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Imprinted DLK1-DIO3 region of 14q32 defines a schizophrenia-associated miRNA signature in peripheral blood mononuclear cells

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MicroRNAs (miRNAs) regulate gene expression at the post-transcriptional level and are important for coordinating nervous system development and neuronal function in the mature brain. We have recently identified schizophrenia-associated alteration of cortical miRNA biogenesis and expression in post-mortem brain tissue with implications for the dysregulation of schizophrenia candidate genes. Although these changes were observed in the central nervous system, it is plausible that schizophrenia-associated miRNA expression signatures may also be detected in non-neural tissue. To explore this possibility, we investigated the miRNA expression profile of peripheral blood mononuclear cells (PBMCs) from 112 patients with schizophrenia and 76 non-psychiatric controls. miRNA expression analysis of total RNA conducted using commercial miRNA arrays revealed that 33 miRNAs were significantly downregulated after correction for multiple testing with a false discovery rate (FDR) of 0%, which increased to 83 when we considered miRNA with an FDR < 5%. Seven miRNAs altered in microarray analysis of schizophrenia were also confirmed to be downregulated by quantitative real-time reverse transcription-polymerase chain reaction. A large subgroup consisting of 17 downregulated miRNAs is transcribed from a single imprinted locus at the maternally expressed DLK1-DIO3 region on chromosome 14q32. This pattern of differentially expressed miRNA in PBMCs may be indicative of significant underlying genetic or epigenetic alteration associated with schizophrenia.

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Introduction

Schizophrenia is a severely debilitating and complex neuropsychiatric disorder afflicting nearly one in 100 people throughout life.¹ Although some gross neuroanatomical and histological features have been associated with schizophrenia, none are consistent enough to be considered diagnostic.^{2–7} The scarcity of tangible pathology means that clinical diagnosis is based on descriptive criteria and requires persistent or episodic presence of certain symptoms over a defined period with associated disability and exclusion of other psychiatric conditions. As early intervention in schizophrenia is thought to be important in improving outcome, any delay in diagnosis or misdiagnosis owing to the inherent limitations of clinical descriptors may have a negative prognostic impact.^{8,9}

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Consequently, accessible biological markers are needed to improve the timing and accuracy of diagnosis, and identify individuals at very high risk of schizophrenia. This information may also provide the opportunity to tailor treatment to the individual and enable more effective early intervention.^{10–13} Despite having a strong genetic component, early expectations of a simple genetic diagnostic for schizophrenia have been thwarted as the heritability appears to involve a complex, heterogeneous mixture of common variants, each with small effect size.¹⁴

More recently, studies have attempted to move closer to the neuropathology of schizophrenia in an attempt to identify common pathways in the disorder. High-throughput gene expression analysis has revealed altered gene expression in post-mortem brain including the dorsolateral prefrontal cortex, superior temporal gyrus and amygdala, which are all functionally relevant to the pathology of schizophrenia.^{15–22} Although no single gene has been found to be consistently altered across studies or brain regions, most report the enrichment of pathways relevant to the putative neurobiology of schizophrenia. In many

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instances, the extent and shape of gene alteration in post-mortem brain may imply the involvement of aberrant regulatory or epigenetic influences. In support of this hypothesis, we recently reported changes in cortical microRNA (miRNA) biogenesis and expression in schizophrenia.^{23,24} These molecules strongly influence gene expression, particularly in the brain where they are thought to be involved in both neurodevelopment and in the regulation of synaptic structure and function.^{25–29}

Although the pathophysiology of changes in cortical miRNA expression remains to be determined, it is plausible that alteration of miRNA expression extends beyond the brain and could provide useful peripheral biomarkers for schizophrenia that would be accessible for clinical use. miRNA molecules are highly stable³⁰ and, owing to their regulatory function (predicted to regulate half of the human genome³¹), are arguably more informative and prognostic than gene expression. Certainly in the field of cancer biology, miRNA may be a more accurate predictor of tumor phenotype than its genotype.^{32,33} miRNA profiling of tissues and body fluids has been instructive in a variety of cancer types including chronic lymphocytic leukemia and lung, prostate, bladder and breast cancer.³⁴⁻³⁷ Perhaps more relevant to brain disorders like schizophrenia, miRNA profiling of whole blood in multiple sclerosis has revealed expression signatures associated with the disease and its remission status.³⁸ Similarly, miRNA analysis in Alzheimer's disease has revealed significant patterns of altered expression in peripheral blood mononuclear cells (PBMCs)³⁹ as well as in the cerebrospinal fluid and brain.40 PBMCs in particular represent an attractive alternative tissue for profiling active disease in living patients at statistically robust numbers. This accessible tissue can reflect global disease-associated changes in an underlying genetic disorder and may have special significance where immunological risk factors are believed to be involved. In schizophrenia, for example, prenatal maternal infection is thought to be an important risk factor^{41–44} that may leave immunologically relevant expression signatures in PBMCs.

In this study, we examined miRNA expression in PBMCs derived from a cohort of schizophrenia patients and non-psychiatric controls, and identified a significant schizophrenia-associated reduction in 83 miRNAs. A large proportion of this expression signature was derived from a cluster of 17 miRNAs residing in a single imprinted domain on 14q32. This discovery has implications for our understanding of epigenetic regulation and inheritance in schizophrenia, and may also have applications as a novel biomarker for the disorder.

Materials and methods

Participant recruitment and clinical assessment protocol

Participants, the majority of whom identified as Caucasian, were recruited by the Australian Schizophrenia Research Bank and the Hunter DNA Bank. In this study, the cohorts consisted of 112 participants with a lifetime diagnosis of schizophrenia and 76 non-psychiatric controls that were deemed eligible for inclusion after assessment by qualified psychologists using the Diagnostic Interview for Psychosis.⁴⁵ Participants were excluded if they were currently diagnosed with any organic brain disorders, substance dependence, brain injury that caused posttraumatic amnesia lasting > 24 h, mental retardation (IQ < 70), movement disorder or if they had electroconvulsive therapy within 6 months. These exclusion criteria were likewise used for non-psychiatric control subjects with the additional criterion that they had no personal or family history of either psychosis or bipolar disorder type 1.46 In addition, participants were required to speak English, and were aged between 18 and 65 years. Participants gave informed consent for inclusion in the study protocol, as approved by the institutional research ethics committees. All cases reported using antipsychotic medication at the time of interview and blood draw. Demographic and clinical variables of the cohort are summarized in Table 1 and described in detail in Supplementary Table 1.

