNA have a higher expression level (higher ranking number) than that in most tissues (more than half of the 41 tissues). A RR value more than one means that the total number of the targets with a rank number lower than 21 is more than the total number of the targets with a rank number higher than 21, and suggests...
Figure 1. Ranking of the expression levels of miRNA targets in human tissues. Each miRNA target was ranked over 41 human tissue samples as described in the ‘Materials and Methods’ according to its expression level. The RR means the ratio of the total number (N_{Rank 1-20}) of the miRNA targets with a lower rank number (between 1 and 20) to the total number (N_{Rank 22-41}) of the miRNA targets with a higher rank number (between 22 and 41). The RR value of the target genes of each miRNA in a tissue is normalized by the RR value of the total genes in the same tissue. 

(A) Ranking results of the 295 targets of miR-128a in liver and brain. 

(B) The RR values of the 55 miRNAs are plotted as a function of human tissues and cell lines. 

(C) The RR values over the 41 tissues and cell lines are plotted as a function of miRNAs. 

(D) For each miRNA, the RR values of its targets were ranked across the 41 samples from higher value to lower value. The miRNAs shown are highly specifically expressed in the indicated tissues as demonstrated previously for miRNA-1 and miRNA-133 in references (51,54); miRNA-122 in reference (60); miRNA-16, miRNA-142, miRNA-181 and miRNA-223 in reference (61); miRNA-155 in references (52,53). The rank positions in the indicated tissues were obtained and shown in the figure.
Figure 2. Ranking of the expression levels of miRNA targets in mouse tissues. Each miRNA target gene was ranked over 55 mouse samples according to its relative expression level in the respective sample as described in the ‘Materials and Methods’. The RR means the ratio of the total number (N_{Rank 1–27}) of the miRNA targets with a lower rank number (between 1 and 27) to the total number (N_{Rank 29–55}) of the miRNA targets with a higher rank number (between 29 and 55). The RR value of the target genes of each miRNA in a tissue is normalized by the RR value of the total genes in the same tissue. (A) Comparison of the target gene expressions (RR values) of the 141 miRNAs in bone marrow and 12.5-day embryo. (B) RR values over 141 miRNAs are plotted as a function of mouse tissues. (C) The percentage of miRNAs, of which the total number (N_{Rank 1–27}) of the targets with lower rank number (between 1 and 27) is more than the number (N_{Rank 29–55}) of the targets with higher rank number (between 29 and 55) (RR value >1).
that the general expression level of the targets of a miRNA in the tissue is most likely to be lower than the median expression level of the targets in all tissues. For example, the RR value for miR-128a is 2.1 (197 targets/92 targets) in liver and 0.57 (104 targets/184 targets) in brain, suggesting a lower expression level of the miR-128a targets in liver than that in brain. Totally, we analyzed 55 miRNAs, each of which have at least 50 targets presented in the microarray dataset (average 180 targets/miRNA), across 41 tissues. We also did the same analysis for total genes present in the microarray dataset. The RR values are shown in Supplementary Table S1. The RR value of the target genes for each miRNA in a tissue was normalized by the RR value of the total genes in the same tissue and then plotted as a function of miRNAs and tissues, respectively (Figure 1B and C). As expected, for each individual miRNA, the RR values in different tissues are equally distributed around one (the number of the tissues with a RR value more than one is similar to the number of the tissues with a RR value less than one) (Figure 1B).

We next studied the correlation of the expression of miRNAs and their targets using several public microRNA microarray expression datasets. Because of the low accuracy of the miRNA microarray expression profiling data, we did not find obvious correlation of the expression of miRNAs and their target expression (data not shown). For example, the miRNAs microarray expression profiles published by different researchers are not consistent (41–50). Therefore, we next focused on the miRNAs, which were highly specifically expressed in one or a few tissues and whose expression was highly confirmed by different researchers through northern blot analysis and/or cloning method (51–54).

