ARTICLE

PAX3-FOXO1 drives miR-486-5p and represses miR-221 contributing to pathogenesis of alveolar rhabdomyosarcoma

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Abstract
Rhabdomyosarcoma is the most common soft-tissue sarcoma in childhood and histologically resembles developing skeletal muscle. Alveolar rhabdomyosarcoma (ARMS) is an aggressive subtype with a higher rate of metastasis and poorer prognosis. The majority of ARMS tumors (80%) harbor a PAX3-FOXO1 or less commonly a PAX7-FOXO1 fusion gene. The presence of either the PAX3-FOXO1 or PAX7-FOXO1 fusion gene foretells a poorer prognosis resulting in clinical re-classification as either fusion-positive (FP-RMS) or fusion-negative RMS (FN-RMS). The PAX3/7-FOXO1 fusion genes result in the production of a rogue transcription factors that drive FP-RMS pathogenesis and block myogenic differentiation. Despite knowing the molecular driver of FP-RMS, targeted therapies have yet to make an impact for patients, highlighting the need for a greater understanding of the molecular consequences of PAX3-FOXO1 and its target genes including microRNAs. Here we show FP-RMS patient-derived xenografts and cell lines display a distinct microRNA expression pattern. We utilized both loss- and gain-of function approaches in human cell lines with knockdown of PAX3-FOXO1 in FP-RMS cell lines and expression of PAX3-FOXO1 in human myoblasts and identified microRNAs both positively and negatively regulated by the PAX3-FOXO1 fusion protein. We demonstrate PAX3-FOXO1 represses miR-221/222 that functions as a tumor suppressing microRNA through the negative regulation of CCND2, CDK6, and ERBB3. In contrast, miR-486-5p is transcriptionally activated by PAX3-FOXO1 and promotes FP-RMS proliferation, invasion, and clonogenic growth. Inhibition of miR-486-5p in FP-RMS xenografts decreased tumor growth, illustrating a proof of principle for future therapeutic intervention. Therefore, PAX3-FOXO1 regulates key microRNAs that may represent novel therapeutic vulnerabilities in FP-RMS.

Introduction
Rhabdomyosarcoma (RMS), the most common soft-tissue sarcoma in children [1], is divided into two major histologic subclasses: embryonal RMS (ERMS) and alveolar RMS (ARMS) [2]. ARMS is notoriously more aggressive and most (~80%) ARMS tumors harbor chromosomal translocations t(2;13)(q35;q14) or t(1;13)(p36;q14) resulting in PAX3-FOXO1 or PAX7-FOXO1 gene fusions respectively [3–5]. The presence of the PAX3/7-FOXO1 fusion gene foretells a worse prognosis and is superior to histology in predicting survival [6–9]. ARMS patients without a PAX3/7-FOXO1 translocation have both molecular features and clinical outcome similar to ERMS [7, 10]. Therefore, molecular classification as fusion-positive RMS (FP-RMS) and fusion-negative RMS (FN-RMS) based on the presence or absence of the PAX3/7-FOXO1 fusion more accurately represents the clinical features and biology of RMS. Despite the recently expanded genomic understanding of RMS, patient survival and the treatment strategies have not improved [11–14].

In both PAX3-FOXO1 and PAX7-FOXO1 fusion proteins, the amino terminus of the paired box (PAX) protein including the DNA-binding domain is fused to the carboxy

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Fig. 1 Differential microRNA expression in FN-RMS and FP-RMS. Heat map of differentially expressed microRNAs based on $P < 0.05$ and fold change $> 2$ in FP-RMS cell lines (green, Rh3, Rh4, Rh28, Rh30, Rh41), FP-RMS patient-derived xenografts (PDX, yellow, SJRH5010468_X1, SJRH5010463_X16), FN-RMS cell lines (blue, RD, Rh2, Rh18), and FN-RMS PDX (red, SJRH5011_X, SJRH5011_Y, SJRH5012_X, SJRH5012_Y, SJRH5012_Z, SJRH5013_X, SJRH5026_X).

The terminus of the forkhead box O1 (FOXO1) protein including the transcriptional transactivation domain. Thus, it is thought that the PAX3/7-FOXO1 fusion proteins function as aberrant transcription factors at PAX3 or PAX7 target genes [15]. FP-RMS cells are addicted to PAX3-FOXO1 expression making it therapeutically attractive [16, 17].
Given targeting transcription factors remains a challenge, many groups have focused on identifying and inhibiting PAX3-FOXO1 target genes [18]. However, little is known about the regulation of non-coding RNAs including microRNAs by PAX3-FOXO1.