Extraction of RNA and DNA from whole blood

Whole blood (20 ml) was collected from subjects and transported to the Australian Schizophrenia Research Bank for processing. PBMCs were extracted by centrifugation using Lymphoprep (Vital Diagnostics, Lincoln, RI, USA). Total RNA was extracted from PBMCs obtained using Trizol, as per the manufacturer's instructions (Invitrogen, Life Technologies, Carlsbad, CA, USA). RNA concentration was determined using a NanoDrop 2000 (Thermo Scientific, Delaware, ME, USA). Genomic DNA was extracted

 Table 1
 Summary statistics for clinical and demographic

 variables characterizing the schizophrenia and control cohort

Demographic/clinical variable	Summary statistics
Diagnosis: SZ/CTRL	112/76
Mean age	39.4
Gender: M/F	101/82
Mean age at the onset of illness	24.2
Family history of schizophrenia (cases): present/none	41/55
Family history of other psychosis (cases): present/none	49/47
Mean duration of illness	16.4
Mean RQI	9.3

Abbreviations: CTRL, non-psychiatric control; F, female; M, male; RQI, RNA quality indicator; SZ, schizophrenia. Family history of other psychosis, reports any other psychiatric mental illness in any first- or second-degree relative. from PBMCs using a standard salt extraction method and quantified by PicoGreen assay (Invitrogen, San Diego, CA, USA). The integrity of randomly selected genomic DNA was checked by agarose gel electrophoresis.

RNA integrity

RNA integrity was assessed using an Experion microcapillary electrophoresis device according to the manufacturer's instructions (Bio-Rad, Hercules, CA, USA). The average RNA Quality Indicator for this cohort was 9.3. Samples with RNA Quality Indicator values ≥ 6.3 were considered intact and therefore suitable for use in miRNA microarray experiments and quantitative real-time reverse transcription polymerase chain reaction (PCR).

miRNA expression profiling

Profiling of human miRNA expression (miRBase version 9.1) was achieved using the commercial miRNA microarray platform (Illumina, San Diego, CA, USA) as described previously.47 Microarray data was extracted and normalized with respect to the geometric mean of U44 and U49 small nucleolar RNA (snoRNA) expression. Analysis of variance was used for the evaluation of association between non-diagnosis variance (batch effects, gender and age) and miRNA expression. Age showed a significant correlation with miRNA expression and was therefore used as a covariate in the subsequent analysis. Differential expression analysis was performed using the significance analysis of microarray statistical analysis program (full academic version 2.23, website http://www-stat.stanford.edu/ ~tibs/SAM/).48 Normalized expression data for the differentially expressed miRNA is shown in Supplementary Table 2. Unsupervised hierarchical clustering was performed in Cluster (Stanford University, Palo Alto, CA, USA; Eisen et al⁴⁹) and a heat map visualized through Java Treeview V.1.1.1.50 Data were log transformed and median centered by genes. Genes and arrays were clustered, correlation un-centered, by average linkage clustering.

Quantitative real-time reverse transcription PCR

Validation of differentially expressed miRNA was performed by quantitative real-time reverse transcription PCR on a subset of the cohort consisting of 91 participants (57 schizophrenia, 34 non-psychiatric controls), similarly to that described previously.24 Briefly, 500 ng of sample RNA was treated with DNAse-I (Invitrogen, Life Technologies) and multiplex reverse transcription performed with Superscript II reverse transcriptase (Invitrogen, Life Technologies), a 3 nM mix of miRNA sequence-specific primers and primers for U44 snRNA and U49 snoRNA (for primer sequences, see Supplementary Table 3). Seven miRNA were chosen for real-time validation on the basis of strong differential expression on the microarray and/or biological significance. Triplicate reactions were set up in a 96-well format using the epMotion 5070 automated pipetting system

(Eppendorf, Hamburg, Germany) and carried out using the Applied Biosystems 7500 real-time PCR machine. Relative miRNA expression was determined with respect to the geometric mean of U44 and U49 snoRNAs by calculating the difference between the average cycle threshold (C_t) of the miRNA of interest and the geometric mean of the average C_t values of housekeeping snoRNA for each sample (ΔC_t). Data were not normally distributed (as determined by GraphPad Prism version 5.0); therefore, the significance of differential expression was determined using a Mann–Whitney U-test (PASW Statistics 18).

Target prediction and functional pathway analysis

Putative target genes for schizophrenia-associated miRNA identified in this study were predicted using the miRanda⁵¹ and TargetScan (release 5.1)⁵² search algorithms (http://www.microrna.org/microrna/home.do; http://www.targetscan.org/). These putative target genes were further filtered by selecting genes containing five or more predicted binding sites either by the same or multiple miRNA as these genes are most likely to be regulated by the differentially expressed miRNA.53 Pathway analysis was then performed on a combined list of targets from both algorithms (union), followed by a more stringent pathways analysis on those targets that were common to both miRanda and TargetScan (intersection) using WebGestalt Gene Set Analysis Toolkit V.2 (http://bioinfo.vanderbilt.edu/webgestalt/).54 Visualization of pathways and networks from this information was facilitated by the construction of an 'enrichment map' of significant pathways (P < 0.05 Benjamini-Hochberg) containing seven or more genes per category (http://baderlab.org/Software/EnrichmentMap).⁵⁵

Analysis of copy-number variation

Genomic copy-number status of the 14q32 region was investigated using commercial TaqMan copy-number assays (Applied Biosystems, Life Technologies, Carlsbad, CA, USA; Hs07397776 cn, Hs03879584 cn and Hs03889902 cn) corresponding to cytobands 14q32.2b, 14q32.2b and 14q32.31a, respectively (refer to Supplementary Figure 1 for details). Reactions were prepared as per the manufacturer's instructions using an EPMotion 5070 automated liquid handling device (Eppendorf) and carried out using an ABI7500 real-time PCR machine (Applied Biosystems). The CopyCaller software V.1.0 (Applied Biosystems) was used to calculate relative copy number at each of the three locations. Each sample was normalized to the copy-number reference assay (RNase P, part number 4316831), known to be present in two copies in a diploid genome.