To do so, for each miRNA, we further ranked the RR values of their targets across the 41 tissues (from high value to low value) and found the rank position for the tissues in which the miRNA is highly specifically expressed. As shown in Figure 1D, most miRNAs have a low rank number of their target RR values in the tissues where they are specifically expressed. This suggests that the expression of miRNAs has a negative correlation with their target expression. However, we noted that two miRNAs (miRNA-124a and miRNA-125b) specifically expressed in brain do not show negative correlation with their targets. For example, the RR values of miRNA-124a and miRNA-125b in brain ranked at 38 and
35 of the 41 tissues, respectively. Interestingly, Sood et al. found that for many miRNAs, their predicted targets are up-regulated in neuronal tissues compared with most other tissue types, without considering the expression patterns of the miRNAs (36). We obtained a similar result (see below). However, it is not clear if the unexpected higher expression level of the targets of miRNA-124a and miRNA-125b in brain, where the miRNAs are highly expressed, is caused by the general high expression level of miRNA targets in this tissue.

When we looked at the distribution of the RR values in each tissue (Figure 1C), to our surprise, we found a dramatic difference between different tissues. In some tissues, the preponderance of miRNAs have a RR > 1. Conversely, in some tissues, RR < 1 for an overwhelming fraction of the miRNAs. This suggests that the overall expression level of the total targets of all miRNAs is quite lower in some tissues and higher in others. For example, in bone marrow, 54 of the 55 miRNAs have a RR value more than one, whereas in brain, none of them has a RR value more than one, indicating that the overall expression level of miRNA targets is lower in bone marrow but higher in brain. Similar results were obtained when using the miRNA targets published by John et al. (19) (Supplementary Figure S1).

Expression levels of miRNA targets are lower in mature tissues than in embryos in both mammalian and fly

We next analyzed the expression of miRNA targets in 55 mouse samples using the gene expression profile data published by Zhang et al. (39) and the dataset of microRNA targets published by John et al. (19). Similar to what we found in human tissues, the RR value for all of the 141 miRNAs in mouse bone marrow is greater than one (Figure 2A), suggesting that the expression level of miRNA targets in this tissue is obviously lower than the median level across the other tissues. A similar result was observed in other hematopoietic cells-rich or lymphocytes-rich tissues, such as thymus, spleen and lymph node (Figure 2B and C). Most importantly, the expression levels of miRNA targets in embryo, embryo head and placenta are significantly higher than that in most mature tissues (Figure 2B and C). For example, in 12.5-day embryo, all of the 141 miRNAs have a RR value below one.

To further confirm the observation, we directly compared the average expression value of the total 2276 miRNA targets in the mouse tissues. Consisting with the result above derived from ranking analysis, the average expression level of miRNA target genes is higher in 12.5-day embryo than that in any mature tissues except for cerebral cortex (Figure 3). For example, the levels in bone marrow, spleen, thymus...
and lymph node are significantly (more than two times) lower than in 12.5-day embryo, whereas, the levels in neuronal tissues, such as whole brain, middle brain, hindbrain, cerebellum and trigeminus are slightly lower than in 12.5-day embryo or even higher (cerebral cortex) than in 12.5-day embryo. This consists with Sood et al.‘s observation (36) and our observation in human brain. This analysis further confirmed that, miRNA target expression levels are

Figure 5. Analysis of the expression of miRNA targets in Drosophila. (A and B) A published microarray dataset (37) was analyzed for the comparison of the expression levels of the miRNA targets during different periods of Drosophila life cycle. The total number of miRNA targets whose expression level is two times lower than that in 23–24 h embryo ($N_{<1.5\times E23–24h}$) was divided by the total number of those with expression level two times higher than that in 23–24 h embryo ($N_{>2\times E23–24h}$). Two sets of Drosophila miRNA targets published by Enright et al. (18) and Stark et al. (18,18,26), respectively were analyzed. As control, the expression levels of the total genes were also analyzed by the same way. E, embryo; L, larva; M, pupae; Am, adult male and Af, adult female. (C) Correlation between the data ($N_{<1.5\times E23–24h} / N_{>2\times E23–24h}$) obtained using two different miRNA target datasets.
significantly lower in most mature mouse tissues, except for the neuronal tissues, than in embryos.