MicroRNAs are single-stranded ~22 nucleotide non-coding RNAs that function to reduce gene expression [19]. Primarily through binding complementary sequences in the 3′ untranslated regions (UTR) of target mRNAs, microRNAs target transcripts for degradation or repress transcript translation [20]. Individual microRNAs can modulate the
expression of hundreds of target genes allowing for regulation of complex biological processes. MicroRNAs have been implicated in normal development, differentiation, cellular homeostasis, and diseases including cancer [21–24]. Methods to block or replace microRNA function with either inhibitors or mimics are now in clinical trials opening the potential to leverage microRNA biology to target multiple nodes in signaling pathways or biological processes [22, 25].

Given the PAX3-FOXO1 fusion protein is thought to function as a rogue transcription factor; transcriptional regulation by PAX3-FOXO1 is thought to be central to FP-RMS pathogenesis. Nothing is known about how the regulation by PAX3-FOXO1 is thought to be central to FP-RMS pathogenesis. We hypothesize that microRNAs are transcriptionally regulated by the PAX3-FOXO1 fusion protein and actively participate in FP-RMS pathogenesis.

Results

Distinct microRNA expression in FP-RMS and FN-RMS

We hypothesized that if microRNAs participate in driving the clinical differences between FN- and FP-RMS microRNA expression patterns should be distinct. To interrogate microRNA expression in RMS, we assessed the microRNA expression profile in a panel of RMS cell lines as well as patient-derived xenografts (PDX) from the Childhood Solid Tumor Network [26]. Our panel included seven FN-RMS PDX samples, three FN-RMS cell lines, two FP-RMS PDX samples, and five FP-RMS cell lines. All of the FP-RMS cell lines and the SJRHBO10463_X16 PDX harbor the PAX3-FOXO1 translocation; however, the SJRHBO10468_X1 PDX contains the PAX7-FOXO1 translocation. Comparing the microRNA expression between FP-RMS and FN-RMS with greater than two-fold difference and P-value less than 0.05, we identified 57 microRNAs decreased and 52 with increased expression in FP-RMS compared to FN-RMS (Fig. 1). Because the PAX3-FOXO1 translocation is the driver of FP-RMS, we hypothesized that microRNAs differentially expressed in FP-RMS are transcriptionally regulated by the PAX3-FOXO1 fusion protein.

Knockdown of PAX3-FOXO1 reduces transformative properties and promotes differentiation of FP-RMS

In order to identify microRNAs regulated by PAX3-FOXO1 in FP-RMS, we generated a doxycycline-inducible lentivirus expressing shRNA directed against PAX3-FOXO1 breakpoint or a scrambled shRNA control. The PAX3-FOXO1 shRNA (shP3F) targets the translocation breakpoint to minimally complement endogenous PAX3 or FOXO1 sequences (Fig. 2a). We utilized the broadly studied FP-RMS Rh30 and Rh41 cell lines harboring the PAX3-FOXO1 translocation to established stable cell lines that express doxycycline-inducible shP3F or scramble control (shScr) shRNA [27, 28]. Treatment with doxycycline in both Rh30 and Rh41 cells leads to a significant knockdown of PAX3-FOXO1 protein (Fig. 2b and Supplementary Figure S1a) and transcript levels (Fig. 2c and Supplementary Figure S1c). Reduced PAX3-FOXO1 protein correlated with a reduction of well-known PAX3-FOXO1 target genes ALK, FGFR4, and MET (Fig. 2c and Supplementary Figure S1c). FOXO1 and PAX3 proteins are not decreased with the shP3F illustrating specificity (Fig. 2b and Supplementary Figure S1a and b).

To confirm functional PAX3-FOXO1 knockdown in our system, we measured the phenotypic consequences in Rh30 and Rh41 cells. The PAX3-FOXO1 knockdown in Rh41 cells markedly decreased the population doubling of cells to the point that cells had senesced or died after 12 days in doxycycline (Fig. 2d). In contrast, PAX3-FOXO1 knockdown in Rh30 significantly reduced population doubling but permitted continued although attenuated growth (Supplementary Figure S1d). PAX3-FOXO1 knockdown reduced cell viability and increased apoptosis (Fig. 2e, f, Supplementary Figure S1e and f). Cell cycle analysis with FACS illustrated PAX3-FOXO1 knockdown results in a G1 arrest (Fig. 2g). PAX3-FOXO1 knockdown in both Rh41 and Rh30 reduced colony formation in clonogenic cell survival assays (Fig. 2h and Supplementary Figure S1g). Since FP-RMS has a higher propensity for metastasis, we assessed the effect of PAX3-FOXO1 knockdown on cell migration and invasion both of which were reduced with PAX3-FOXO1 knockdown (Fig. 2i, j). We noted that with PAX3-FOXO1 knockdown both Rh30 and Rh41 cells elongated and formed myosin heavy chain-positive multinucleated myotubes and expressed mature skeletal muscle gene CKM indicative of myogenic differentiation (Supplementary Figure S2a–f). These results illustrate that our inducible shP3F system reduces expression of known PAX3-FOXO1 target genes, reduces the oncogenic capacity and invasion potential while inducing myogenic differentiation of FP-RMS cell lines.