Results

miRNA expression analysis

miRNA expression was analyzed in PBMCs of 112 patients with schizophrenia and 76 non-psychiatric controls using human miRNA array matrices (Illumina). After normalization and background

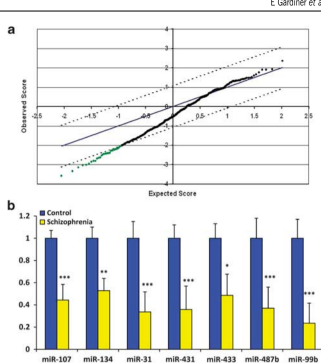


Figure 1 Schizophrenia associated microRNA (miRNA) expression in peripheral blood mononuclear cells (PBMCs). (a) Significance analysis of microarray (SAM) plot of differentially expressed miRNA in PBMCs from participants with schizophrenia (n = 112) compared with non-psychiatric controls (n = 76). The central solid blue line indicates equal expression and the upper and lower dashed black lines indicate levels for significantly altered expression (false discovery rate (FDR) of 0%). (b) Quantitative real-time reverse transcription polymerase chain reaction (Q-PCR) validation of miRNA significantly downregulated by miRNA array analysis. Bars indicate mean fold change ± s.e.m. of 91 samples (57 schizophrenia cases and 34 non-psychiatric controls). The control cohort is set at 1. *P < 0.05; **P < 0.01; ***P < 0.001 by Mann–Whitney U-test.

subtraction, signals corresponding to 358 miRNAs or approximately 40% of the array content remained. Statistical analysis revealed 33 significantly downregulated miRNA at a false discovery rate (FDR) of 0% (Figure 1a) increasing to 83 at an FDR <5% (Table 2). This included a number of brain-enriched miRNAs, miR-134, miR-128 and miR-181b. By contrast, there were no miRNAs that were significantly increased according to these criteria. Seven of the differentially expressed miRNA were further analyzed by quantitative real-time reverse transcription PCR in a representative sample consisting of 57 schizophrenia cases and 34 controls. miRNA expression was normalized to the geometric mean of two constitutively expressed snoRNAs U44 and U49. These results were highly concordant with the microarray analysis with all seven of the miRNAs confirmed to be significantly downregulated in schizophrenia compared with controls (Figure 1b) (two-tailed Mann–Whitney U-test). These included miR-31 (2.97-fold, *P* < 0.001), miR-431 (2.79-fold, P < 0.001), miR-433 (2.06-fold, P = 0.017),miR-107 (2.25-fold, P<0.001), miR-134 (1.89-fold,

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P=0.007), miR-99b (4.24-fold, *P*<0.001) and miR-487b (2.06-fold, *P*<0.001).

Cluster analysis of schizophrenia-associated miRNA

The miRNAs shown to be associated with schizophrenia in the microarray analysis were also subjected to hierarchical clustering (Figure 2). This segregated the participants into two main groups in which reduced expression was enriched within the schizophrenia cohort (Figure 2, left side) in contrast to the majority of controls, which were visually distinguished by their relatively higher expression (Figure 2, right side). Strikingly, a cluster of 17 of the most substantially downregulated miRNAs were also structurally associated by their genomic position on the long arm of chromosome 14 (14q32). These miRNA all reside within two closely neighboring segments each spanning about 40 kb and separated by ~110 kb, at 14q32.2 and 14q32.31 (Figure 3). This cluster encodes at least 48 miRNA genes, 56,57 of which 30 mature miRNAs were expressed in the PBMCs. This indicated that approximately 53% of the miRNAs expressed at this locus were significantly downregulated in schizophrenia. Moreover, when we examined miRNAs expressed in this cluster that were not significantly altered after correction for multiple testing, a further 10 miRNAs or 33% (including miR-493*, miR-665, miR-379, miR-411, miR-376a, miR-376b, miR-376c, miR-495, miR-539, miR-889) also displayed an average reduction in schizophrenia consistent with the trend seen in the majority of miRNAs expressed from this region. This pattern of expression among the clustered miRNAs associated with schizophrenia was also highly consistent within individuals, suggesting that their transcription may be under the influence of shared regulatory mechanisms.

Copy-number analysis

To investigate whether the downregulation of miRNA on chromosome 14q32.31–32.2 is related to structural variation, we analyzed a subset of samples including 57 controls and 81 cases using commercially available TaqMan copy-number assays (Applied Biosystems). Amplicons specific for sequence within maternally expressed gene 3, cluster A and cluster B of the 14q32 region, were analyzed relative to the endogenous control RNase-P. No significant deletion or amplification events were detected in this region in either cases or controls (Supplementary Figure 1).

Target gene and pathway analysis

To gain some appreciation of the biological implications of the schizophrenia-associated changes in miRNA expression, we examined predicted miRNA targets and their associated pathways to see whether any patterns emerged. The more miRNA binding sites a gene has within its 3'-untranslated region, the more likely it is to be post-transcriptionally regulated.⁵³ The target prediction algorithms miRanda⁵¹ and TargetScan⁵² are capable of identifying those miRNA

 Table 2
 miRNA significantly downregulated in schizophrenia compared with non-psychiatric controls

