MicroRNA and gene co-expression networks characterize biological and clinical behavior of rhabdomyosarcomas

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Abstract

Rhabdomyosarcomas (RMS) in children and adolescents are heterogeneous sarcomas broadly defined by skeletal muscle features and the presence/absence of PAX3/7-FOXO1 fusion genes. MicroRNAs are small non-coding RNAs that regulate gene expression in a cell context specific manner. Sequencing analyses of microRNAs in 64 RMS revealed expression patterns separating skeletal muscle, fusion gene positive and negative RMS. Integration with parallel gene expression data assigned biological functions to 12 co-expression networks/modules that reassuringly included myogenic roles strongly correlated with microRNAs known in myogenesis and RMS development. Modules also correlated with clinical outcome and fusion status. Regulation of microRNAs by the fusion protein was demonstrated after PAX3-FOXO1 reduction, exemplified by miR-9-5p. MiR-9-5p levels correlated with poor outcome, even within fusion gene positive RMS, and were higher in metastatic versus non-metastatic disease. MiR-9-5p reduction inhibited RMS cell migration. Our findings reveal microRNAs in a regulatory framework of biological and clinical significance in RMS.

Keywords: MicroRNAs, Next generation sequencing, Co-expression modules, Rhabdomyosarcoma, Fusion protein

Introduction

Rhabdomyosarcomas (RMS) are the most frequent soft tissue sarcoma in children and RMS cells resemble those in the early stages of myogenic differentiation [1,2]. Histologically there are two major subtypes, embryonal (ERMS) accounting for 60–70% of cases and alveolar (ARMS). Approximately 70% of ARMS harbor fusion genes that join the 5′ and 3′ UTR of PAX3 or PAX7 to the 3′ sequence of FOXO1 gene [3,4]. The encoded fusion proteins play an oncogenic role in the development of RMS in association with co-operating events, notably overexpression of MYCN that is a transcriptional target of the fusion protein and genomically amplified in approximately 20% of cases [5–8]. The presence of PAX3-FOXO1 has been correlated with aggressive clinical behavior in several studies [9–12] and gene expression profiles and genomic imbalances are similar in fusion gene negative ARMS and ERMS [10,13]. MicroRNAs (miRNAs) are small (20–22 nt) RNA molecules that can alter cellular processes including proliferation, differentiation and apoptosis [14]. MiRNAs inhibit protein synthesis of specific genes by degrading specific mRNA species or hindering translation. Specific mRNA degradation involves base-pairing to partially complementary sequences within the 3′ UTR [15] or other less frequently studied mechanisms including their binding to the 5′ UTR [16] and gene promoters regions [17]. Each miRNA potentially regulates hundreds of target gene products and it is suggested that the entire protein coding genome is regulated by miRNAs [14]. Individual miRNAs have been ascribed oncogenic or tumor suppressor roles in cancer and their roles can clearly be cell context specific [18]. Roles for individual miRNAs in RMS have been described such as miR-1, miR-206, miR-133a/b and miR-29 that is described in a differentiated myogenic state [19–25] and miR-29 that is described with a tumor suppressor role [26]. However, a comprehensive view of how miRNAs shape the gene expression and biological features of a large number of RMS has not been described but could contribute to identification of prognostic markers/signatures and ultimately therapeutic targets, as indicated by previous studies of other tumor types [27–29].
Here we derived miRNA expression profiling data from high-throughput sequencing analyses of 64 primary RMS samples and took a systems biology approach to integrate this with parallel gene expression profiling data. This allowed us to identify co-expression miRNA networks (modules) with assigned biological functions that correlated with both patient outcome data and the presence of PAX3-FOXO1. This led us to demonstrate that PAX3-FOXO1 regulates the expression of many miRNAs, including miR-9-5p that was indirectly altered by PAX3-FOXO1 and shown to be of clinical and functional relevance.