Identification of candidate microRNAs regulated by PAX3-FOXO1

To identify microRNAs regulated by PAX3-FOXO1 in FP-RMS, we utilized both loss- and gain-of-function approaches. First, shP3F and shScr Rh41 knockdown and control cells respectively were treated with doxycycline for 5 days (before population doubling decreases) and RNA was
prepared for microRNA expression profiling (Fig. 3a). To complement the loss-of-function PAX3-FOXO1 knockdown, PAX3-FOXO1 was stably expressed in the FN-RMS RD cell line with correspondingly increased expression of PAX3-FOXO1 target genes (Supplementary Figure S3a and b). We hypothesized that microRNAs transcriptionally activated by PAX3-FOXO1 would have decreased expression in the shP3F knockdown Rh41 cells and increased expression in the PAX3-FOXO1 forced expression RD cells. In contrast, microRNAs transcriptionally repressed by PAX3-FOXO1 would have increased expression in the shP3F Rh41 cells and decreased expression in the RD cells with PAX3-FOXO1 forced expression.

MicroRNA microarray analysis identified a total of 44 microRNAs with greater than two-fold expression difference with PAX3-FOXO1 knockdown (Fig. 3a). Seventeen microRNAs had decreased expression and 27 had increased expression with PAX3-FOXO1 knockdown. To further define microRNAs specifically regulated by the PAX3-FOXO1 fusion protein, we compared the microRNA expression in the RD cells expressing PAX3-FOXO1. Of the 17 microRNAs with decreased expression with shP3F, 8 microRNAs (miR-18a-5p, miR-301b, miR-326, miR-335-3p, miR-455-3p, miR-486-5p, miR-486-3p, and miR-5100) increased expression in RD cells with PAX3-FOXO1 overexpression. We hypothesized that these microRNAs are transcriptional target genes of PAX3-FOXO1. In
Fig. 4 miR-221 functions as a tumor suppressor in FP-RMS and targets several known oncogenes. a Expression of miR-221 by qRT-PCR in patient-derived xenografts from the Childhood Solid Tumor Network (CSTN), including FN-RMS (n = 13, SJRH010463_X16, SJRHB013759_X1, SJRH010468_X1 and SJRHB013757_X2). b Expression of predicted miR-221 targets by qRT-PCR in Rh30 and Rh41 cells transfected with NC or miR-221 mimic. c Luciferase activity in C2C12 cells co-transfected with miR-221 or control vector and wild type or mutant miR-221 site 3’UTRs. Luciferase activity represented as mean S.E.M. (n = 4), the Renilla/Firefly luciferase ratio normalized to empty reporter (no miR-221). d Expression of miR-221 targets CCND2, CDK6, and ERBB3 by immunoblot analysis in cells as in (g). *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001 for FN-RMS vs FP-RMS or miR-221 mimic vs NC.
contrast, 23 of the 28 microRNAs with increased expression with PAX3-FOXO1 knockdown had decreased expression in RD cells with PAX3-FOXO1 expression, suggesting they are repressed by PAX3-FOXO1.

We validated six of the eight microRNAs with decreased expression with PAX3-FOXO1 knockdown in FP-RMS cells and concordant increased expression with forced expression of PAX3-FOXO1 in the FN-RMS, RD cell line. We excluded miR-5100 as it is poorly conserved. Both miR-486-5p and miR-486-3p had decreased expression with PAX3-FOXO1 knockdown on the microarray; however, we restricted our experiments to miR-486-5p as it is upregulated by PAX3-FOXO1 in LHCN-M2 cells (Fig. 3c). Regulated microRNAs except miR-301b and miR-335 were upregulated (Supplementary Figure 3a and b). All of the RH41 cells; however, PAX3-FOXO1 target genes were expressed as a cluster on the X chromosome and share an identical seed sequence, thus share the same target mRNAs. Since they are transcriptionally co-regulated and target the same mRNAs, we focused on miR-221 and validated it to be inversely correlated with PAX3-FOXO1 expression by qRT-PCR (Fig. 3d, e, Supplementary Figure S3e). Thus we identified CCND2, CDK6, and ERBB3 as potential miR-221 targets for further analysis. MiR-221 overexpression in Rh30 and Rh41 cells decreased the expression of all the selected targets except KIT by qRT-PCR, protein levels on immunoblot, and we confirmed direct miR-221 regulation of CCND2, CDK6, and ERBB3 with 3′ UTR luciferase reporter assays in C2C12 cells (Fig. 4g–i). Importantly, luciferase activity was rescued following mutation of the miR-221-binding site, although the ERBB3 rescue was not complete suggesting additional non-predicted sites in the 3′ UTR (Supplementary Figure S4). Collectively these data suggest that miR-221 is negatively regulated by PAX3-FOXO1 and functions as a tumor suppressor in FP-RMS partially through targeting CCND2, CDK6, and ERBB3 decreasing proliferation and migration/invasion.

