Growth-Promoting Role of the miR-106a∼363 Cluster in Ewing Sarcoma

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Abstract

MicroRNAs (miRs) have been identified as potent regulators of both normal development and the hallmarks of cancer. Targeting of microRNAs has shown to have preclinical promise, and select miR-based therapies are now in clinical trials. Ewing Sarcoma is a biologically aggressive pediatric cancer with little change in clinical outcomes despite improved chemotherapeutic regimens. There is a substantial need for new therapies to improve Ewing Sarcoma outcomes and to prevent chemotherapy-related secondary sequelae. Most Ewing Sarcoma tumors are driven by the EWS/Fli-1 fusion oncogene, acting as a gain-of-function transcription factor causing dysregulation of a variety of targets, including microRNAs. Our previous studies, and those of others, have identified upregulation of miRs belonging to the related miR-17~92a, miR-106b~25, and miR-106a~363 clusters in Ewing Sarcoma. However, the functional consequences of this have not been characterized, nor has miR blockade been explored as an anti-cancer strategy in Ewing Sarcoma. To simulate a potential therapeutic approach, we examined the effects of blockade of these clusters, and their component miRs. Using colony formation as a read-out, we find that blockade of selected individual cluster component miRs, using specific inhibitors, has little or no effect. Combinatorial inhibition using miR “sponge” methodology, on the other hand, is inhibitory to colony formation, with blockade of whole clusters generally more effective than blockade of miR families. We show that a miR-blocking sponge directed against the poorly characterized miR-106a~363 cluster is a particularly potent inhibitor of clonogenic growth in a subset of Ewing Sarcoma cell lines. We further identify upregulation of miR-15a as a downstream mechanism contributing to the miR-106a~363 sponge growth-inhibitory effect. Taken together, our studies provide support for a pro-oncogenic role of the miR-106a∼363 cluster in Ewing Sarcoma, and identify miR-106a∼363 blockade, as well as miR-15a replacement, as possible strategies for inhibition of Ewing Sarcoma growth.

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Introduction

Ewing Sarcoma (EWS) is the second most common solid bone and soft tissue malignancy in children and young adults. With mesenchymal progenitor cells (MPCs) as the presumed cell of origin, the vast majority of Ewing Sarcoma tumors are driven by EWS/Ets fusion oncogenes, with the EWS/Fli-1 fusion, arising from the t(11;22)(q24:12) translocation, being the most common. [1,2] EWS/Fli-1 is a non-physiologic, gain-of-function transcription factor that activates and represses a number of pathways including regulation of cell cycle progression, and TGF-1 Receptor (IGF-1R) pathway, Cyclin D1 and p21 regulation of mitotic and apoptosis, and IGF-1R targeted therapy, many EWS patients either fail to respond to treatment or develop resistance. One alternative avenue of targeted therapies with pre-clinical promise in a number of cancers, but relatively unexplored in Ewing Sarcoma, is modulation of microRNAs [14–16].

MicroRNAs (miRs) are short non-coding RNAs that bind a 2–7 nucleotide sequence (“seed sequence”) within the 3‘ untranslated regions (UTR) to mediate gene repression via degradation or sequestration of the targeted mRNA. [17] MiRs have tremendous regulatory power, potentially regulating over 60% of the genome, including regulation of normal cellular functions and the hallmarks of cancer. [18] The exact role of any given miR depends on the cell type and possibly other, currently largely unknown, regulatory factors.

While EWS is a chemosensitive-tumor, these treatments place primary outcomes and long-term quality of life. [11] However, with the exception of IGF-1R targeted blockade, understanding EWS/Fli-1-mediated tumorigenesis has yielded few therapeutic alternatives to date. [12,13] Furthermore, despite the promise of IGF-1R targeted therapy, many EWS patients either fail to respond to treatment or develop resistance. One alternative avenue of targeted therapies with pre-clinical promise in a number of cancers, but relatively unexplored in Ewing Sarcoma, is modulation of microRNAs [14–16].

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Several studies examining miR alterations in Ewing Sarcoma have identified upregulation of miRs belonging to three paralogous clusters – miR-17~92a, miR-106b~25, and miR-106a~363. [4–6] These clusters are highly conserved across...
species, are thought to have arisen from genetic duplications during evolution, and have been shown to be pro-oncogenic in a number of malignancies. [19–27] In EWS, tumor expression of multiple members of these clusters has recently been shown to be negatively correlated with both 5-year event-free survival and overall survival. [28] Thus, manipulation of these clusters, or their component miRs, may have therapeutic benefit in Ewing Sarcoma. The purpose of the present study was to determine the requirements for miR cluster upregulation in Ewing Sarcoma, and identify strategies for their blockade to inhibit Ewing Sarcoma oncogenesis.