Materials and methods

Primary tumor samples and cell lines

Sixty-four samples (36 ARMS and 28 ERMS) were available from RMS patients as part of a previously described cohort [30]. Clinicopathological features of the cases used in this study for miRNA profiling are summarized in Supplementary Table S1A. Three normal skeletal muscle samples were also obtained. Samples from a similarly treated cohort of 154 RMS patients were used for miRNA validation analyses and their clinicopathological details are summarized in Supplementary Table S1B. RMS from U.K. centres were collected through the Children’s Cancer and Leukaemia Group (Local Research Ethics Committee protocol Nos. 1836 and 2015 and Multi-Regional Research Ethics Committee 06/4/71 with consent, where required). Human cell lines derived from embryonal (RH2, RH36, RD, RMSYM, RUCH2, RUCH3, CT10) and alveolar (RH3, RH4, RH5, RH18, RH41, RMS, SCMCRMZ, CW9019) were cultured and validated by DNA fingerprinted as previously described [7,31].

miRNA library preparation, sequencing and analyses

RNA extraction was performed using Trizol reagent following manufacturer’s instructions (Invitrogen – Carlsbad, CA). Size selection and library preparation was performed by Fasteris, Plan-les Ouates, Switzerland including duplicates for quality control using 5 μg of RNA in a maximum volume of 10–15 μl that was sent to the company. Briefly, small RNA fragments were purified by acrylamide gel purification and 3’ and 5’ adapters were ligated to the ends of single stranded small RNA molecules. Reverse transcription and PCR amplification was performed to generate the DNA colonies template library, which was diluted to 10 nM. The miRNA expression profiling was performed by high-throughput sequencing analysis using a Solexa/Illumina Genome Analyzer II. The samples were analyzed by grouping in pools of four per channel. Each sample specific sequence was then retrieved using a bar-coding system made of 4 nucleotides. The Solexa fasta-Q format sequence of each sample was aligned to 2 databases: mature miRNA sequence (miRBase release 21) and EST (NCBI). The mature miRNA sequence was extended with Ns at each end to allow the alignment with segments of RNA one or two nucleotides longer than the wild type sequence. All the alignments were performed using novoalign 2.06.09 with default parameters, after removing the 3’ adapter and cutting the sequence to 30 nt. Following alignment, the output files were imported into R for further analysis. First of all the sequence with low Solexa quality score (which did not passed the cut-off in the novoalign software) and those having less than 5 reads in at least 2 samples were filtered out from the analysis. When multiple alignments with the same score were detected, the sequence was randomly assigned to one of them. After the alignment each sequence was assigned to an RNA class based on Entrez Gene type. The variance of the counts data were stabilized using a power transformation function and normalized by applying a non-linear function (lowess). Differentially expressed miRNAs were identified by fitting a linear model (limma) and adjusting for multiple testing by computing FDR values from p-values using the Benjamini-Hochberg procedure (limma package). Clinical correlations between levels of miRNA expression and patient outcome data were performed by fitting a
Functional annotation, predicted by GeneGo software, of the miR/gene co-expression modules for RMS identi-