miR-221 decreases proliferation and invasion by regulating CCND2, CDK6, and ERBB3

To further explore the role of miR-221 in FP-RMS, we first validated its expression level in an expanded cohort of PDX samples consisting of four FP-RMS PDX samples with two PAX3-FOXO1 fusions (SJRHB010463_X16 and SJRHB013759_X1) and two PAX7-FOXO1 fusions (SJRHB010468_X1 and SJRHB013757_X2) as well as 13 FN-RMS from the Childhood Solid Tumor Network [26]. MiR-221 is significantly reduced in FP-RMS compared to FN-RMS (Fig. 4a) consistent with a recent study in 64 RMS patients [31]. Transient transfection of miR-221 mimic led to overexpression as evaluated by qRT-PCR, reduction of cell viability, and an increase in apoptosis in both Rh30 and Rh41 cells (Fig. 4b–d). We next determined if miR-221 overexpression could contribute to FP-RMS pathogenesis beyond proliferation and found a striking reduction in the migratory capacity and invasive potential of the cells (Fig. 4e, f). Thus, miR-221 expression decreased proliferative, survival and metastatic potential of FP-RMS.

Given that overexpression of miR-221 decreased proliferation and the invasiveness of FP-RMS cells, we sought to identify the downstream targets of miR-221 responsible for the observed phenotypes. We interrogated the microRNA target prediction algorithms TargetScan [32] as well as previously published miR-221 targets with a focus on known oncogenes in RMS [33–36]. Thus we identified CCND2, CDK6, KIT, and ERBB3 as potential miR-221 targets for further analysis. MiR-221 overexpression in Rh30 and Rh41 cells decreased the expression of all the selected targets except KIT by qRT-PCR, protein levels on immunoblot, and we confirmed direct miR-221 regulation of CCND2, CDK6, and ERBB3 with 3′ UTR luciferase reporter assays in C2C12 cells (Fig. 4g–i). Importantly, luciferase activity was rescued following mutation of the miR-221-binding site, although the ERBB3 rescue was not complete suggesting additional non-predicted sites in the 3′ UTR (Supplementary Figure S4). Collectively these data suggest that miR-221 is negatively regulated by PAX3-FOXO1 and functions as a tumor suppressor in FP-RMS partially through targeting CCND2, CDK6, and ERBB3 decreasing proliferation and migration/invasion.

Candidate oncogenic microRNAs contribute to FP-RMS pathogenesis

We hypothesized that microRNAs positively regulated by PAX3-FOXO1 could contribute to FP-RMS pathogenesis functioning as oncogenes. To this end, we assessed cell viability and apoptosis after transfection with antimiRs for microRNAs positively regulated by PAX3-FOXO1 and upregulated in FP-RMS cell lines and patient-derived
Fig. 5 Contribution of candidate oncogenic microRNAs to FP-RMS. 

**a** qRT-PCR microRNA expression in Rh41 cells 3 days after transfection with indicated negative control (NC) or antimiR. 

**b** Cell Titer Glo cell viability assay in Rh41 cells as in (**a**). 

c qRT-PCR in Rh30 cells transfected with antimiR-486-5p or NC 3 days after transfection. 

d Cell Titer Glo cell viability and e Caspase 3/7 Glo apoptosis assay in Rh30 cells as in (**c**). *P < 0.05, **P < 0.01, ***P < 0.001 for antimiRs vs NC.
xenografts (Supplementary Table S1). We included miR-506 and miR-510 given they had the largest expression change with PAX3-FOXO1 knockdown although there was no expression change with PAX3-FOXO1 expression in RD cells (Fig. 3a). We also included miR-9 given it is uniformly increased in FP-RMS cell lines and PDXs although not regulated by PAX3-FOXO1 knockdown (Fig. 1). We excluded miR-455 since it is enriched in FN-RMS compared to FP-RMS (Fig. 1). AntimiRs transiently transfected in the Rh41 cells resulted in a significant decrease in microRNA level with the exception of antimiR-9 and antimiR-335 (Fig. 5a). Since antimiRs could function through sequestering the mature microRNA without degradation, we continued with phenotypic analysis despite the lack of decreased miR-9 and miR-335 expression. However, only miR-486-5p and miR-326 inhibition reduced cell viability (Fig. 5b) while none of the antimiRs increased apoptosis in Rh41 cells (Supplementary Figure S5). Because miR-486-5p had the largest effect, we validated the findings in Rh30 cells, revealing similar knockdown of miR-486-5p, significant reduced cell viability, and increased apoptosis (Fig. 5c–e).

