#### www.nature.com/oncsis

# **SHORT COMMUNICATION**

# Comparative transcriptomic analysis reveals the oncogenic fusion protein PAX3-FOXO1 globally alters mRNA and miRNA to enhance myoblast invasion

JM Loupe<sup>1,4,6</sup>, PJ Miller<sup>1,5,6</sup>, BP Bonner<sup>1,6</sup>, EC Maggi<sup>1</sup>, J Vijayaraghavan<sup>1</sup>, JS Crabtree<sup>1</sup>, CM Taylor<sup>2</sup>, J Zabaleta<sup>3</sup> and AD Hollenbach<sup>1</sup>

Rhabdomyosarcoma, one of the most common childhood sarcomas, is comprised of two main subtypes, embryonal and alveolar (ARMS). ARMS, the more aggressive subtype, is primarily characterized by the t(2;13)(p35;p14) chromosomal translocation, which fuses two transcription factors, PAX3 and FOXO1 to generate the oncogenic fusion protein PAX3-FOXO1. Patients with PAX3-FOXO1-postitive tumors have a poor prognosis, in part due to the enhanced local invasive capacity of these cells, which leads to the increased metastatic potential for this tumor. Despite this knowledge, little is known about the role that the oncogenic fusion protein has in this increased invasive potential. In this report we use large-scale comparative transcriptomic analyses in physiologically relevant primary myoblasts to demonstrate that the presence of PAX3-FOXO1 is sufficient to alter the expression of 70 mRNA and 27 miRNA in a manner predicted to promote cellular invasion. In contrast the expression of PAX3 alters 60 mRNA and 23 miRNA in a manner predicted to inhibit invasion. We demonstrate that these alterations in mRNA and miRNA translate into changes in the invasive potential of primary myoblasts with PAX3-FOXO1 increasing invasion nearly 2-fold while PAX3 decreases invasion nearly 4-fold. Taken together, these results allow us to build off of previous reports and develop a more expansive molecular model by which the presence of PAX3-FOXO1 alters global gene regulatory networks to enhance the local invasiveness of cells. Further, the global nature of our observed changes highlights the fact that instead of focusing on a single-gene target, we must develop multi-faceted treatment regimens targeting multiple genes of a single oncogenic phenotype or multiple genes that target different oncogenic phenotypes for tumor progression.

Oncogenesis (2016) 5, e246; doi:10.1038/oncsis.2016.53; published online 25 July 2016

# INTRODUCTION

Rhabdomyosarcoma (RMS), which accounts for nearly half of childhood soft tissue sarcomas, is comprised of two main subtypes: embryonal rhabdomyosarcoma and alveolar (ARMS), each defined by its unique histology, clinical presentation and prognosis.<sup>1</sup> ARMS, the more aggressive subtype, is primarily defined by the t(2;13)(p35;p14) chromosomal translocation,<sup>2</sup> which generates the oncogenic fusion protein PAX3-FOXO1.<sup>3,4</sup> PAX3-FOXO1 has altered molecular activities relative to wild-type PAX3, including being a more potent transcriptional activator,<sup>5</sup> being unresponsive to normal PAX3 co-regulators<sup>6</sup> and having greater post-translational stability upon the induction of myogenic differentiation.<sup>7</sup> These aberrant molecular activities are believed to contribute to altered gene regulation, including the activation of genes not normally regulated by PAX3<sup>8</sup> and increased expression of other genes relative to PAX3,<sup>9,10</sup> which taken together is believed to contribute to ARMS tumor phenotypes.<sup>11</sup>

Patients diagnosed with PAX3-FOXO1-positive ARMS have a 4-year survival rate of 8%. <sup>12</sup> This poor prognosis stems in part from these tumor cells having a higher incidence of localized

invasion,<sup>12</sup> which may then lead to heightened aggressiveness and an increased propensity for metastasis. The presence of PAX3-FOXO1 is known to enhance the invasive potential of cells, 13 possibly through its ability to alter the expression of multifunctional genes that contribute, in part to invasion in other tumor types, including MET,<sup>10</sup> FGFR4,<sup>14</sup> IGF2<sup>15</sup> and IGFBP5.<sup>15</sup> Despite these circumstantial correlations, to date only a single report demonstrates that the PAX3-FOXO1 altered expression of a gene, the cannabinoid receptor 1, directly contributes to the invasive capacity in ARMS. 16 However, these results were derived from the expression of the oncogenic fusion protein in established tumor cell lines<sup>13</sup> or in primary myoblasts that genetically contained compensatory oncogenic mutations.<sup>16</sup> Further, these reports either did not examine altered gene expression 13 or focused their study on changes in the expression of a single gene. 16 While these reports are noteworthy and of importance, they provide little information to describe how the expression of PAX3-FOXO1 in the absence of any other compensatory mutations globally alters mRNA expression patterns to contribute to invasion. Further, to date no studies have directly examined how the presence of

<sup>1</sup>Department of Genetics, Louisiana State University Health Sciences Center, New Orleans, LA, USA; <sup>2</sup>Department of Microbiology, Immunology, and Parasitology, Louisiana State University Health Sciences Center, New Orleans, LA, USA and <sup>3</sup>Department of Pediatrics and Stanley S. Scott Cancer Center, Louisiana State University Health Sciences Center, New Orleans, LA, USA. Correspondence: Dr AD Hollenbach, Department of Genetics, Louisiana State University Health Sciences Center, 533 Bolivar Street, CSRB 6th floor, New Orleans, LA 70112, USA.