We then focused on the comparison of mouse mature tissues to embryo. To do so, we counted the total number of miRNA targets whose expression level is lower in a given tissue than that in 12.5-day embryo (N_{E12.5}) and divided it by the total number of miRNA targets whose expression level is higher than that in 12.5-day embryo (N_{>E12.5}). As shown in Figure 4A, in all mature tissues, the lower expressed target number is more than higher expressed target number (N_{<E12.5}/N_{>E12.5} > 1). As a control, we carried out the same calculations for all genes. We see that, for all tissues, the N_{<E12.5}/N_{>E12.5} value of total genes is lower than that of miRNA targets (Figure 4A). Resambling statistical tests (see ‘Materials and Methods’ for details) demonstrated that the difference is significant (P < 0.0002 for almost all of the tissues, Supplementary Table S2). To further confirm the observation, we performed the same analysis with the total miRNA targets published by Lewis et al. (23) and Krek et al. (21), respectively. We found the similar patterns (Figure 4B and C). Figure 4D shows that the data obtained using the miRNA targets published by the three groups respectively, are highly correlated. This supports both the quality of the original data and our analysis method.

To determine if the observed expression pattern is conserved in other species, we analyzed the published gene expression profile over 75 stages of the whole life cycle of Drosophila (37). As shown in Figure 5A and B, compared to 23–24 h embryo, while the ratio of lower expressed miRNA targets to higher expressed miRNA targets remains the same during embryo period, it dramatically increases starting from larval periods and lasting to adulthood (P < 0.0002 for all larva and male adult, see Supplementary Table S3 for more details). There is a clear correspondence (R^2 = 0.93) between the data calculated using the target sets published by Enright et al. (18) and Stark et al. (26), respectively (Figure 5C). This data supports the important role of miRNAs for determining the timing of tissue differentiation during larva period of Drosophila development and maintaining the tissue identity during the adulthood.

It should be noted that the human microarray dataset (38) we used in this study does not contain human embryo and a large scale gene expression profile containing human embryo is not available. Consequently, we could not perform the comparison of the expression levels of human miRNA targets between human mature tissues and human embryos.

**MiRNAs more frequently target ubiquitously expressed genes than tissue-specific genes**

We analyzed the tissue-specificity of miRNA target expression using the microarray data representing 21622 mouse genes.
In this study, we found that the expression level of miRNA targets in mature tissues is significantly lower than that in embryos in both mammalian and fly and that miRNAs more frequently target ubiquitously expressed genes than tissue-specific genes. These findings support the recent observations that some miRNAs play a most important role in driving tissue terminal differentiation and maintaining tissue identity. Previous studies suggest that 10 to 30% of human genes are potential miRNA targets (19,22). However, analysis of the specific gene ontology (GO) molecular function classification among the predicted targets could not reveal any specific biological functions of animal miRNAs since the animal miRNA targets populated many GO functional categories (19,22). Only ~13% of mammalian miRNA targets predicted by Lewis et al. were involved in development according to the GO biological process categories (22). Failing to predict the functions of miRNA targets through GO analysis may be simply caused by the evolving stage of the classification of GO function categories. Alternatively, on principle, the functions of miRNAs could not be predicted by the GO function categories of their targets because the expression and therefore the functions of their targets are proposed to be turned down but not induced by miRNA expression and GO analysis can only tell the function of a group of genes in a GO function category when they are expressed or up-regulated but not that when they are down-regulated. Our studies demonstrate that statistical analysis of the expression of miRNA targets may reflect the global functions of miRNAs. However, the biological significance of the observation is not clear right now. One proposal is that the statistically lower expression level of miRNA targets in matured tissues than in embryos remains to be elucidated. One potential reason is that the miRNA expression level is lower in embryos. This is true for zebrafish. Recently, Wienholds et al. have reported that most zebrafish miRNAs were not detected during early development (55). However, the results regarding to the global expression patterns of miRNAs during mouse or Drosophila embryo development are currently not available although the differential expression of miRNAs in different tissues were clearly demonstrated. Another possibility is that the activity of miRNA machinery is lower in embryos than in other tissues. For example, Yang et al. (56) reported that dicer, an important protein for miRNA biogenesis and miRNA function, starts expression in 7-day old mouse embryos and remains stable through 17-day embryos. However, maternal Dicer protein is probably present from fertilization (57–59). In addition, a possible delay from miRNA processing to target gene inactivation might also contribute to the high level expression of miRNA targets in embryos.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

ACKNOWLEDGEMENTS

Funding to pay the Open Access publication charges for this article was provided by National Research Council Canada.

Conflict of interest statement. None declared.

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