**miR-486-5p is a direct transcriptional target of PAX3-FOXO1**

We focused on miR-486-5p as it is a muscle-enriched microRNA and antimiR-486-5p treatment in Rh41 cells had a strong antiproliferative phenotype. miR-486-5p is transcribed from intron 41 of ANK1 (Fig. 6a). ANK1 is an erythroid-specific ankyrin-repeat protein linking the plasma membrane and the cytoskeleton. A muscle-specific promoter upstream of an alternative exon 39a drives the expression of a muscle-specific ANKI protein called small ANK1 (sANK1) that links the sarcoplasmic reticulum to the sarcomere [37]. The promoter region preceding sANK1...
contains multiple E-Box binding sites for the myogenic transcription factor MYOD1 [38]. Comparable to miR-486-5p, sANK1 expression increases with PAX3-FOXO1 expression in LHCN-M2 and RD cells and decreases with PAX3-FOXO1 knockdown in FP-RMS cells (Fig. 6b, c). To determine if PAX3-FOXO1 transcriptionally regulates sANK1 and miR-486 we cloned the 3 kb promoter region 5' of ANK1 exon 39a containing the MYOD1 responsive E-boxes upstream of luciferase for reporter assays. Increasing PAX3-FOXO1 overexpression led to a dose-dependent increase in sANK1 promoter-driven luciferase activity comparable to luciferase activity with MyoD (Fig. 6d). Thus, both sANK1 and miR-486-5p are transcriptionally regulated by PAX3-FOXO1 through the sANK1 promoter.

miR-486-5p promotes invasion of FP-RMS cell lines

We confirmed miR-486-5p overexpression in FP-RMS compared to FN-RMS in patient-derived xenografts (Fig. 7a). Because miR-486-5p expression is positively regulated by PAX3-FOXO1 and upregulated in FP-RMS PDXs, we hypothesized that miR-486-5p could contribute to phenotypes of FP-RMS cells other than promoting proliferation. Similar to PAX3-FOXO1 knockdown, Rh41 cells with antimiR-486-5p transfection reduced both cell migration and invasion in transwell assays (Fig. 7b, c). In addition, miR-486-5p inhibition reduced clonogenic colony formation capacity of Rh41 cells (Fig. 7d). Thus, inhibition of miR-486-5p phenocopies PAX3-FOXO1 knockdown illustrating that miR-486-5p contributes to cell proliferation, cell migration and invasion, and clonogenic growth in FP-RMS.

To elucidate the molecular mechanisms of miR-486-5p in FP-RMS, we sought to explore the miR-486-5p target genes in FP-RMS. First, we identified 2755 putative target genes of miR-486-5p using the target prediction algorithm TargetScan irrespective of target site conservation. To narrow the list of putative target genes, we compared the putative target genes from the TargetScan algorithm to that of genes with increased expression with PAX3-FOXO1 knockdown (GSE73483) [39]. We identified 35 miR-486-5p putative target genes with increased expression with PAX3-FOXO1 knockdown (Supplementary Figure S6a). Of the 35 putative miR-486-5p target genes, 5 have tumor suppressive roles in cancer, including ARID1A, CYLD, GRIN20, PNPLA4, and SOCS2. We interrogated miR-486-5p regulation by transfecting Rh41 cells with NC or antimiR-486-5p and performing qRT-PCR (Supplementary Figure S6b). However, none of these five genes were upregulated with miR-486-5p inhibition. We also assessed previously identified miR-486-5p targets, including ARID4B, EMP, FOXO1, PIK3AP1, PTEN, SMARCD2, TOB1, and TWF1. Interestingly, only subtle but significant changes in gene expression were detected for several predicted target genes, including ARID4B, PIK3AP1, SMARCD2, and TWF1 (Supplementary Figure S6b). Despite the known role of miR-486-5p in regulating the...
PI3-kinase/AKT pathways in skeletal muscle and lung cancer, miR-486-5p inhibition did not result in increased expression of previously identified target genes PTEN, FOXO1, or PIK3R1 or activation of the PI3K/AKT pathway (Supplementary Figure S6b and c) [38, 40, 41]. One of the top predicted targets of miR-486-5p is Twinfilin1 (TWF1). TWF1 is an actin monomer binding protein that contributes to motility of cells through actin cytoskeleton rearrangement. Inhibition of miR-486-5p also led to an increase in TWF1 protein levels in both Rh30 and Rh41 cells (Supplementary Figure S6d). We confirmed that miR-486-5p directly regulates TWF1 using a luciferase reporter with the 3′UTR of TWF1 (Supplementary Figure S6e and f).