Results

The miR-17–92a, miR-106b–25, and miR-106a–363 Clusters are Overexpressed in Ewing Sarcoma Cell Lines Compared to Human Mesenchymal Progenitor Cells, the Presumed Cells of Ewing Sarcoma Origin

Our previous studies profiling miRs downstream of the EWS/Fli1 fusion oncoprotein identified several members of the miR 17–92a, 106b–25 and 106a–363 clusters as candidate upregulated miRs in Ewing Sarcoma. [4] These miR clusters have been demonstrated to play important roles, largely pro-oncogenic, in a broad array of cancers. [19–27] We were thus very interested in further understanding their potential role in Ewing Sarcoma. We began by surveying expression levels of all the component miRs in a panel of Ewing Sarcoma cell lines (A673, EWS502, TC71, Sk-N-Mc, Sk-ES-1, and RD-ES) relative to two different human mesenchymal progenitor cell (hMPC) lines, the presumed cells of Ewing Sarcoma origin, using qRT-PCR (Figure 1b–d). Overall, we found that Ewing Sarcoma cell lines consistently upregulated all the cluster component miRs, with the exception of miR-363, which was only minimally expressed in both EWS cells and hMPCs. Some cell line differences were evident, with A673 and EWS502 cells tending toward lower levels of upregulation, and TC71, Sk-N-Mc, Sk-ES-1 and RD-ES cells tending toward greater upregulation of the clusters. We next compared the relative expression levels of individual miR cluster members within the various cell lines, by correcting for any differences in qRT-PCR primer efficiency and examining the absolute copy number of each miR relative to the copy number of an endogenous U6 RNA control. This revealed miR-92a, miR-93, miR-25, and miR-106a to be the most highly expressed miRs between studied miRs in both hMPCs and EWS cell lines (Figure 2a and 2b).

MiR Blocking Experiments Identify a Growth-promoting Role for the 106a–363 Cluster in Ewing Sarcoma

We next sought to determine the functional consequences of miR overexpression in Ewing Sarcoma. Overexpression of the miR clusters and/or component miRs has been shown to promote growth in a number of cancer types. [22,25–27,29–35] We thus postulated that miR cluster overexpression would be growth-promoting in Ewing Sarcoma. We began by examining the effects of blockade of select individual cluster component miRs. We chose a clonogenic assay as a stringent assay of cell growth to screen for effects of miR inhibition. MiR inhibition was achieved using either hairpin inhibitor (HI) or locked nucleic acid (LNA) methodology, and verified using the psiCHECK2 dual luciferase reporter system. We compared these two methodologies (HI and LNA) for select miRs and did not observe differences in the effectiveness of miR blockade (Figure S1). Inhibition of miRs 17, 92a, 93 and 25 was chosen because of the robust overexpression of these miRs in EWS cells compared to hMPCs and/or their high expression levels in both cell types. In addition, miR-19b inhibition was also selected because, while not expressed at high levels, this miR has been shown in other systems to be sufficient for the pro-oncogenic activity of the entire miR-17–92a cluster. [25,29,36] Sk-ES-1 cells were chosen because of their tendency toward high miR cluster overexpression. Individual blockade of miR-93, miR-25, miR-17, miR-92b, or miR-92a did not show a significant effect on Sk-ES-1 clonogenic growth (Figure S2). Individual blockade of miRs 25 and 93 was also tested in two other EWS cell lines (TC71 and Sk-N-MC), but, similar to the findings in Sk-ES-1 cells, did not significantly affect clonogenic growth (Figure S2b–c). Inhibition of miRs 25 and 93 in combination tended to show a very slight inhibition of growth in some experiments, but this did not reach statistical significance (Figure S2a). In all cases, the dosage of miR inhibitor used appeared to be sufficient to effect miR blockade in Sk-ES-1 cells, as determined using the psiCHECK2 reporter system (Figure S2d and S2f).