Table 1

| Module     | No. of genes | Go processes                                      | GeneGo networks                                      |
|------------|--------------|---------------------------------------------------|------------------------------------------------------|
| Black      | 116          | Epidermis development, Ectoderm development, Epithelial cell differentiation, Epithelium development | Immune response, Th17-derived cytokines, Cytoskeleton, Intermediate filaments, Inflammation, Amphoterin signaling, Chemotaxis |
| Blue       | 516          | Cellular macromolecule metabolic process, Regulation of metabolic process, Nucleic acid metabolic process, Regulation of gene expression | Signal Transduction, TGF-beta, GDF and Activin signaling, Signal Transduction, BMP and GDF signaling, Cell cycle, Mitosis, Cytoplasmic microtubules, Inflammation, Cytoskeleton, Microtubules, Chemotaxis |
| Brown      | 460          | Immune system process, Immune response, Defense response, Response to stimulus | Inflammation, Interferon signaling, Chemotaxis, Cell adhesion, Leukocyte chemotaxis, Immune response, Phagocytosis |
| Green      | 206          | M phase, Cell cycle phase, Mitosis, Nuclear division | Cell cycle, Core, Cell cycle, M phase, Cell cycle C2-M, Cytoskeleton, Spindle microtubules |
| Green/yellow | 41          | Regulation of atrial cardiomyocyte membrane depolarization, Keratan sulfate biosynthetic process, response to nickel ion, multicellular organismal process | Neurophysiological process, Long-term potentiation, Signal transduction, Neureptide signaling pathways, Reproduction, FSH-beta signaling pathway, Reproduction, Feeding and Neurohormone signaling, Cell adhesion, Cell-matrix interactions, Cell adhesion, Platelet-endothelium-leucocyte interactions, Development, EMT, Regulation of epithelial-to-mesenchymal transition, Development, Skeletal muscle development, Cell adhesion, Cell-matrix interactions, Proteolysis, ECM remodelling, Development, Cartilage development, Signal transduction, Cholecystokinin signaling |
| Magenta    | 79           | Tissue development, Wound healing, Extracellular matrix organization, Skeletal system development | Development, Hematopoiesis, Regulation of skeletal muscle differentiation, Development, Bone development, Development, Skeletal muscle development, Extracellular matrix organization, Bone development, Development, Skeletal muscle development, Development, Skeletal muscle development |
| Pink       | 103          | Extracellular matrix organization, Skeletal system development, positive regulation of dendritic spine development, positive regulation of smooth muscle cell chemotaxis, ANH: Anatomical structure morphogenesis | Development, EMT, Regulation of epithelial-to-mesenchymal transition, Development, Skeletal muscle development, Cell adhesion, Cell-matrix interactions, Proteolysis, ECM remodeling, Development, Cartilage development, Signal transduction, Cholecystokinin signaling |
| Purple     | 64           | Cardiac muscle contraction, Heart contraction, Striated muscle contraction, Muscle contraction | Cytoskeleton, Actin filaments, Development, Skeletal muscle, Development, Neuromuscular junction |
| Red        | 156          | Protein localization, Cellular process, Macromolecule localization, Protein transport | Immune response, Phagosome in antigen presentation, Transcription, mRNA processing, Cell cycle, Mitosis, Development, Hedgehog signaling |
| Tan        | 31           | Positive regulation of gene expression, Response to external stimulus, Positive regulation of transcription, DNA-dependent, Positive regulation of RNA metabolic process | Reproduction, Gondadotropin regulation, Immune response, Th17-derived cytokines, Inflammation, IL-6 signaling, Cell cycle, G1/S Interleukin regulation, Development, Neurogenesis in general, Development, Hedgehog signaling, Development, Blood vessel morphogenesis, Development, Ossification and bone remodeling |
| Turquoise  | 529          | Nervous system development, Developmental process, Anatomical structure development, Multicellular organismal development | Development, Hedgehog signaling, Development, Blood vessel morphogenesis, Development, Ossification and bone remodeling |
| Yellow     | 209          | Muscle system process, Muscle contraction, Striated muscle contraction, Myofibril assembly | Development, Skeletal muscle development, Cytoskeleton, Actin filaments, Cytoskeleton, Regulation of cytoskeleton rearrangement |
Fig. 2. Relationship between modules and with other clinic-molecular variables. WGCNA analysis identified 12 modules based on gene co-expression patterns. Each module was uniquely color labeled. The expression profile of the genes included in each module was summarized by a module eigengene (first principal component of the expression matrix). These eigengenes were used to assess relationships between modules as well as with other variables. A. Hierarchical clustering of module eigengenes and a heatmap showing the relatedness of these to each other. In the heatmap, red indicates modules are highly related and blue that they are not related. Modules that cluster closer together tend to be more related. In B, we averaged the statistics computed using the Kruskal-Wallis rank sum test of the genes within each module — higher values indicate the presence of a larger number of genes contained in that module which are associated to a specific patient-specific parameter. The horizontal bars mark the threshold level for significance. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
**Results**

**miRNA profiles in skeletal muscle and RMS**

MiRNA expression profiling was performed by high-throughput sequencing analysis of 68 samples (3 skeletal muscle, 1 myoblast and 64 RMS patient samples (Supplementary Table S1)). On average, around 2,069,412 tags were sequenced for each sample (range: 753,614–7,390,288). After sequence alignment and filtering, each tag was assigned to a RNA class based on Entrez Gene type and 78% and 73% were shown to be miRNAs in RMS and skeletal muscle, respectively, confirming their successful enrichment (Supplementary Fig. S1A). Since miRNAs are often included in transcription clusters, their expression is more coordinated when located in close proximity compare to those further apart [35]. This feature was clearly observed in our dataset (Supplementary Fig. 1B).

Direct comparisons of levels of expression across molecular (fusion positive versus negative RMS), histological (skeletal muscle versus RMS), and clinical (metastatic M1 versus non metastatic M0) entities produced lists of miRNAs significantly over or under-expressed in these groups (Supplementary Tables S2-5). Correlation between miRNA expression and overall survival (OS) and progression free survival (PPS) was also assessed by Cox proportional hazard regression model (Supplementary Tables S6-9).