**miR-486-5p inhibition reduces tumor growth in mouse xenografts**

Given that miR-486-5p inhibition in FP-RMS cell lines reduces proliferation, we sought to determine the effect of long-term miR-486-5p inhibition in vitro as well as in vivo tumor growth. We inhibited miR-486-5p with transduction of lentivirus expressing miRZip-486-5p (lenti-miRZip-486-5p) that stably express single-stranded anti-microRNAs that
bind the endogenous microRNA target, thus inhibiting its function. Transduction of Rh30 and Rh41 with lentimiRZip-486-5p significantly decreased miR-486-5p expression and cell viability (Fig. 8a, b). Transduction efficiency was evaluated by GFP fluorescence expressed from the lenti-miRZip and greater than 80% of the cells were GFP positive 48 h after transduction (Supplementary Figure S7a). Transduction of additional FP-RM cell lines, including Rh3, Rh4, and Rh28 led to similar transduction efficiencies and decreases in miR-486-5p expression and cell viability (Supplementary Figure S7a–c).

Rh30 and Rh41 cells were transduced with lenti-miRZip-486 or lenti-miRZip-Scr control and subcutaneously injected into immunocompromised SCID-Beige mice 48 h after transduction. A striking decrease in xenograft growth was observed in both Rh30 and Rh41 xenografts transduced with miRZip-486-5p (Fig. 8c, d, and Supplementary Figure S8a and b). In fact, measurable tumors developed in only 4 of 10 mice in Rh41 and 6 of 10 mice in Rh30 cells with miRZip-486-5p, compared to 9 of 10 mice with miRZip-Scr transduced cells. The expression of miR-486-5p was assessed in the resulting xenografts by qRT-PCR. No significant decreased miR-486-5p expression was observed suggesting a negative selection for untransduced cells or for cells with low expression of the miRZip-486-5p (Fig. 8e and Supplementary Figure S8c). Indeed, whole-mount imaging of the xenografts illustrates that miRZip-486-5p-transduced xenografts expressed significantly less GFP (Fig. 8f and Supplementary Figure S8d). The histology and immunohistochemistry (IHC) staining of the miRZip-Scr and miRZip-486 were indistinguishable in regard to MYOGENIN and Ki67 staining (Fig. 8g and Supplementary Figure S8e). However, the GFP staining on tumor sections was consistent with the whole-mount imaging with remarkably less GFP staining in the miRZip-486-5p-transduced xenografts. Taken together, miR-486-5p promotes FP-RMS tumorigenesis with a strong selection to maintain high expression of miR-486-5p.

**Discussion**

In this study, we found distinct expression of microRNAs in FP-RMS and FN-RMS and interrogated the role of microRNAs regulated by PAX3-FOXO1 in FP-RMS. Using microRNA microarrays, we found many microRNAs positively and negatively correlated with PAX3-FOXO1 in patient-derived xenografts and cell lines as well as gain- and loss-of-function RMS cells. We found miR-221/222 and miR-486-5p as novel candidates of tumor suppressive and oncogenic microRNAs, respectively, in FP-RMS.

We found miR-221/222 among the most downregulated microRNAs in FP-RMS. These microRNAs, clustered on the X chromosome, are expressed as a single pri-microRNA and contain identical seed sequences [42]. One of the first studies on miR-221/222 found that they function as tumor suppressors through the modulation of the KIT receptor in erythroleukemia cells and acute myeloid leukemia [42, 43]. In addition, miR-221 is downregulated in cholangiocarcinoma [44], gastrointestinal stromal tumors [45, 46], and pancreatic cancer where it targets IRF2 and SOCS3 [47, 48]. In endothelial cells, miR-221/222 inhibit proliferation, migration, and angiogenesis [49]. Consistent with our findings, miR-221/222 are downregulated specifically in FP-RMS cell lines and patients [31, 50]. Interestingly miR-221/222 expression correlates with overall survival and progression-free survival of RMS [31]. In contrast, miR-221/222 are overexpressed and function as oncogenes in breast cancer, hepatocellular carcinoma, glioblastoma, and others [51, 52]. Furthermore, miR-221/222 are highly expressed in myoblasts promoting proliferation and delaying differentiation [30, 53]. The function of miR-221/222 is highly context dependent on the expression and regulation of its targets [54]. The context-dependent oncogenic roles of miR-221/222 abrogate the temptation to proceed with miR-221/222 replacement therapy in RMS.