Given the lack of effects by targeting individual miRs, but the possible slight effect seen by inhibition of multiple miRs, we examined the possibility that miRs may be acting cooperatively. MicroRNAs can be grouped into clusters containing miRs that are co-expressed based on their locations within the genome, as well as into families, which contain miRs with a shared seed sequence. We explored the possibility that miRs with related seed sequences from different clusters compensate for each other, or, alternatively, that miRs unrelated by sequence, but co-expressed from the same cluster, may work together. To further probe this in Ewing Sarcoma, and distinguish between these two scenarios, we took advantage of the recently developed miR “sponge” technology, which allows simultaneous blockade of multiple miRs. [37] Specifically, an RNA molecule is engineered with miR complementarity to sequences resembling a perfect target, including a mismatched bulge to prevent degradation of the miR-RNA duplex. This system has been used effectively in a number of studies to understand miR function. [37] For our experiments, we employed the lentiviral pGreen expression system to drive sponge expression; this system uses the RNA polymerase III-driven H1 promoter driving the expression of multiple bulged miR binding sites. The modular composition of the different miR sponge constructs used is shown in Figure 3. Specifically, the sponge constructs generated were: s-3-miR-18, s-z-miR-19, s-z-miR-20, and s-z-miR-25, to target all the members of the specified miR family, as defined by shared seed sequence; and s-z-miR-17–92a, s-z-miR-106b–25, and s-z-miR-106a–363, to target all the members of a given miR cluster. Additionally, given the established importance of the miR-17–92a and miR-106b–25 clusters in other cancers, the combination sponge construct s-z-17–92a/106b–25 was generated to target these clusters in tandem. For negative controls, two different constructs were used, s-Empty, lacking any miR-binding sites, and s-Neg, a non-targeting control that is based on a sequence that no miRs are predicted to bind, but that still resembles a potential miR binding site, as done by others previously. [37] Sk-ES-1 cells were again selected for screening experiments, as a cell line with robust endogenous miR cluster overexpression (see Figure 1). Cells were stably infected with the above constructs and, following antibiotic (Puromycin) selection, levels of expression of each were determined by qRT-PCR, using the same primer pair to permit comparison between constructs (Figure 4a). These analyses revealed all of the miR-targeting sponges to be expressed at comparable levels to each other. The relative level of the s-Neg control was more variable and overall higher than the levels of the sponge constructs, possibly reflecting some degree of negative selection against the miR-targeting constructs.
We next examined the phenotypic effects of miR-blocking sponge expression on Sk-ES-1 clonogenic cell growth (Figure 4b). Clonogenic growth was similar for both the s-Empty and s-Neg expressing cells. Of the sponges targeting miR families (with related seed sequence), only s-a-miR-25 had an effect on clonogenic growth, causing a 20% decrease in colony formation, compared to both controls (s-Empty and s-Neg). This was interesting given the absence of a phenotype upon specific blockade of miRs 25 and 92a alone (see Figure S2), suggesting that members of the miR-25 family may indeed be able to compensate for one another. The other miR family targeting sponges, s-a-miR-18, s-a-miR-19 and s-a-miR-20, had no significant effect on clonogenic growth of Sk-ES-1 cells. Of the miR cluster targeting sponges (targeting co-expressed miRs, but with unrelated seed sequences), the one targeting the miR-17,92a cluster caused a 20% decrease in colony formation. The sponge targeting the miR-106b–25 cluster had no effect on clonogenic growth. Furthermore, the effect on growth of the combination miR-17,92a/miR-106b–25 sponge was essentially identical to the miR-17,92a targeting sponge. Interestingly, of the cluster-targeting sponges, the one targeting the miR-106a–363 cluster showed the most potent inhibitory effect, with an average 50% decrease in clonogenic growth. The s-a-miR-106a–363 sponge also yielded the most potent inhibition of colony formation in a soft agar assay of anchorage-independent growth (Figure 4c). To verify that the s-a-miR-106a–363 sponge was acting through a miR-blocking mechanism, a construct was generated containing mutated seed sequence for each of the miRs in the cluster. Stable expression of this seed-mutated sponge construct resulted in loss of inhibition of clonogenic growth relative to the (non-mutated) s-a-miR-106a–363 sponge in Sk-ES-1 cells (Figure S3). Thus, miR cluster blockade appears to be overall more effective as a means of inhibiting cell growth than miR family blockade in Ewing Sarcoma, with blockade of the miR-106a–363 cluster showing the highest potency.

We next tested the inhibitory activity of the s-a-miR-106a–363 sponge in additional EWS cell lines. Similar to Sk-ES-1 cells, stable expression of s-a-miR-106a–363 inhibited both clonogenic and anchorage-independent growth of RD-ES cells (Figure 5a–c). Introduction of the s-a-miR-106a–363 sponge into A673 or TC71 cells, on the other hand, did not affect clonogenic growth. Interestingly, both cell lines vulnerable to the sponge effect appeared to restrict sponge expression (Sk-ES-1 and RD-ES), further suggesting that introduction of s-a-miR-106a–363 is deleterious in these cells (Figure 5d).