miRNA ID	Fold change	q-Value	Chromosomal coordinates
hsa-miR-329	0.753	0	hsa-miR-329-1 14: 101 493 122–101 493 201 [+] hsa-miR-329-2 14: 101 493 437–101 493 520 [+]
hsa-miR-31	0.754	0	9: 21 512 114–21 512 184 [–]
hsa-miR-409-3p	0.757	0	14: 101 531 637–101 531 715 $[+]$
hsa-miR-224	0.762	0	X: 151 127 050–151 127 130 [–]
hsa-miR-432	0.763	0	14: 101 350 820–101 350 913 $[+]$
hsa-miR-487b	0.764	0	14: 101 512 792–101 512 875 [+]
hsa-miR-134	0.764	0	14: 101 521 024–101 521 096 [+]
hsa-miR-431 hsa-miR-150*	0.765	0 0	14: 101 347 344–101 347 457 [+]
hsa-miR-150" hsa-miR-99b	$0.771 \\ 0.783$	0	$\begin{array}{c} 19:\ 50\ 004\ 042 {=} 50\ 004\ 125\ [{-}]\\ 19:\ 52\ 195\ 865 {=} 52\ 195\ 934\ [+] \end{array}$
hsa-miR-1275	0.798	0	13. 32 133 803 - 32 133 334 [+] 6: 33 967 749 - 33 967 828 [-]
hsa-miR-335*	0.799	0	7: 130 135 952 - 130 136 045 [+]
hsa-miR-200c	0.813	0	12:7072862-7072929[+]
hsa-miR-486-3p	0.815	0	8: 41 517 959–41 518 026 [–]
hsa-miR-29b-1*	0.817	0	7: 130 562 218–130 562 298 [–]
hsa-miR-16-2*	0.821	0	3:160122533-160122613[+]
hsa-miR-877	0.83	0	6: 30552109– 30552194 [+]
hsa-miR-107	0.835	0	10: 91352504– $91352584[-]$
hsa-miR-130b*	0.839	0	22: 22 007 593–22 007 674 [+]
hsa-miR-544	0.843	0	14: 101 514 995–101 515 085 [+]
hsa-miR-342-5p	0.844	0	14: 100 575 992–100 576 090 [+]
hsa-miR-148b	0.844	0	12: 54 731 000–54 731 098 [+]
hsa-miR-625*	0.856	0 0	14:65937820-65937904[+]
hsa-miR-28-3p hsa-miR-576-5p	$\begin{array}{c} 0.858 \\ 0.86 \end{array}$	0	$3: 188\ 406\ 569-188\ 406\ 654\ [+]$ $4: 110\ 409\ 854-110\ 409\ 951\ [+]$
hsa-miR-151-3p	0.864	0	8: 141 742 663–141 742 752 [–]
hsa-miR-28-5p	0.869	0	3: 188 406 569 - 188 406 654 [+]
hsa-miR-664	0.871	0	1: 220 373 880–220 373 961 $[-]$
hsa-miR-128	0.874	0	hsa-miR-128-1 2: 136 422 967–136 423 048 [+]
			hsa-miR-128-2 3: 35 785 968–35 786 051 [+]
hsa-miR-584	0.879	0	5: 148441876– 148441972 [–]
hsa-miR-574-3p	0.883	0	4: 38869653– $38869748[+]$
hsa-miR-181a	0.913	0	hsa-miR-181a-1 1: 198 828 173–198 828 282 [–] hsa-miR-181a-2 9: 127 454 721–127 454 830 [+]
hsa-miR-30e*	0.917	0	1: 41 220 027–41 220 118 $[+]$
hsa-miR-433	0.664	1.555	14: 101 348 223–101 348 315 [+]
hsa-miR-654-5p	0.73	1.555	14: 101 506 556-101 506 636 [+]
hsa-miR-193b hsa-miR-485-3p	0.806	1.555	16: 14 397 824–14 397 906 [+]
hsa-miR-465-5p hsa-miR-370	$0.816 \\ 0.831$	$1.555 \\ 1.555$	14: 101 521 756–101 521 828 [+] 14: 101 377 476–101 377 550 [+]
hsa-miR-340*	0.854	1.555	5: 179 442 303–179 442 397 [–]
hsa-miR-1271	0.854	1.555	5: 175 794 949 - 175 795 034 [+]
hsa-miR-151	0.855	1.555	8: 141 742 663–141 742 752 [–]
hsa-miR-15b*	0.86	1.555	3: 160 122 376–160 122 473 [+]
hsa-miR-502-3p, hsa-miR-500*	0.86	1.555	X: 49779206–49779291 [+]
hsa-miR-27b	0.905	1.555	9: 97 847 727–97 847 823 [+]
hsa-miR-199a-3p, hsa-miR-199b-3p	0.922	1.555	hsa-miR-199a-1 19: 10 928 102–10 928 172 [–]
			hsa-miR-199a-2 1: 172 113 675–172 113 784 [–]
	0.5		hsa-miR-199b 9: 131007000–131007109 [–]
hsa-miR-151-5p	0.933	1.555	8: 141 742 663-141 742 752 [-]
hsa-miR-146a	0.95	1.555	5: 159 912 359–159 912 457 [+]
hsa-miR-21 hsa-miR-30d	$0.957 \\ 0.959$	$1.555 \\ 1.555$	17: 57 918 627–57 918 698 $[+]$ 8: 135 817 119–135 817 188 $[-]$
hsa-miR-127-3p	0.939	2.131	14: 101349316 - 101349412 [+]
hsa-miR-98	0.82	2.131	X: 53 583 184–53 583 302 [–]
hsa-miR-328	0.864	2.131	$16: 67\ 236\ 224-67\ 236\ 298\ [-]$
hsa-miR-181b	0.875	2.131	hsa-miR-181b-1 1: 198 828 002–198 828 111 [–]
			hsa-miR-181b-2 9: 127 455 989–127 456 077 [+]
hsa-miR-378	0.882	2.131	5: 149 112 388–149 112 453 [+]
hsa-miR-150	0.963	2.131	19:50004042– $50004125[-]$
hsa-miR-323-3p	0.796	2.74	14: 101 492 069–101 492 154 $[+]$
hsa-miR-874	0.797	2.74	5: 136 983 261–136 983 338 [–]
hsa-miR-330-3p	0.857	2.74	19: 46 142 252–46 142 345 [-]
hsa-miR-500	0.862	2.74	X: 49773039–49773122 [+]
hsa-miR-181a-2*	0.878	2.74	9: 127 454 721–127 454 830 [+]

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Table 2	Continued
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miRNA ID	Fold change	q-Value	Chromosomal coordinates				
hsa-miR-146b-5p	0.912	2.74	10: 104 196 269–104 196 341 [+]				
hsa-let-7b	0.952	2.74	22: 46509566 – $46509648[+]$				
hsa-miR-25	0.956	2.74	7: 99691183–99691266 [–]				
hsa-miR-92a	0.957	2.74	hsa-miR-92a-1 13: 92 003 568–92 003 645 [+]				
hsa-miR-410	0.812	3.288	hsa-miR-92a-2 X: 133 303 568–133 303 642 [–] 14: 101 532 249–101 532 328 [+]				
hsa-miR-221*	0.812	3.288	X: 45 605 585–45 605 694 [-]				
hsa-miR-942	0.825	3.288	1: $117637265-117637350[+]$				
hsa-miR-664*	0.845	3.288	1: 220 373 880–220 373 961 [–]				
hsa-miR-20b	0.865	3.288	X: 133 303 839–133 303 907 [–]				
hsa-miR-628-3p	0.867	3.288	15: 55 665 138-55 665 232 [-]				
hsa-miR-152	0.892	3.288	17: 46114527-46114613[-]				
hsa-let-7d	0.957	3.288	9: 96 941 116-96 941 202 [+]				
hsa-miR-154	0.766	4.484	14: 101 526 092–101 526 175 $[+]$				
hsa-miR-337-3p	0.87	4.484	14: 101 340 830–101 340 922 [+]				
hsa-miR-505	0.884	4.484	X: 139006307-139006390 [-]				
hsa-miR-625	0.897	4.484	14: 65937820 – 65937904 [+]				
hsa-miR-22*	0.898	4.484	17: 1617197–1617281 [–]				
hsa-let-7g	0.965	4.484	3:52302294– $52302377[-]$				
hsa-miR-1301	0.869	4.853	2: 25 551 509–25 551 590 [-]				
hsa-let-7d*	0.905	4.853	9:96941116-96941202[+]				
hsa-miR-766	0.93	4.853	X: 11 8780 701–118 780 811 [–]				
hsa-let-7a	0.971	4.853	hsa-let-7a-1 9: 96 938 239–96 938 318 [+]				
			hsa-let7a-2 11: 12 2017 230–122 017 301 [–]				
			hsa-let7a-3 22: 46 508 629–46 508 702 [+]				

Abbreviation: miRNA, microRNA.