In FP-RMS, we identified CCND2, CDK6, and ERBB3 as novel targets of miR-221. Interestingly, CCND2 is the predominant D-Type Cyclin expressed in RMS and its downregulation by miR-1/206 decreases RMS proliferation [34, 55, 56]. Furthermore, Cyclin D and CDK4/6 activity is tightly regulated in myoblast proliferation and differentiation [57, 58] and overexpressed/hyperactive in cancer and RMS [33]. ERBB3 is significantly upregulated in FP-RMS, promotes RMS, and may represent a therapeutic target [35, 59]. Interestingly, ERBB3 upregulation may contribute to MEK inhibition resistance in RMS [60]. However, the context-dependent function of miR-221/222 suggests that therapeutic manipulation with replacement or inhibition of miR-221 needs to be rigorously evaluated since it can function as both an oncogene and tumor suppressor.

In evaluating the phenotypic effect of inhibition of 13 microRNAs positively regulated by PAX3-FOXO1, we found miR-486-5p to have the strongest proliferative phenotype. miR-486-5p is overexpressed and can function as an oncogene in KRAS-mutated colorectal cancer [61], pancreatic cancer [62], gliomas [63], Sezary syndrome lymphoma [64], and leukemias [29, 65, 66]. In contrast, miR-486-5p is downregulated in gastric cancer [67], lung cancer [40], and hepatocellular carcinoma [68]. The tumor suppressing phenotype is primarily through downregulation of IGF signaling pathway components. Similar to the promiscuous roles of miR-221 in cancer, the varied functions
of miR-486-5p in different tumors highlight the context dependence of microRNA function likely secondary to differential target gene regulation in different cell types.

miR-486-5p is a muscle-enriched microRNA that is upregulated by MRTFs and can promote myogenic differentiation [38, 69]. miR-486-5p also induces proliferation and increases myotube size in primary mouse myoblasts [70]. Furthermore, miR-486-5p induces skeletal muscle hypertrophy in Myostatin knockout mice and ameliorates muscular dystrophy through Pten regulation of Akt signaling [41, 71]. In addition, miR-486-5p can reduce skeletal muscle atrophy in chronic kidney disease [72]. Consistent with increased proliferation in myoblasts overexpressing miR-486-5p [70], we find that inhibition of miR-486-5p reduces proliferation and xenograft growth of FP-RMS. In addition, we find that miR-486-5p inhibition decreases migration and invasion of FP-RMS, indicating that miR-486-5p contributes to the increased metastatic potential of FP-RMS. We were unable to identify a single specific miR-486-5p target gene in our model system responsible for the phenotypes observed with miR-486-5p inhibition. This is consistent with idea that the “1 microRNA-1 target” model is insufficient to explain the complex phenotypes regulated by microRNAs. Modest regulation of many target genes collectively are likely responsible for the complex phenotypes induced by microRNAs. The subtle gene expression changes we report here subsequent to inhibition of predicted miR-486-5p target genes are consistent with this model.

Despite knowing the molecular driver of FP-RMS, targeted therapies have yet to make an impact for patients, highlighting the need for a greater understanding of the molecular consequences of PAX3-FOXO1 expression. Our data suggest that PAX3-FOXO1 regulates the expression of key microRNAs involved in FP-RMS pathogenesis. The hypothesis that microRNAs can target hundreds of mRNAs and therefore be more therapeutically valuable than agents against a single molecule or pathway suggest that identifying crucial microRNAs may lead to therapeutics to improve patient outcomes. miR-221/222 and miR-486-5p may represent novel therapeutic vulnerabilities in FP-RMS with inhibition of miR-486-5p and replacement of miR-221/222.

Materials and methods

Cell lines

Cells were from the following sources: RD (CCL-136) and Rh30 (CRL-2061) from ATCC (Manassas, VA, USA), Rh3, RH4, Rh28, and Rh41 (Gerard Groverseld, St. Jude), 293T (Martine F. Roussel, St. Jude), and LHCN-M2 (Woodring Wright, University of Texas Southwestern Medical Center) [73]. Cells were authenticated with short tandem repeat profiling (Supplementary Table S2) [56].

RNA and gene expression

Total RNA was isolated, reverse transcribed, and quantitative real-time PCR (qRT-PCR) performed as previously described using primers and probes in Supplementary Table S3 [56]. For microRNA microarrays, total RNAs were labeled and hybridized to The Human microRNA Microarray Release 19.0, 8 × 60k (G4872A-046064; Agilent, Santa Clara, CA, USA). Normalized signal data were log2 transformed and each probeset was compared by unequal variance t-test (Partek Genomics Suite 6.6, St. Louis, MO, USA). The microarray data have been deposited to GEO (GSE97553). Differential microRNA expression was considered significant if P < 0.05 and the fold change was greater than 2. MicroRNA targets predicted based on TargetScan (Release 7.1) [32].