Upregulation of miR-15a Contributes to the Growth Inhibitory Effects of miR-106a–363 Blockade in Ewing Sarcoma

MicroRNAs have many targets in the cell, and the relative importance of individual miR-target interactions varies among cell types. A number of targets have been identified for the miR-17–92a and miR-106b–25 clusters in other cancers. Less is known about the biology of the miR-106a–363 cluster and few miRs have been functionally evaluated in sarcomas. To probe for relevant mechanisms of action of the s-a-miR-106a–363 sponge in Ewing Sarcoma, we examined the number of targets, and/or related pathway activity, of miR-106a–363, as
well as the closely related miR-17~92a, cluster, which have been identified in other cancers as regulators of growth and apoptosis. These targets/pathways included PTEN and the PI3K/Akt pathway, Erk1/2 and MAPK signaling, TGF \( \beta \) signaling, and Wnt signaling. However, this approach failed to reveal a candidate mechanism of s-a-miR-106a,363 sponge action. We thus employed an unbiased approach to identify targets and/or pathways affected by s-a-miR-106a,363 sponge expression in Ewing Sarcoma. Expression profiling using Affymetrix whole transcript arrays was performed on Sk-ES-1 cells expressing either s-Neg or s-a-miR-106a,363. Interestingly, this analysis revealed the transcript containing miR-15a as significantly upregulated in the context of s-a-miR-106a,363 sponge action. We thus used an LNA approach to determine whether miR-15a blockade can reverse the growth-inhibitory effect of the s-a-miR-106a,363 sponge. In both Sk-ES-1 and RD-ES cells, treatment with LNA-anti-miR-15a resulted in at least partial rescue of growth inhibition by s-a-miR-106a,363 (Figure 7). The effect was more robust in RD-ES cells where colony formation by s-a-miR-106a,363/anti-miR-15a LNA cells was both similar to s-Neg/negative control LNA cells, and statistically greater than colony formation by s-a-miR-106a,363/negative control LNA cells (Figure 7b). In Sk-ES-1 cells, colony formation by s-a-miR-106a,363/anti-miR-15a LNA cells was similar to colony formation by s-Neg/negative control LNA cells; however, while, there was a trend toward increased colony formation relative to the s-a-miR-106a,363/negative control LNA group, this did not reach statistical significance (Figure 7a). Interestingly, LNA inhibition of miR-15a, as determined using the psiCHECK2 reporter system, was more potent in RD-ES cells (Figure 8a), suggesting this as the possible basis for the more robust rescue in this cell line. Taken together, these findings suggest that miR-15a upregulation contributes to the growth-inhibitory effects of miR-106a~363 blockade in Ewing Sarcoma.

To further explore the role of miR-15a in Ewing Sarcoma, we asked how miR-15a manipulation affects clonogenic growth in the absence of miR-106a~363 inhibition. Inhibition of miR-15a in Sk-ES-1 and RD-ES cells, using the same LNAs as above, resulted in increased clonogenic growth compared to a non-targeting
negative control LNA (Figure 8b). As in the rescue experiments above, LNA-anti-miR-15a treatment resulted in a greater increase in colony formation in RD-ES cells compared to Sk-ES-1 cells. Again, this may be due in part to greater potency of LNA-mediated miR-15a inhibition in RD-ES cells compared to Sk-ES-1 cells (Figure 8a). Conversely, transient overexpression of miR-15a, using a miR mimic, resulted in reduction of Sk-ES-1 and RD-ES clonogenic growth compared to a scrambled negative control miR mimic (Figure 8c and 8d). As in the LNA experiments, the more potent phenotypic effect of miR-15a mimic treatment in RD-ES cells correlated with higher miR-15a overexpression levels in this cell line. These findings further support a growth suppressive role of miR-15a in Ewing Sarcoma.

Discussion

Targeting of miRs through delivery of LNAs or antagomiRs has recently been identified to have substantial promise in pre-clinical models, due to the specificity and relative lack of off-target effects as are seen with traditional chemotherapeutics. Currently, one LNA-based therapy is in phase 2 clinical trials and showing substantial promise. [43] Our previous studies identified members of the three paralogous oncomiR clusters, miR-17–92, miR-106b–25, and miR-106a–363, among the most strongly EWS/Fli1-upregulated miRs. [4] Other miR profiling studies in Ewing Sarcoma have made similar observations, and also verified oncomiR cluster overexpression in patient tumors. [5–8] Moreover, in a recent study, overexpression of several members of these clusters has been shown to correlate with both poor 5-year event free survival and overall survival. [28] Similarly, in Alveolar Rhabdomyosarcoma, another pediatric sarcoma, tumor expression of multiple members of these clusters has been shown to be negatively correlated with patient prognosis. [24] In the present study, we undertook the first functional analysis of these clusters in Ewing Sarcoma, as well as the first systematic analysis of the effects of blockade of miR genomic clusters versus their seed sequence-related components. These studies have identified an important role for the miR-106a–363 cluster in the promotion of clonogenic and anchorage-independent growth in Ewing Sarcoma, and have
Figure 4. Effects of miR-blocking sponge expression on Ewing Sarcoma cell growth. (A) Sponge expression levels in Sk-ES-1 cells as determined by qRT-PCR. Results represent the mean and SEM of three independent experiments, each performed in triplicate. (B) Colony formation by Sk-ES-1 cells stably transduced with the indicated miR-blocking sponge constructs, in a clonogenic assay. Data represent the mean and SEM of two independent experiments, each performed in triplicate. *p<0.05 compared to s-Empty and s-Neg, individually. (C) Colony formation by Sk-ES-1 cells stably transduced with the indicated miR-blocking sponge constructs, in a soft agar assay. Results represent the mean and SEM of three independent experiments, each performed in triplicate. *p<0.05 compared to s-Empty and s-Neg, individually; †p<0.05 compared to s-Empty only based on an unpaired student’s t-test. doi:10.1371/journal.pone.0063032.g004