Differentially expressed miRNA were ranked by significance and then by fold change. miRNA transcribed from the genomic region encompassing 14q32.2–q32.31 were identified with bold font. *Denotes star form of the mature miRNA.

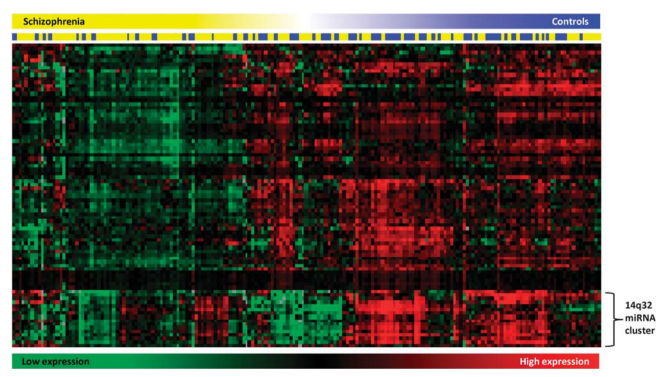


Figure 2 Heatmap showing hierarchical clustering of significantly downregulated microRNA with diagnosis. Samples are color coded according to diagnosis (blue = control; yellow = schizophrenia). Green indicates low expression and red indicates high expression (Java Treeview).

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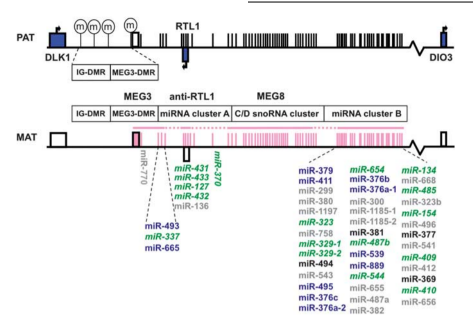


Figure 3 Genomic organization of 14q32 microRNA (miRNA) clusters neighboring the DLK1-DIO3 imprinted domain. DLK1, RTL1 and DIO3 (blue rectangles) are paternally expressed genes. The maternally expressed non-coding RNA genes (pink) include maternally expressed gene 3 (MEG3), anti-RTL1 (encodes the smaller miRNA cluster A), maternally expressed gene 8 (MEG8) (encodes small nucleolar RNA (snoRNA) cluster) and the larger miRNA cluster B. The intergenic germline derived differentially methylated region (IG-DMR) is methylated on the paternal chromosome denoted by the encircled letter m. The 14q32 miRNA clusters are arranged in two segments: each separated by a C/D snoRNA cluster. miRNA in dark green italic font were significantly downregulated in schizophrenia and miRNA in blue exhibited a trend for downregulation in schizophrenia. miRNA in black were expressed but not differentially expressed, and in gray were not expressed. This figure was adapted from Royo and Cavaille⁶⁴ and Hagan *et al.*¹⁰³

that bind multiple times within the one 3'-untranslated region. As such, this information was used to predict target genes that were most likely to be heavily regulated by the differentially expressed miRNA. Owing to a high degree of false positives, the list of target genes was filtered to only include those genes that were targeted by five or more miRNA, or contained five or more binding sites for a particular miRNA within its 3'-untranslated region. This target analysis process was performed on the 17 miRNA displaying altered expression in the 14q32 region. The miRanda and TargetScan lists were then combined to form a union of targets predicted by both algorithms in addition to a more stringent analysis involving the intersection of targets identified by both miRanda and TargetScan. Selected genes were then collectively subjected to pathway analysis using the Gene Set Analysis Toolkit V.2 from WebGestalt (http://bioinfo.vanderbilt.edu/webgestalt).54 The union of miRanda and TargetScan revealed a number of significantly enriched pathways with relevance to neural connectivity and synaptic plasticity, including axon guidance, long-term potentiation, focal adhesion, neurotrophin, ErbB, calcium and mitogenactivated protein kinase signaling (Table 3 and Supplementary Figure 2). Furthermore, the more stringent intersection approach, using genes predicted to be targets by both algorithms, also suggested the involvement of neurologically relevant pathways

as well as pathways relevant to immune system function (Table 4 and Supplementary Figure 3).

Discussion

A subset of brain-enriched miRNA genes located within a large imprinted cluster on chromosome 14q32 were differentially expressed in schizophrenia

Analysis of post-mortem miRNA expression has previously reported schizophrenia-associated changes in both the superior temporal gyrus and prefrontal cortex.^{24,47,58} Although miRNA expression typically displays a stage- and tissue-specific expression pattern, we identified several brain-enriched miRNAs including miR-134, miR-128 and miR-181b that were also differentially expressed in PBMCs from individuals with schizophrenia. It is plausible that these represent a neurologically relevant signature reflecting the prevailing neuropsychiatric phenotype. The reduction of miR-134 was particularly significant not only because of its role in brain development and function, 28,59,60 but also as a consequence of its position in a cluster of differentially expressed miRNAs on chromosome 14q32.

A large proportion of the miRNAs found to be differentially expressed in schizophrenia are encoded within this large cluster on chromosome 14q32, which collectively comprises over 5% of known human miRNA genes^{61–63} and about 40–50% of all

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Table 3 Pathways enriched with target genes predicted by a union of miRanda and TargetScan target lists for the 14q32 cluster