Molecular cloning, viral transduction, and luciferase assays

Doxycycline-inducible shRNAs (shP3F and shScr) were generated by ligating annealed overlapping oligonucleotides for PAX3-FOXO1 or Scrambled to AgeI and EcoRI digested Tet-pLKO-Puro (gift from Dmitri Wiederschain, Addgene plasmid #21915) [74]. Primers used in cloning are described in Supplementary Table S4. pBabe-PAX3-FOXO1-puro was generated by PCR amplifying PAX3-FOXO1 from pcDNA-PAX3-FOXO1 (gift from Rene Galindo) digesting with BamHI and SalI (New England Biolabs (NEB), Ipswich, MA, USA) and ligating to pBabe-puro (gift from Jay Morgenstern and Hartmut Land Addgene plasmid #1764) [75]. Retrovirus was packaged as described previously [24]. Cells were selected and maintained in 0.5 μg/mL puromycin. shRNA expression was induced with 50 ng/mL doxycycline. miRZip-486-5p (MZIPO486-5p-PA-1; System Bioscience, Palo Alto, CA, USA) and miRZip-Scr (MZIPO000-PA-1) were packaged following manufacturers’ instructions. Transduction efficiency of miRZip was determined by counting the number of GFP-positive cells divided by the total number of cells in four random fields for each condition.

458 bp of the human genomic DNA sequence containing miR-221 was synthesized (gene block; Integrated DNA Technologies (IDT), Coralville, IA, USA), digested with EcoRI and SalI and ligated to pCMV6. 337 bp of the human genomic DNA sequence containing miR-486 was PCR amplified using primers detailed in Supplementary Table S4, digested with EcoRI and SalI and ligated to pCMV6.
The psiCHECK2-miR-221-sensor and psiCHECK2-miR-486-5p-sensor were constructed by annealing oligonucleotides with the reverse complement sequence of the mature microRNA with 5' XhoI and 3' NolI overhangs (IDT) and ligating to psiCHECK2 reporter. CCND2 and TWF1 3' UTRs were described previously [56]. Gene blocks (IDT) for wild type and miR-221 site mutant 3'UTRs (Supplementary Table S5) for CDK6 (6643-7109 bp) and ERBB3 (1-478 bp) were digested with XhoI and NolI and ligated to psiCHECK2 reporter. The miR-221 or miR-486-5p recognition sites in CCND2 and TWF1 were mutated with Quick Change II Site Directed Mutagenesis (200524; Agilent Technologies, Santa Clara, CA, USA). Luciferase reporter ligating to psiCHECK2 reporter.

Transfection of siRNA, microRNA mimics and inhibitors

Cells were transfected with 30 nM miRVana microRNA antimiRs or mimics (Thermo Fisher Scientific, Waltham, MA, USA) with inhibitors and mimics (Supplementary Table S7) as described previously [56]. The NC Dicer-substrate siRNA (51-01-14-03, IDT) and PAX3-FOXO1 siRNA (detailed in Supplementary Table S4) were transfected with 3 nM duplex as above.

In vivo tumorigenesis

Rh30 and Rh41 cells were transduced with miRZip-Scr or miRZip-486-5p, 48 h after infection. 1 × 10^6 Rh41 cells or 7.5 × 10^5 were subcutaneously injected in the flank of 8-week-old age-matched, randomized female SCID-Beige mice (n = 10 per group). Tumor growth measured every other day blinded to treatment once palpable tumors formed. Tumor volume calculated as previously described [24]. The tumor volume at each time point was compared using an unpaired Student's t-test. All studies involving animal studies were reviewed and approved by the St. Jude Children's Research Hospital Institutional Animal Care and Use Committee.

Immunohistochemistry

Histology and IHC performed following standard protocols using antibodies detailed in Supplementary Table S6 [24].

Statistical analysis

All results expressed as the mean ± SEM. All pair-wise comparisons utilized a two-tailed, unpaired Student’s t-test using GraphPad Prism Version 5 (Graph Pad Software, Inc., San Diego, CA, USA). P-values <0.05 considered significant. The number of mice required (n = 10 per group) for in vivo xenograft experiments was determined by power analysis using G Power 3.1 [78] with two sided, 90% power, Type I error of 1%.

Disclaimer

The content is solely the responsibility of the authors and does not necessarily represent the views of the NIH.

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**Compliance with ethical standards**

**Conflict of interest** The authors declare that they have no competing interests.

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