Figure 5. Growth effects of the s-α-miR-106a~363 sponge in other Ewing Sarcoma cell lines. Comparison of colony formation in a panel of EWS cell lines (Sk-ES-1, RD-ES, A673, and TC71) stably transduced with s-Neg or s-α-miR-106a~363, in a clonogenic assay (A) and in a soft agar assay (B). Results represent the mean and SEM of a minimum of three independent experiments, each performed in triplicate. *p<0.05 compared to s-Neg based on an unpaired student’s t-test. (C) Sponge expression in EWS cell lines was determined by qRT-PCR. Data represent the mean and SEM of a minimum of three independent experiments, each performed in triplicate. doi:10.1371/journal.pone.0063032.g005
uncovered miR-15a as a novel contributor to oncomiR cluster action in cancer.

MicroRNAs can be grouped into miR clusters based on their locations within the genome or into miR families based on the presence of shared seed sequences. Within a given family, most miRs differ by only a few nucleotides, leaving the potential for redundancy among family members. There is also a high degree of similarity between the miR-17–92a and miR-106a–363 clusters, with both miR-19b and miR-92a being absolutely duplicated between the two clusters. To date, there have been few systematic analyses of these clusters and their component miR families. In developmental studies, the miR-106a–363 cluster is dispensable for normal mouse development. [44] However, double knock out experiments with just the miR-17–92a and the miR-106a–363 clusters were not performed, only the triple knock out experiment for all three clusters and the double knock out for both miR-17–92a and miR-106b–26. This leaves the possibility that the exaggerated developmental phenotypes seen in the miR-17–92a/miR-106b–25/miR-106a–363 triple knock out mice could still be partially contributed to by the loss of the miR-106a–363 cluster. Our data using microRNA-blocking sponges to examine the effects of inhibition of entire miR clusters compared to inhibition of miR families, the first analysis of this type in a pediatric cancer, suggest that, at least in Ewing Sarcoma, inhibition of all the members of a given cluster may be more potent than inhibition of paralogous miRs with shared seed sequences. This in turn suggests that the co-expressed functionally distinct miRs cooperate to promote oncogenesis and that targeting these miRs together may provide the most potent blockade of oncogenesis.

The miR-17–92a cluster has been implicated in numerous cancer types, while the role of the miR-106a–363 cluster is present relatively obscure. We were, thus, somewhat surprised by the more potent inhibitory activity of the miR-106a–363-targeting sponge, relative to the sponge targeting the miR-17–92a cluster. One possibility is that miR-106a–363 is indeed more important than miR-17–92a in Ewing Sarcoma pathogenesis. However, it is also possible that s-α-miR-106a–363 and s-α-miR-17–92a cross-react, at least to some extent, given how closely related the miRs are from the two clusters. Thus, we cannot exclude the possibility that the s-α-miR-106a–363 sponge works at least in part by blocking miRs from the miR-17–92a cluster. Even if the sponges do to some extent cross-react, however, the more potent activity of s-α-miR-106a–363 supports a role for the miR-106a–363 cluster in Ewing Sarcoma.

While the miR-106a–363 cluster is by far the least studied of the paralogs, an increasing number of studies are emerging which support the idea that it plays an important role in tumorigenesis. In a bronchial epithelial chemical carcinogenesis model and in gastric cancer models, miR-106a promotes both in vitro and in vivo tumorigenesis. [22,31] Furthermore, three members of the miR-17 family (miR-17, miR-106b and miR-106a) each directly inhibit p21 in Diffuse Large B-cell Lymphoma and in Burkitt’s Lymphoma to increase cellular proliferation. [23] Importantly, in this context, specific blockade of miR-106a alone also resulted in increased expression of Bim and decreased expression of CDK4/CDK6. Together, these studies support an important and distinct role for miR-106a in tumorigenesis. Interestingly, in our studies, inhibition of miR-106a alone, or in combination with inhibition of miR-92a or miR-20b, using LNAs, did not significantly inhibit Sj ES-1 clonogenic growth. This suggests that inhibition of the entire miR-106a–363 cluster may be required for the growth-inhibitory effects of the s-α-miR-106a–363 sponge in Ewing Sarcoma.