**miRNA profile distinguishes RMS according to fusion gene status**

Unsupervised hierarchical clustering analysis performed using miRNAs with at least 100 tags in 50% of the samples showed the presence of distinct patterns of expression across RMS samples (Fig. 3). Notably, miRNA expression patterns mostly distinguished RMS histological subtypes showing sets of miRNAs positively and negatively associated with fusion gene status. The well-known muscle specific miRNAs including miR-1, miR-133 and miR-206 showed higher expressed in skeletal muscle versus RMS subtypes.

Application of class prediction analysis based on nearest shrunken centroid classifier [32] showed that miRNA expression can distinguish fusion positive and negative patients with around 89% accuracy (Supplementary Fig. S2) using the expression level of

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**Migration and viability assays**

At 72 h post transfection, cells were re-suspended in DMEM plus 1% FCS and placed in triplicate into cell culture inserts (BD, NJ, USA) before being submerged into specially adapted 24 well plates (BD, NJ, USA) containing DMEM plus 10% FCS. 24 h later, cells that had not migrated were removed using a cotton bud, and cells that had migrated to the base of the inserts were fixed overnight in 100% methanol at −20 °C and then stained for 2 min with 2% crystal violet solution (Sigma-Aldrich, MO, USA). Cells were photographed at 10× magnification using a Leica DM IRB microscope with a Leica 420C camera attached (Leica Microsystems, Milton Keynes, UK). Viability was assessed at 72 h post transfection using Cell Titer Aqueous ONE reagent (Promega, WI, USA) as previously described [7].
**Fig. 4. Validation of clinical correlations.**

A. Validation of NGS results on larger cohort of RMS. Box and whisker plots representing the expression levels (QRT-PCR) of miR-9 in 13 normal skeletal muscles, 66 ERMS, 31 ARMS fusion negative (ARMS, NEG), 45 ARMS PAX3-FOXO1A (ARMS, P3F), 12 ARMS PAX7-FOXO1A (ARMS, P7F), 8 ERMS cell lines (CL_ERMS), 8 ARMS cell lines (CL_ARMS) and 2 myoblast samples (Myoblasts).

B. Silencing of MYCN using siRNAs resulted in a reduction in the expression level of miR-9 in RH30 cells when compared to control cells. Data represent a triplicate measurement for each sample at 72 h post addition of negative control and MYCN siRNAs. **p < 0.01; ***p < 0.001.

E. Missiaglia et al. / Cancer Letters 385 (2017) 251–260
C. Kaplan-Meier plots for miR-9 expression and overall survival (OS) Expression levels within the seen using a larger population of 132 publicly available RMS sam-
tients [30] and identi-
WGCNA using parallel gene expression data from the 64 RMS pa-
associated with similar biological function/pathways, we employed
modules (miR-221, DP-0.24, -0.043; miR-222, DP-0.2, -0.041, respectively). Both modules had functional annotations linked to skeletal muscle development. Additionally mir-1 and -133a showed inverse correlation with green module (miR-1, DP-0.31; miR-133a, DP-0.37) that was linked to cell cycle, indicating that low levels of these miRNAs correlated with genes involved in cell cycle progression. These miRNAs/gene expression correlations associated with differentiation processes strongly support the ability of the approach to identify key roles for miRNAs in RMS.

**PAX3-FOXO1 alters the profile of miR expression**

Consistent with miRNA expression accurately distinguishing fusion positive from negative patients, four of the 12 modules correlated with fusion gene positivity (turquoise, purple, green/yellow, and pink) with turquoise correlating most strongly (Fig. 2B and indicated in Fig. 3). MiRNAs were also overexpressed in PAX3/7-FOXO1 positive cases compared to RMS without the fusion gene (Supplementary Table S3). Based on these results, we hypothesized that the fusion protein could regulate miRNA expression. To identify miRNAs altered by PAX3-FOXO1, RNA interference (RNAi) was used to reduce expression levels of the fusion gene in the cell line RH4 (Supplementary Fig. S3A), and corresponding miRNAs levels screened for in duplicate (Supplementary Table S11). To prioritize miRNAs of likely relevance in RMS patients for further investigation we identified those that were reduced by knockdown of the fusion protein, were most strongly positively correlated with the turquoise and purple modules and that were also differentially overexpressed comparing fusion gene positive with negative RMS. This identified miRNAs as candidates for further investigation (Table 2), including miR-9-5p that was additionally correlated with metastatic behavior and poor outcome (Supplementary Tables S4-59).