Category	Term	n	%	P-value	Benjamini
KEGG_PATHWAY	Pathways in cancer	108	0.3	< 0.000	< 0.000
KEGG_PATHWAY	Axon guidance	54	0.1	< 0.000	< 0.000
KEGG_PATHWAY	Adherens junction	36	0.1	< 0.000	< 0.000
KEGG_PATHWAY	Regulation of actin cytoskeleton	72	0.2	< 0.000	< 0.000
KEGG_PATHWAY	Focal adhesion	68	0.2	< 0.000	< 0.000
KEGG_PATHWAY	Chronic myeloid leukemia	32	0.1	< 0.000	< 0.000
KEGG_PATHWAY	Insulin signaling pathway	48	0.1	< 0.000	< 0.000
KEGG_PATHWAY	Renal cell carcinoma	30	0.1	< 0.000	< 0.000
KEGG_PATHWAY	ErbB signaling pathway	34	0.1	< 0.000	0.001
KEGG_PATHWAY	MAPK signaling pathway	79	0.2	< 0.000	0.001
KEGG_PATHWAY	Pancreatic cancer	29	0.1	< 0.000	0.0016
KEGG_PATHWAY	Ubiquitin-mediated proteolysis	46	0.1	< 0.000	0.0018
KEGG_PATHWAY	Wnt signaling pathway	49	0.1	< 0.000	0.0024
KEGG_PATHWAY	Prostate cancer	33	0.1	< 0.000	0.0023
KEGG_PATHWAY	Long-term potentiation	27	0.1	< 0.000	0.0028
KEGG_PATHWAY	Glioma	25	0.1	< 0.000	0.0049
KEGG_PATHWAY	Neurotrophin signaling pathway	41	0.1	< 0.000	0.0046
KEGG_PATHWAY	Colorectal cancer	30	0.1	< 0.000	0.0078
KEGG_PATHWAY	mTOR signaling pathway	21	0.1	0.0011	0.0110
KEGG_PATHWAY	Endocytosis	54	0.1	0.0012	0.0110
KEGG_PATHWAY	Melanogenesis	33	0.1	0.0015	0.0130
KEGG_PATHWAY	Calcium signaling pathway	50	0.1	0.0039	0.0320
KEGG_PATHWAY	Non-small-cell lung cancer	20	0.1	0.0047	0.0370
KEGG_PATHWAY	Progesterone-mediated oocyte maturation	28	0.1	0.0053	0.0400
KEGG_PATHWAY	TGF-β signaling pathway	28	0.1	0.0063	0.0460
KEGG_PATHWAY	Melanoma	24	0.1	0.0064	0.0450
KEGG_PATHWAY	T-cell receptor signaling pathway	33	0.1	0.0067	0.0450
KEGG_PATHWAY	Jak-STAT signaling pathway	44	0.1	0.0070	0.0450
KEGG_PATHWAY	Dilated cardiomyopathy	29	0.1	0.0073	0.0460
KEGG_PATHWAY	HCM	27	0.1	0.0089	0.0540
KEGG_PATHWAY	Oocyte meiosis	33	0.1	0.0090	0.0530

Abbreviations: HCM, hypertrophic cardiomyopathy; KEGG, Kyoto Encyclopedia of Genes and Genomes; MAPK, mitogenactivated protein kinase; mTOR, mammalian target of rapamycin; TGF, tumor growth factor.

Enriched KEGG pathways identified by WebGestalt gene set analysis using predicted target genes. The list was compiled from an union of the miRanda and TargetScan predicted targets of 17 members of the miR-14q32 cluster differentially expressed in schizophrenia. *n*, number of input genes in pathway (filtered to only show pathways represented by \geq 7 target genes). %, percentage of total target genes analyzed (~3600 gene IDs) that are represented in this pathway.

known eutherian-specific miRNA.⁶⁴ Significantly, this locus on 14q32, known as the DLK1-DIO3 region, is imprinted such that the associated miRNA cluster is only expressed from the maternal chromosome.⁶⁴ In mice, the syntenic region on chromosome 12 also resides within an imprinted domain expressed exclusively from the maternal chromosome.⁵⁷ The miRNAs are encoded by two clusters located in close proximity to each other, the smaller of the two at 14q32.2 contains 10 miRNA, which reside within the retrotransposon-like gene RTL1 (in antisense), whereas the larger cluster also known as the miR379-410 cluster contains approximately 40miRNAs. Although the nature of their organization is not entirely clear in human beings, some of the miRNA in this region could be processed from a single large polycistron such as Mirg (miRNA containing gene, also known as Meg9), which appears to be the case in the mouse. However, it seems more likely

that the majority of miRNA in this cluster are encoded by tandem arrays of related intronic sequences.^{57,65}

Regulation of the 14q32 miRNA cluster

As more than 85% of the miRNAs expressed in this region showed a trend for downregulation in schizophrenia, it is reasonable to suggest that these molecules may be co-regulated by a shared mechanism associated with the disorder. This could relate to a number of common factors that affect transcriptional activity or miRNA processing in this region. Why the remaining miRNAs expressed from this cluster were neutral with respect to disease status remains to be determined, but it could be that the genes encoding these molecules are responding more to differences in their local transcriptional environment, which over-ride the prevailing cluster-wide influence. They may also display isolated differences in their splicing or primary transcript processing,

Table 4	Pathways	enriched	with	target	genes	predicted	by	an	intersection	of	miRanda	and	TargetScan	target	lists for	the
14q32 clu					0	-							0	0		