The growth inhibitory effects of s-α-miR-106a–363 demonstrate cell specificity, affecting growth of Sk-ES-1 and RD-ES, but not A673 and TC71 cells. The degree of miR-106a–363 overexpression (high in Sk-ES-1 and RD-ES cells, but lower in A673 cells) may explain the difference in phenotypes between these cell lines. However, other factors must be responsible for the lack of phenotype in TC71 cells, which manifest miR-106a–363 overexpression resembling Sk-ES-1 and RD-ES cell. Interestingly, both Sk-ES-1 and RD-ES cells contain the EWS/Fli-1 type 2 fusion, while A673 and TC71 cells bear the EWS/Fli-1 type 1 fusion. This suggests the intriguing possibility that fusion type may be a contributory factor to the observed cell type specificity. The EWS/Fli-1 type 1 fusion accounts for approximately 60% of all EWS/Fli-1 translocations, with the type 2 fusion accounting for an additional 25%. [45] These translocations differ in the number of exons from the C-terminus of Fli-1 (type 1 containing exons 6–9 and type 2 containing exons 5–9). Originally, EWS/Fli-1 fusion subtype was studied in part because a type 1 fusion appeared to confer a better prognosis. [46] Additionally, in mouse xenograft experiments, cell lines bearing the type 1 fusion had delayed tumor initiation and resulted in fewer primary tumors and metastases. [45] In microarray expression profiling of type 1 fusion compared to non-type 1 fusion bearing cells revealed 41 genes were differentially expressed, with all being downregulated in the non-type 1 fusion bearing cell lines. [47] These genes included genes involved in muscle development, proliferation, and calcium-ion binding, among others. However, the functional consequences of these differences have not been studied in detail. Moreover, more recent patient data suggest no difference in prognosis or chemosensitivity between type 1 and non-type 1 fusion-bearing tumors with current treatment protocols. [48] Our data suggest possible differences in microRNA biology between these groups, but this possibility awaits further experimental exploration.

Our studies demonstrate a growth-inhibitory role for miR-15a in Ewing Sarcoma. MiR-15a has been shown to be tumor suppressive in other systems, primarily through regulation of a variety of cell cycle targets, including Wee1 and multiple cyclins. [38,39,41,42] In mouse genetic models, miR-15a deletion significantly accelerates the development of Chronic Lymphocytic Leukemia, in part through de-repression of multiple miR-15a-
targeted cyclins (Cyclin D1 and D3, Cyclin E), CDK6 and the anti-apoptotic factor Bcl-2. In prostate cancer, inhibition of miR-15a leads to increased anchorage-independent growth and migration in vitro, as well as transforming a non-tumorigenic prostate cell line in vivo. On the other hand, intratumoral injection of a miR-15a inhibitor induces tumor necrosis and regression. Again, the identified miR-15a targets included Cyclin D1 and Bcl-2. Not only does miR-15a negatively regulate cell cycle progression through inhibition of CDKs and cyclins, but its inhibition of two cyclin kinases, Wee1 and Chk1, is associated with Cisplatin resistance in cancer cell lines and this resistance is reversed upon re-expression of miR-15a. Finally, in Ewing Sarcoma, treatment with MLN4924, a compound that inhibits neddylation, and subsequent degradation, of cullins in cullinRING ubiquitin ligase complexes leading degradation of a variety of proteins, leads to a G2 cell cycle arrest and apoptosis. Interestingly, two miR-15a targets, Wee1 and Cyclin E, were increased under such conditions, with Wee1 expression being required to mediate the G2 arrest. The exact mechanism of Wee1 accumulation was not determined. In our microarray expression profiling of s-Neg and s-a-miR-106a~363 expressing Sk-ES-1 cells, GSEA phenotypic analysis revealed the KEGG cell cycle pathway as being differentially expressed. Somewhat unexpectedly, many genes in this pathway were overall upregulated by s-a-miR-106a~363 expression, including two known miR-15a targets, Cyclin E1 and E2, and E2F1. This is not entirely surprising, however, in light of data from osteosarcoma, where E2Fs, Cyclin E, and miR-15a form a complex regulatory loop, whereby E2Fs induce both Cyclin E and miR-15a expression, and miR-15a limits proliferation by inhibiting Cyclin E expression during G1/S.