**Validation of miR-9-5p levels in RMS driven by PAX3-FOXO1 via MYCN**

miR-9-5p expression levels were analyzed by quantitative RT-PCR in a larger cohort of RMS patients (n = 154. Supplementary Table S1B) and, consistent with the sequencing data (Supplementary Table S3), results showed that levels of miR-9-5p were significantly higher in fusion gene positive versus negative

### Table 2

| miRNA     | Turquoise module (Pearson correlation coefficient) | Purple module (Pearson correlation coefficient) | log2 fold change expression fusion positive vs negative RMS | RH4 set 1 fold expression relative to non targeting control | RH4 set 2 fold expression relative to non targeting control |
|-----------|---------------------------------------------------|-------------------------------------------------|----------------------------------------------------------|----------------------------------------------------------|----------------------------------------------------------|
| miR-9-5p  | 0.67                                              | 0.51                                            | 1.14                                                     | -2.49                                                   | -1.15                                                   |
| miR-660-3p | 0.64                                              | 0.41                                            | 0.53                                                     | -3.54                                                   | -1.22                                                   |
| miR-9-3p  | 0.70                                              | 0.53                                            | 1.32                                                     | -1.42                                                   | -1.32                                                   |
| miR-532-3p | 0.67                                              | 0.49                                            | 0.46                                                     | -1.40                                                   | -1.37                                                   |
| miR-135a-5p | 0.57                                             | 0.45                                            | 1.45                                                     | -13.01                                                  | -8.85                                                   |
| miR-193b-5p | 0.58                                             | 0.19                                            | 0.43                                                     | -3.18                                                   | -2.14                                                   |
| miR-362-5p | 0.72                                              | 0.51                                            | 0.28                                                     | -4.98                                                   | -1.28                                                   |

11 miRNAs. These data suggest a strong interplay between fusion protein and miRNA expression that potentially contributes to the malignant phenotype of RMS.

**RMS gene co-expression-modules are assigned biological functions and correlate with clinicopathological features**

Based on the principle that co-expressed genes are commonly associated with similar biological function/pathways, we employed WGCNA using parallel gene expression data from the 64 RMS patients [30] and identified 12 groups of co-expressing genes/modules. Each module was color coded and tested for GO term enrichment in order to associate each module with biological functions (Table 1). Consistent modular gene structure was also seen using a larger population of 132 publicly available RMS samples [13] suggesting that these modules are a good representation of the biological characteristic of RMS tumors (data not shown). In addition, an eigengene per module was generated for each patient to summarize the expression of all the genes contained in each module. These values were used to correlate modules to each other as well as to clinico-pathological features (Fig. 2). Modules did not show significant correlation with IRS stage, metastasis, tumor size or age of patient at diagnosis (data not shown). However notably, the turquoise, green/yellow, pink and purple modules showed a significant correlation fusion status and histological subtype (Fig. 2B). The association between modules and overall survival (OS) or progression free survival (PFS) was also tested. The pink and turquoise modules showed a significant correlation with the OS and PFS (Supplementary Table S10).

**RMS gene-expression-modules associated with skeletal muscle development correlate with known myomiRs**

As miRNAs are known to modulate many of their targets through miRNA transcript degradation [14,36], we correlated the expression of miRNAs to the module eigengenes in order to evaluate their level of association. Fig. 3 represents a heatmap of the correlation between module eigengenes and a subset of miRNAs selected based on their coefficient values (absolute value above 0.4) and significance (p value < 0.001). The power of this approach is demonstrated by well-known myogenic miRs (myomiRs) -1, -206 and -133a that are most highly expressed in more differentiated skeletal muscle. These myomiRs correlated positively with yellow (miR-1, direct pearson (DP) 0.59; miR-206, DP 0.49; miR-133a, DP 0.58), and purple modules (miR-1, DP 0.73; miR-206, DP 0.46; miR-133a, DP 0.70) (Fig. 4). In an inverse manner, miR-221 and -222 are myoD-associated myomiRs most highly expressed early in myogenic differentiation [37] and show a negative correlations with the yellow and purple modules (miR-221, DP-0.24, -0.043; miR-222, DP-0.2, -0.041, respectively). Both modules had functional annotations linked to skeletal muscle development. Additionally mir-1 and -133a showed inverse correlation with green module (miR-1, DP-0.31; miR-133a, DP-0.37) that was linked to cell cycle, indicating that low levels of these myomiRs correlated with genes involved in cell cycle progression. These myomiRs/gene expression correlations associated with differentiation processes strongly support the ability of the approach to identify key roles for miRNAs in RMS.