Category	Term	n	%	P-value	Benjamini
KEGG_PATHWAY	Pathways in cancer	31	3.8	< 0.0001	< 0.0001
KEGG_PATHWAY	Prostate cancer	13	1.6	< 0.0001	< 0.0001
KEGG_PATHWAY	MAPK signaling pathway	21	2.6	< 0.0001	< 0.0001
KEGG_PATHWAY	Focal adhesion	18	2.2	< 0.0001	< 0.0001
KEGG_PATHWAY	Melanoma	11	1.4	< 0.0001	< 0.0001
KEGG_PATHWAY	Adherens junction	11	1.4	< 0.0001	< 0.0001
KEGG_PATHWAY	Endocytosis	16	2.0	< 0.0001	< 0.0001
KEGG_PATHWAY	mTOR signaling pathway	9	1.1	< 0.0001	< 0.0001
KEGG_PATHWAY	Non-small-cell lung cancer	9	1.1	< 0.0001	< 0.0001
KEGG_PATHWAY	Long-term potentiation	10	1.2	< 0.0001	< 0.0001
KEGG_PATHWAY	Chronic myeloid leukemia	10	1.2	< 0.0001	< 0.0001
KEGG_PATHWAY	Insulin signaling pathway	13	1.6	< 0.0001	< 0.0001
KEGG_PATHWAY	Regulation of actin cytoskeleton	16	2.0	< 0.0001	< 0.0001
KEGG_PATHWAY	Glioma	9	1.1	< 0.0001	< 0.0001
KEGG_PATHWAY	Progesterone-mediated oocyte maturation	10	1.2	< 0.0001	< 0.0001
KEGG_PATHWAY	ErbB signaling pathway	10	1.2	< 0.0001	< 0.0001
KEGG_PATHWAY	Renal cell carcinoma	9	1.1	< 0.0001	< 0.0001
KEGG_PATHWAY	Jak-STAT signaling pathway	13	1.6	< 0.0001	< 0.0001
KEGG_PATHWAY	Pancreatic cancer	9	1.1	< 0.0001	< 0.0001
KEGG_PATHWAY	Neurotrophin signaling pathway	11	1.4	< 0.0001	< 0.0001
KEGG_PATHWAY	Colorectal cancer	9	1.1	< 0.0001	< 0.0001
KEGG_PATHWAY	Small-cell lung cancer	9	1.1	< 0.0001	< 0.0001
KEGG_PATHWAY	Calcium signaling pathway	13	1.6	< 0.0001	< 0.0001
KEGG_PATHWAY	TGF-β signaling pathway	9	1.1	< 0.0001	< 0.0001
KEGG_PATHWAY	Dilated cardiomyopathy	9	1.1	< 0.0001	< 0.0001
KEGG_PATHWAY	Phosphatidylinositol signaling system	8	1.0	< 0.0001	0.0001
KEGG_PATHWAY	Fcγ R-mediated phagocytosis	9	1.1	< 0.0001	0.0001
KEGG_PATHWAY	Acute myeloid leukemia	7	0.9	< 0.0001	0.0002
KEGG_PATHWAY	HCM	8	1.0	0.0001	0.0002
KEGG_PATHWAY	Melanogenesis	9	1.1	< 0.0001	0.0002
KEGG_PATHWAY	Axon guidance	10	1.2	0.0001	0.0002
KEGG_PATHWAY	Apoptosis	8	1.0	0.0002	0.0003
KEGG_PATHWAY	Neuroactive ligand–receptor interaction	14	1.7	0.0002	0.0003
KEGG_PATHWAY	Vascular smooth muscle contraction	9	1.1	0.0002	0.0003
KEGG_PATHWAY	Long-term depression	7	0.9	0.0003	0.0004
KEGG_PATHWAY	Wnt signaling pathway	10	1.2	0.0004	0.0006
KEGG_PATHWAY	ARVC	7	0.9	0.0004	0.0006
KEGG_PATHWAY	Fce RI signaling pathway	7	0.9	0.0005	0.0007
KEGG_PATHWAY	GnRH signaling pathway	8	1.0	0.0005	0.0007
KEGG_PATHWAY	Tight junction	9	1.1	0.0007	0.0009
KEGG_PATHWAY	Chemokine signaling pathway	11	1.4	0.0007	0.0009
KEGG_PATHWAY	T-cell receptor signaling pathway	8	1.0	0.0007	0.0009
KEGG_PATHWAY	Ubiquitin-mediated proteolysis	9	1.1	0.0009	0.0011
KEGG_PATHWAY	Oocyte meiosis	8	1.0	0.001	0.0012
KEGG_PATHWAY	Gap junction	7	0.9	0.0012	0.0014
KEGG_PATHWAY	Leukocyte transendothelial migration	8	1.0	0.0013	0.0015
KEGG_PATHWAY	Natural killer cell-mediated cytotoxicity	8	1.0	0.0033	0.0037
KEGG_PATHWAY	Lysosome	7	0.9	0.0052	0.0056
KEGG_PATHWAY	Huntington's disease	9	1.1	0.0063	0.0067
KEGG_PATHWAY	CAMs	7	0.9	0.0106	0.011
KEGG_PATHWAY	Cytokine–cytokine receptor interaction	10	1.2	0.0227	0.0231
KEGG_PATHWAY	Metabolic pathways	29	3.6	0.0268	0.0268

Abbreviations: ARVC, arrhythmogenic right ventricular cardiomyopathy; CAM, cell adhesion molecules; GnRH, gonadotropin-releasing hormone; HCM, hypertrophic cardiomyopathy; KEGG, Kyoto Encyclopedia of Genes and Genomes; MAPK, mitogen-activated protein kinase; mTOR, mammalian target of rapamycin.

Enriched KEGG pathways identified by WebGestalt gene set analysis using predicted target genes. The list was compiled from an intersection of the miRanda and TargetScan predicted targets of 17 members of the miR-14q32 cluster differentially expressed in schizophrenia. n, number of input genes in pathway (filtered to only show pathways represented by ≥ 7 target genes). %, percentage of total target genes analyzed (~810 gene IDs) that are represented in this pathway.

or pre-miRNA cleavage and maturation. It is also possible that the schizophrenia-associated regulatory influence seen for most of the expressed miRNA extends beyond the miRNA clusters to affect protein coding and other non-coding mRNAs in the region; however, there are no reports to this effect in the schizophrenia literature. A genome-wide study of MS from our laboratory suggests that the proteincoding genes in this domain (DLK1, RTL1, DIO3 and MEG3) are not expressed in lymphocytes.⁶⁶

Studies in the mouse suggest that the expression of miRNA clustered on 14q32 in human beings may be particularly sensitive to changes in the miRNA biogenesis pathway. For example, DiGeorge syndrome critical region gene 8 (DGCR8) heterozygous knockout mice were shown to have significantly reduced levels of mature miRNA expression from the syntenic region on chromosome 12 in both the prefrontal cortex and hippocampus.⁶⁷ The product of this gene, one of many depleted in the 22q11 microdeletion syndromes, is an important component of the microprocessor complex involved in pri-miRNA cleavage in the nucleus. Interestingly, these miRNA were also over represented ($\sim 25\%$) among miRNA showing downregulation in dopaminergic neurons from the striatum of Ago2-deficient mice.⁶⁸ A large proportion of miRNA from this cluster also contain primary and secondary structural features associated with dicerindependent/Ago2-slicer activity-dependent processing.⁶⁹ This includes the preponderance for an A/U base at the 5' terminal in the mature sequence (80%)and the preference for miRNA residing in the 3p side of precursor hairpins (\sim 70%).^{68,70} This suggested that Ago2-dependent pre-miRNA processing^{69,71} is particularly important for the biogenesis of miRNA in this cluster and may be associated with changes we observed in schizophrenia. Another possibility is that a 14g32 cluster-associated transcription factor or other regulatory molecule is responsible for the change observed in schizophrenia. One example of this is the Ca²⁺-activated transcription factor known as myocyte enhancer factor-2 (MEF2), which has been shown to positively regulate expression of miRNA in this cluster.⁵⁶ MEF2 isoforms have also been shown to regulate neuronal development through their role in supporting newly differentiated neurons and synapse formation.⁷²⁻⁷⁶ We have also shown that expression of the MEF2D isoform is upregulated in differentiating neurons in response to a reduction in miR-17 family miRNA expression.77 Lower expression of MEF2 in PBMC could potentially contribute to the reduction in transcription of this cluster in schizophrenia.