E-mail: aholle@lsuhsc.edu

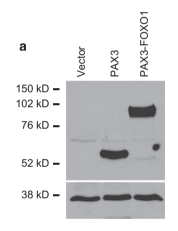
<sup>&</sup>lt;sup>4</sup>Current address: Center for Human Genetic Research, Massachusetts General Hospital, Richard B. Simches Research Center, Boston, MA 02114, USA.

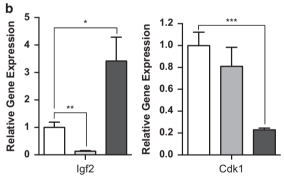
<sup>&</sup>lt;sup>5</sup>Current address: Tulane University, New Orleans, LA 70112, USA.

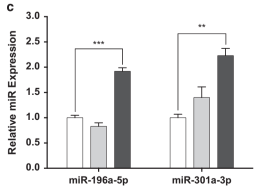
<sup>&</sup>lt;sup>6</sup>These authors contributed equally to this work.

PAX3-FOXO1 affects microRNA (miRNA) expression and how these changes contribute to the invasive capacity of myoblasts.

In this study we utilized physiologically relevant wild-type primary myoblasts along with large-scale comparative transcriptomic analyses to examine how the expression of PAX3-FOXO1 or PAX3 alters global mRNA and miRNA expression profiles and how these changes contribute to the invasive potential of these cells. We report here that the expression of the oncogenic fusion protein is sufficient to alter the expression of 70 mRNA and 27 miRNA in such a way that would be expected to promote cellular invasion. In contrast, the expression of PAX3 elicits mRNA and miRNA expression changes that would be expected to inhibit cellular invasion. We found that these mRNA and miRNA changes translate into biological effects, with the expression of PAX3-FOXO1 enhancing and the expression of PAX3 inhibiting primary myoblast invasion. Taken together, these results provide a more expansive picture to describe the increased localized invasion seen with t(2;13)(q35;q14) positive ARMS tumors, and describes how the presence of PAX3-FOXO1 may contribute to higher levels of metastasis in these patients.







#### RESULTS AND DISCUSSION

To understand how PAX3-FOXO1 affects global mRNA and miRNA expression, we stably transduced passage-matched wild-type mouse primary myoblasts with the MSCV-IRES-puromycin retroviral vector (negative control), or the same retroviral vector expressing FLAG epitope-tagged PAX3 (FLAG-PAX3) or FLAG-PAX3-FOXO1, a tag previously shown to not affect Pax3 or Pax3-FOXO1 function.<sup>6,17</sup> The puromycin selected cells were harvested from three independent transductions and pooled, resulting in a single mixed population for each individual construct, which removes the potential for variability that may occur from clonal effects. The level of PAX3-FOXO1 expression is equivalent to the level of expression of the fusion protein in ARMS tumor cell lines (Figure 1 and Dietz *et al.*<sup>18,19</sup>) and is therefore directly relevant to the role of the oncogenic fusion protein in ARMS. This model allows us to use a physiologically relevant cell system in the absence of any complimentary transforming mutations to determine the specific effects of PAX3-FOXO1 on oncogenic phenotypes.

We performed mRNA and miRNA deep-sequencing analyses on total RNA isolated from three independent growths of stably transduced cells and utilized the resulting data to perform comparative transcriptomic analyses to understand how each protein alters expression profiles to exert their effects on the invasive capacity of cells. For both the mRNA and miRNA analyses the data used for subsequent studies were limited to (1) those genes or miRNA displaying statistically significant differences (*P* < 0.05, as determined by the Galaxy Cuffdiff program (mRNA) or miRNAKey (miRNA)), (2) transcripts whose expression was present in both data sets being analyzed to rule out potential artifactual differences resulting from depth of read and (3) transcripts or miRNA that exhibited at least 2-fold differences in expression either up or downregulated.

We found a total of 480 mRNA whose expression changed in a PAX3-FOXO1-dependent manner (276 downregulated and 204 upregulated) relative to the empty vector negative control (data