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Discussion

Here we describe the sequencing analyses of miRNAs in a large series (n = 64) of well-characterized RMS samples. We were broadly able to distinguish tumor from skeletal muscle and also fusion gene positive from fusion gene negative RMS. This is consistent with a previous study of sarcomas that included only 7 RMS cases in which cluster analyses of miRNA data separated skeletal muscle from RMS and other types of sarcoma and placed 3 alveolar RMS in a different group from one embryonal case and 3 adult pleomorphic cases [40]. Furthermore, our integrated analyses of miRNA expression data with parallel gene expression levels using WGCNA identified co-expression modules that we were able to assign with biological functions and also correlate with clinical outcome data and fusion gene status. Given that miRNAs regulate the expression of many genes in a dynamic and cell context specific manner [34], these results provide a valuable framework and resource for further investigations of RMS.

Reassuringly, modules identified as associated with skeletal muscle development (Fig. 3) included correlations with miRNAs that have been previously described as miRNAs involved in myogenesis (myomiRs) and also shown to functionally impact on the phenotype of RMS cell lines. Specifically these included miRs -1, -206, -133a and miR-378 that correlated positively with muscle associated modules (yellow and purple, Table 1, Fig. 3) [19–25]. In addition, we previously showed that lower levels of miR-206 expression correlated with poorer outcome in samples from RMS patients that were linked to their undifferentiated myogenic phenotype, indicating relevance to the clinical behavior of patients [22]. MiR-221 and miR-222 are also described as myomiRs and their overexpression and reduction is reported to enhance and decrease myoD protein levels, respectively [37]. As predicted from this, our analyses of miRs-221-3p and -222-3p inversely correlated their expression with the skeletal muscle associated modules (yellow and purple) in contrast to miRs -1, -133 and -378 which were positively correlated to these functional modules. MiR-128-3p shows a similar expression pattern in these modules to miRs -1, -133 and -378 and is described with a clear role in muscle development [41]. A definitive role for miRs -221, -222 and 128-3p in RMS remains to be determined, as well as other miRNAs correlated with the modules defined with myogenic functions, such as miR-96-3p, miR-152-3p and miR-156-5p.

Interestingly, miR-1, 133, 206, 378 and 128-3p negatively, and miR-221 and -222 positively, correlated with gene expression processes for cell adhesion, chemotaxis and extracellular organization (pink module). This is consistent with the differentiation status of RMS cells shown on oncogenic KRAS (G12D) induced zebrafish models of RMS where embryonal RMS cells recapitulate normal myogenesis and their associated migratory and proliferative capacity [42]. Also, targets of these miRNAs in these networks are likely to contribute to these invasive and migratory phenotypes in RMS, for example miR-206 targets the MET tyrosine-kinase receptor which is highly expressed in RMS and enhances RMS cell migration [24]. Similarly, miR-221 and -222 have strong evidence to support that they target TIMP3 and MMP1 proteins known to influence the extracellular matrix (http://mirTarBase.mbc.nctu.edu.tw/index.php).

The myomiRs -1, -133a and miR-378 also inversely correlated with modules linked to cell cycle progression (green and red).
consistent with their role in cell cycle arrest when overexpressed in RMS [21,23] and the requirement of this prior to normal myogenic differentiation [43]. These cell cycle related modules were positively correlated with other miRNAs that may play a role in RMS, including miR-106b, that is overexpressed and associated with proliferation in medulloblastoma [44], miR-32 that when down-regulated in mesenchymal stem cells inhibits cell cycle progression [45] and miR-130b that enhances proliferation through inactivating HIPPO signaling in glioma [46]. Mir-19a-3p, as part of the polycistronic miRNA cluster miR-17-92, also positively correlated with these cell cycle modules as well as a module associated with fusion genes (turquoise). The latter and red modules were both linked to Sonic hedgehog (SHH) signaling. Together this is consistent with high expression of the miR17-92 cluster being genomically amplified in around 20% of RMS and MYCN driven, mostly in those with PAX7-FOXO1 fusion genes, and possible involvement of the cluster in proliferation through SHH signaling [47,48].