Function of 14q32 miRNA cluster in brain function and development

The miR-379/410 cluster has been shown to follow a similar stage and tissue-specific expression pattern as a colocalized cluster of C/D snoRNAs derived from a single intronic transcript known as maternally expressed gene 8 (RNA imprinted and accumulated

in the nucleus on mouse chromosome 12).⁶² This is detectable throughout the developing mouse embryo and the placenta, is enriched in the brain in adult mice^{29,57,62,78} and may be significant for the development of higher brain functions in eutherian mammals.⁷⁹ The neural function of this cluster is supported by recent research suggesting that coordinated expression of these miRNA, including miR-134 and miR-329, is important for brain function and development. Neuronal activity in rats trigger MEF2, a neuronal-activity Ca2+-dependent transcription factor (described above), to bind to sequence upstream of the larger of the two clusters and induce expression of all pre-miRNAs within it.⁵⁶ At least three members of the co-regulated cluster, including miR-329 and miR-134, that we see altered in schizophrenia, appear to regulate hippocampal neuron dendrite morphology.⁵⁶ MiR-134 has also been shown repress dendritic development by targeting to LimK1²⁸ and to activate it through targeting Pumilio2.⁵⁶ Interestingly, miR-134 expression is upregulated in the rat hippocampus and amygdala following acute stress, but is decreased during chronic stress.⁶⁰ A predicted target of miR-134, splicing factor SC35, has been shown to be upregulated after stress and is believed to promote alternative splicing of acetylcholinesterase mRNA. Therefore, it is possible that miR-134 and other related miRNAs from the 14q32 cluster function as stress-dependent regulators of neurotransmission and synaptic plasticity. This is supported by the pathway analysis of target genes predicted for differentially expressed miRNA in this cluster, which included axon guidance, regulation of the actin cytoskeleton, long-term potentiation, long-term depression, neuroactive ligand-receptor interaction, focal adhesion, neurotrophin, mammalian target of rapamycin, calcium, mitogen-activated protein kinase and ErbB signaling pathways. It was also broadly consistent with previous pathway analysis of the human 14q32 cluster and the syntenic miRNA c luster in the mouse, with enrichment of target genes involved in neurogenesis (including the biological processes in axon guidance), embryonic development and more general developmental processes.⁷⁹ Another interesting theme emerging from the pathway analysis was the large number of immunologically relevant processes including endocytosis, Fcy R-mediated phagocytosis, Fcc RI signaling pathway, chemokine signaling pathway, T-cell receptor signaling pathway, leukocyte transendothelial migration and natural killer cell-mediated cytotoxicity. Given the evidence suggesting a link between early environmental exposure to infection/inflammation and schizophrenia, the alteration of miRNA expression in this tissue and their influence in these pathways could have functional significance for the immune system in individuals with the disorder. The expression of miRNA in this cluster was also shown to be highly associated with cellular pluripotency.⁸⁰ If this cluster is involved in the regulation and maintenance of stem cell behavior, the reduction observed in schizophrenia

could have broad developmental implications for both the immune and nervous systems.

Differentially expressed miRNA are contained within an imprinted locus

Maternal specific expression of this region is regulated by imprinting centers defined by an intergenic germline derived differentially methylated region (DMR) and the post-fertilization derived MEG3-DMR (shown in Figure 3).⁸¹ Differential expression of miRNA clustered in this imprinted region raises the possibility that there is a schizophrenia-associated parent of origin effect influencing the miRNA expression from this cluster similar to that observed at a number of other genes including the 5-hydroxytryptamine type 2A receptor⁸² and GABA_A receptor $\beta 2.^{83}$ Imprinting at the 14q32 region has also been implicated in bipolar disorder^{84,85} and anxiety.⁸⁶ Although no gender bias was observed in the expression of the miRNA cluster, specific details about participants' parental disease status were not sufficient to derive parent of origin effect.

Many characteristics of the imprinted locus on 14q32 parallel the genetic landscape at 15q11-13, known for its involvement in Prader-Willi Syndrome). This neurodevelopmental disorder arises from monoallelic expression at 15q11-13 as a result of deletion of the paternal copy of the imprinted region (delPWS), maternal uniparental disomy (mat-upd-15), as well as abnormal imprinting (reviewed by Buiting⁸⁷). The 14q32 and 15q11-13 regions are similar in that they are both developmentally regulated via an imprinting center, and contain both maternally and paternally expressed genes. The imprinted regions of both loci also encode C/D snoRNAs that function as guides for epigenetic modification of other RNAs. The snoRNAs in these loci are atypical in that they lack complementarity to their neighboring RNA. These small non-coding RNA molecules, exclusive to eutherians, are also brain enriched and transcribed in the same direction, possibly by a single transcription unit.62,64,88-90 In further support of a role for these imprinted loci in neuronal development, Leung et al.⁹⁰ observed developmentally regulated chromatin decondensation of snoRNA clusters at 15q11-13 and 14q32 in postnatal mouse and human neurons.

Individuals with Prader-Willi Syndrome also share some phenotypic features with schizophrenia such as cognitive deficits, behavioral problems and, in some cases, autistic symptoms and psychosis.91-97 The 14q32 locus has also been implicated in other disorders with impaired development and behavioral phenotypes.98 Rare copy-number variations at 14q32 and 15q11-13 have also been reported in schizophrenia.⁹⁹⁻¹⁰² Although our analysis of copy number in this region did not identify structural variants, this genomic region clearly has the potential to convey risk for neurodevelopmental disorders including schizophrenia.

Conclusions

In this study, we examined schizophrenia-associated miRNA expression and identified a significant correlation between miRNAs clustered in a small imprinted domain on 14q32. Although the cause of this anomaly is not known, it may be due to an alteration in transcriptional regulation or processing. The miRNAs encoded in this cluster evolved in eutherian mammals and appear to have a significant role in brain development and function. And while the significance of their downregulation in PBMCs in schizophrenia is less clear, there are implications for lymphocyte function that could contribute to the risk for the disorder. Alternatively, this phenomenon may be a vestige of broader alteration to post-transcriptional regulation, with more relevance to neurodevelopment. Regardless of the functional significance, the miRNA expression signatures observed in PBMCs may have the potential to serve as biomarkers of schizophrenia and associated phenotypes.

Conflict of interest

The authors declare no conflict of interest.

Acknowledgments

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