Figure 7. MiR-15a blockade reverses growth inhibition by s-a-miR-106a~363. Clonogenic growth assay in Sk-ES-1 (A) and RD-ES (B) cells stably expressing s-Neg or s-a-miR-106a~363 that were untreated or treated with 50 nM of a negative control LNA or an LNA targeting miR-15a. Results represent the mean and SEM of three independent experiments, each performed in triplicate. *p<0.05; n.s. = not significant; based on an unpaired student’s t-test.

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The miR-17–92 paralogous clusters, are also induced by E2F1 expression and can act to limit proliferation. [35,52,53] Thus, perturbation of individual components of these regulatory loops may have complex effects on the network as a whole.

Our studies indicate that miR-15a levels are also sensitive to perturbation of oncomiR cluster levels. The precise mechanism(s) of miR-15a upregulation by miR-106a~363 blockade remain to be determined. MiR-15a can be regulated transcriptionally. [41,54] However, we did not observe consistent increases in Dleu2, the miR-15a primary transcript, by qRT-PCR in s~a-miR-106a~363 sponge expressing cells, arguing against a major role for a transcriptional mechanism. Interestingly, miR-15a expression in the mouse has recently been shown to be regulated by another microRNA, miR-709, at the level of processing, and similar mechanisms may play a role in Ewing Sarcoma. [55,56].

In summary, we demonstrate that members of the miR-17–92a, miR-106b~25, and miR-106a~363 clusters are upregulated in EWS. Our systematic functional analysis of these paralogous clusters, using miR blocking sponge methodology, identifies the miR-106a~363 cluster as a potentiator of EWS growth. This potentiation manifests cell type specificity, possibly in part related to cellular context related to EWS/Fli1 fusion type, as well as likely other factors. In cell lines sensitive to the growth inhibitory effects of miR-106a~363 blockade, modulation of miR-15a contributes to these effects. Thus, blockade of the miR-106a~363 cluster and/or replacement of miR-15a represent possible new strategies for inhibition of Ewing Sarcoma growth.

Materials and Methods

Cell Lines and Culture

Ewing Sarcoma cell lines A673, Sk-N-MC, Sk-ES-1 and RD-ES were obtained from ATCC. Ewing Sarcoma cell lines EWS502 and TC71 were obtained from Steve Lessnick at the University of...
Utah. A673, Sk-N-MC, Sk-ES-1, and EWS502 cells were grown in DMEM supplemented with 10% fetal bovine serum (FBS). TC71 cells were grown in RPMI supplemented with 10% FBS. RD-ES cells were grown in RPMI supplemented with 15% FBS. Low-passage primary human mesenchymal progenitor cells (hMPCs) were obtained from SciCell and Lonza, and cultured in proprietary media.

MicroRNA Expression Analysis

Total cellular RNA was isolated using the TRIzol reagent, per manufacturer instructions. For each group, RNA was harvested in biological triplicates from plates at similar confluence (50–70%). cDNA was synthesized from 1 μg total cellular RNA using the Qiagen miScript II Reverse Transcriptase kit (Qiagen, Cat #219061), and quantification of microRNA expression levels was performed using the Qiagen SYBRgreen qRT-PCR system (Qiagen, Cat #219075). The relative degree of miR expression between cell lines for a single miR was calculated using the equation: 

\[ \Delta \Delta CT = (C_{TXM_iR} - C_{U6}) \]

In order to correct for primer efficiency and compare the absolute level of different miRs within a cell line, the best fit linear equation generated by the amplicon standard curve (1.2 x 10^2 to 7.7 x 10^6 copies/μl) was used to determine the number of copies of the individual miRs and U6 in 333 ng of cDNA as previously described [57].

In vitro Cell Growth Assays

For clonogenic assays, EWS cells were plated at a density of 500 cells/well in 6-well plates. After 10–14 days, the cells were washed with PBS and then stained with 0.1% crystal violet in 10% methanol. Colonies were quantified using NIS-Elements System Software. For anchorage-independent colony formation assays, 50,000 cells/well in 6-well plates were grown in 0.5% agar (Difco Agar Noble BD 214230) and growth medium containing 20% FBS. Colonies were stained with Nitroblue Tetrazolium Chloride, as previously described [4] and quantified using the NIS-Elements System Software. Statistical significance between the control group(s) and the indicated treatments was determined based on an unpaired student’s t-test.