The strongest positive and negative correlations between genes and miRNA expression levels were associated with fusion gene positive RMS cases (Figs. 2 and 3). In addition, the turquoise and pink modules linked to fusion gene positivity were also most significantly associated with poor outcome (Supplementary Table S10). This is consistent with several studies, including our own, demonstrating the prognostic value of the presence of a fusion gene in RMS [9–12]. Our analyses indicate that the fusion protein impacts on the expression of miRNAs and through screening changes to miRNA levels following reduction of PAX3-FOXO1 expression in a cell line, we were able to show that the fusion protein alters the expression levels of microRNAs. Interestingly, miR-9-5p was one of the top hits in the screen, was highly correlated with the modules associated with the presence of the fusion gene and it was more highly expressed in fusion gene positive RMS compared to fusion gene negative RMS (Table 2). Although no PAX3-FOXO1 binding sites were identified near miR-9-5p, we showed miR-9-5p to be indirectly driven by PAX3-FOXO1 via MYCN. MYC/MYCN have previously been shown to bind to the miR-9 locus and activate expression in other cell types [39]. MYCN is known to be a direct transcriptional target of the fusion protein [5] and to operate in a positive feedback loop mechanism [7]. This overlap in miRNA targets for PAX3-FOXO1 and MYCN is to the gene expression changes identified through altering MYCN levels [7] and PAX3-FOXO1 expression in RMS that show considerable overlap [13]. Taken together, this indicates that the fusion protein together with MYCN, and other transcription factors downstream of PAX3-FOXO1, such as MYOD1, regulate the expression of miRNAs and their many target genes, to massively re-programme gene expression in RMS cells towards a tumorigenic phenotype [49]. The contribution of these altered levels of miRNAs is predicted to impact on the phenotype of fusion positive RMS in ways that include the developmental, myogenic, immunological and cell adhesion processes indicated in the co-expression modules (Fig. 3).

As representative of a miRNA associated with the presence of PAX3-FOXO1, miR-9-5p levels are shown to contribute to the clinical and biological behavior of RMS. MiR-9-5p levels were higher in metastatic versus non-metastatic cases and correlated with a poor outcome of patients, including within fusion gene positive group. Through reducing miR-9-5p we deduce a role for miR-9-5p in enhancing RMS cell migration and this cellular phenotype potently facilitates metastatic behavior. This is similar to the roles for miR-9-5p in the proliferation, differentiation and migration of neural progenitor cells but miR-9-5p may alter different gene targets dependent on cell context [50]. Also, aberrant miR-9-5p levels have been indicated in several cancer types where it may be either overexpressed or underexpressed. Overexpression has been described in Hodgkin’s lymphoma [51], brain tumors [52], neuroblastoma [39] and gastric cancers, in the latter affecting proliferation via the miR-9 target CDX2 [53]. Reduced levels of miR-9-5p play a role in ovarian cancer cells where this, in part, enhances cell proliferation through NFκappaB1 [54]. Furthermore, in normal epithelial and breast cancer cells, miR-9-5p induces epithelial to mesenchymal transition by targeting E cadherin (CDH1) [39]. Recently, reducing miR-9-5p in an osteosarcoma cell line and identification of high levels of expression in primary tumors suggests a role for this miRNA in regulating the progression of osteosarcoma [55]. This role in osteosarcomas in addition to RMS described here, indicates a role for miR-9-5p in mesenchymal malignancies further to that previously described epithelial and neural cancers.

MiRNAs altered by the fusion protein, including indirectly miR-9-5p (Table 2, Supplementary Tables S3 and S11), are likely to play an important role in RMS. Furthermore, the expression levels and the functional framework presented have assigned likely functions for miRNAs in RMS for further investigation. These could identify additional tumor biomarkers and, depending on their expression in bodily fluids, useful markers for disease monitoring, as exemplified by the detection of the myomiR miR-206 in serum from RMS patients [56]. Achieving good in vivo delivery for therapeutically targeting non-inducing miRNAs that are not detectable currently is still challenging [29]. However, further understanding of miRNAs in RMS may ultimately enable targeting their key roles via regulation of multiple genes and pathways in ways that could improve outcome for patients.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.canlet.2016.10.011.

Conflict of interest

None.

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