Stable miR Sponge Experiments

CMV-d2eGFP sponge plasmids for CXCR4 and miR-18 were obtained from the Sharp lab at MIT [37]. The CXCR4 control is a non-targeting miR sponge based on a sequence from CXCR4 that no miRNAs are predicted to target, as described [37] and used in a number of studies [58]; in our manuscript, we refer to this control sponge as s-Neg. MiR targeting sequences with a nucleotide bulge were designed as described [37] and obtained from Integrated DNA Technologies for the miR-17~92a, miR-106b~25, and/or miR-106a~363 clusters and the miR-19, miR-20, and miR-25 families. Complementary oligos were annealed and subcloned into CMV-d2eGFP, using standard molecular methods and sequence verified. The bulged miR-binding sites were then subcloned into the pGreen-lentiviral expression vector (System Biosciences), using standard molecular methods and sequence verified. Infectious virus was prepared as previously described. [4] EWS cells were infected with similar titers of virus and selected with Puromycin (2 μg/ml for A673, and 0.5 μg/ml for Sk-ES-1, TC71, and RD-ES). Following 3–7 days of selection, cells were used for experiments and RNA was harvested to check sponge expression.

Gene Expression Profiling

Biological triplicates of Sk-ES-1, RD-ES, and TC71 cells stably transduced with either s-CXCR4 or s-s-miR-106a~363 were harvested at ~70–80% confluence. Total RNA was harvested using the Qiagen RNeasy Mini Kit according to the manufacturer's protocols. RNA concentration was determined spectrophotometrically. The quality and integrity of RNA were verified using the Agilent 2100 Bioanalyzer. For microarray analysis, 250 ng of total RNA was processed using the Whole Transcript Expression Kit (Ambion) and Whole Transcript Terminal Labeling kit (Affymetrix). Samples were hybridized to Human Gene 1.1 ST array strips (Affymetrix) and washed, stained, and imaged using the Gene Atlas Personal Microarray System (Affymetrix). Resulting CEL files were RMA normalized and Log2 transformed using Partek Genomics Suite, and differentially expressed genes were identified using Significance Analysis of Microarrays (SAM) version 4.0 in Excel (www-stat.stanford.edu/~tibs/SAM) with a false discovery rate of 25%. [59] Normalized data were also analyzed using Gene Set Enrichment Analysis Software (http://www.broadinstitute.org/gsea/doc/GSEAUserGuideFrame.html), available as a stand-alone Java application. The expression profiling data discussed in this publication have been deposited in NCBI's Gene Expression Omnibus and are accessible through GEO Series accession number GSE45205 (http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE45205).

Supporting Information

Figure S1 Comparison of miR inhibition using a Hairpin Inhibitor or LNA targeting miR-19b. Dual Renilla/Luciferase assay with miR-19 target in the 3’ UTR of renilla. Sk-ES-1 cells were transfected with 20 nM of a negative control Li, 20 nM of a miR-19b targeting Hi, or 50 nM of a miR-19b targeting LNA. Results represent the mean and standard deviation of two experiments performed in triplicate. (TIF)

Figure S2 Effects of individual miR blockade on Ewing Sarcoma clonogenic growth. (A–C) Clonogenic assay in Sk-ES-1 (A), TC71 (B), or Sk-N-Mc (C) cells transfected with 20 nM of a negative control hairpin inhibitor or a hairpin inhibitor (Hi) targeting miR-25, miR-93, or miR-25 and miR-93. (D) Dual Renilla/Luciferase assay performed in Sk-ES-1 cells transfected
with 20 nM Hi-miR-K25 and/or Hi-miR-93, and the psiCHECK2 dual luciferase reporter with a corresponding complementary binding site in the 3' UTR of Renilla. (E) Clonogenic assay in Sk-ES-1 cells transfected with 100 nM of a negative control LNA or an LNA targeting mir-17, mir-19b, or mir-92a. (F) Dual Renilla/Luciferase transfected with 100 nM of a negative control LNA or an LNA targeting miR-17, miR-19b, or miR-92a, and the psiCHECK2 dual luciferase reporter with a corresponding complementary binding site in the 3' UTR of renilla. All values represent the mean and SEM of a minimum of two independent experiments, each performed in triplicate.

**Figure S3** Seed sequence mutation of miR-106a~363 binding sites abolishes growth inhibitory effects of s-a-miR-106a~363. (A) Sponge expression was determined in Sk-ES-1 cells stably transduced with s-Neg, s-miR-106a~363 seed mut. or s-a-miR-106a~363 by qRT-PCR. Results represent the mean and SEM of three independent experiments, each performed in triplicate. *p<0.05 compared to s-Empty according to an unpaired student’s t-test.

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**Author Contributions**
Conceived and designed the experiments: LD PJ. Performed the experiments: LD. Analyzed the data: LD PJ. Contributed reagents/materials/analysis tools: LD PJ. Wrote the paper: LD PJ.

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