Figure 1. Protein expression (a) and quantitative RT-PCR analyses for (b) select mRNA and (c) select miRNA. Mouse primary myoblasts were isolated from 2- to 4-day-old C57/Bl6 mice as previously described.<sup>53</sup> Cells were grown as previously described<sup>7,17-19,53</sup> and were passage-matched to prevent possible differences due to passage conditions. Mouse primary myoblasts were stably transduced as previously described<sup>6,53</sup> with the MSCV-IRES-puromycin empty vector, vector containing FLAG epitope-tagged Pax3 (FLAG-Pax3) or FLAG-PAX3-FOXO1. Three days post transduction, cells were selected using puromycin, as previously described. 19 transduced cells were harvested and pooled from three independent transductions to create a single population that express each construct. (a) Total cell extracts made, as previously described. 17-19,53 Equal amounts of total cell lysates (12 µg) were separated by 8% SDS-PAGE and analyzed by western blot analysis using antibodies specific for Pax3,<sup>54</sup> as previously described. <sup>18,19</sup> (**b,c**) Total RNA was isolated from the stably transduced proliferating primary myoblasts (empty vector (white bars), PAX3 (gray bars) or PAX3-FOXO1 (black bars)) using the miRNeasy mini kit (Qiagen, Madison, WI, USA), allowing for the isolation of RNA < 30 bp in length, according to the manufacturer's specifications. Equal amounts of total RNA (100 ng) were used for cDNA synthesis using the iScript cDNA synthesis kit (Bio-Rad, Hercules, CA, USA) for mRNA (b) or the Tagman miRNA reverse transcription kit (Applied Biosystems, Foster City, CA, USA) for miRNA. (c) The qRT-PCR was performed on the resulting cDNA using the CFX96 Touch Real-Time PCR Detection System (Bio-Rad) using commercially available primer/probe sets and the Applied Biosystems Universal Master Mix (Applied Biosystems), according to the manufacturer's specifications. All results were normalized for GAPDH (mRNA) or the U6 small nuclear RNA (miRNA) and reported as fold expression relative to the results obtained for cells stably transduced with the empty vector. In all cases, analyses were performed comparing each sample with the empty vector control (\*P = 0.009, \*\*P = 0.001, \*\*\*P = 0.0001).

not shown). We performed a PubMed search on each of the 480 mRNA, using the gene name and the search term 'invasion' to determine if they were experimentally proven to contribute to cellular invasion. We found that 70 of the 480 altered genes (14.5%) are involved in regulating the invasive capacity of cells (Table 1). Forty-three of these genes have literature evidence demonstrating their role in promoting cellular invasion, with these altered genes being split nearly equally between being uprequlated (19/43; 44%) or downregulated (24/43; 56%) in a PAX3-FOXO1-dependent manner. In a similar manner, 27 genes have literature evidence to support their role in inhibiting cellular invasion, with 21 of these genes (nearly 80%) being downregulated in a PAX3-FOXO1-dependent manner. Finally, 17 of the 70 differentially expressed genes (nearly 25%) contain PAX3-FOXO1 binding sites in their proximal promoters, as previously described<sup>20</sup> (Table 1, c), four of these genes were previously demonstrated to be regulated by PAX3-FOXO1, including cannabinoid receptor 1, <sup>16</sup> FGFR4, <sup>20</sup> IGF2<sup>21</sup> and IGFBP5<sup>15</sup> (Table 1, b), and 21 of the 70 (30%) genes have altered gene expression levels consistent with changes seen in human tumor samples<sup>22–25</sup> (Table 1, a).

An initial examination of the distribution of mRNA whose levels are altered upon the expression of the fusion protein would suggest that PAX3-FOXO1 would primarily exert its invasive effect<sup>13</sup> by decreasing the expression of genes important for inhibiting invasion. However, it is interesting to note that although only 44% of the genes that promote invasion are upregulated, nearly half of these 19 upregulated genes (8/19-42%) are increased > 6-fold, including the previously reported cannabinoid receptor 16 (cannabinoid receptor 1-6.92-fold), with the top four genes being upregulated > 20-fold. Therefore, this data suggest that PAX3-FOXO1 exerts its effects on invasive capacity by not only decreasing the expression of a large number of inhibitory genes, but by simultaneously greatly increasing the expression of key genes that promote invasion, including genes that encode for proteins involved in cytoskeletal organization (CAP6-33.45-fold), cadherins (CDH6-23.23-fold), extracellular matrix metalloproteases (ADAMTS1-21.98-fold) and cell adhesion proteins (MSLN - 20.93fold).

A similar analysis found 399 mRNA change in a PAX3dependent manner (276 downregulated and 123 upregulated) relative to the empty vector negative control (data not shown). A similar PubMed search revealed that 60 of the 399 genes (15%) are involved with regulating invasion (Table 1). Thirty-eight of these genes have a demonstrated role in promoting invasion, with a majority of these genes (25/38; 66%) being downregulated. Further, 22 mRNA were demonstrated to inhibit invasion, with 6 of these genes being upregulated and 16 being downregulated. Finally, four of the differentially expressed genes were demonstrated in the literature to be directly regulated by PAX3, including Ahr,<sup>26</sup> IGF1R,<sup>20</sup> EPHA2<sup>27</sup> and MET<sup>28</sup> (Table 1, b). Although a smaller number of these inhibitory genes are upregulated, one of them is upregulated >15-fold (metallopeptidase Mme—15.17-fold). In contrast to the results seen with PAX3-FOXO1, these data suggest that PAX3 would be expected to inhibit invasive capacity, primarily through the downregulation of genes that promote this biological event.

A comparative transcriptomic analysis of the miRNA data identified a total of 84 miRNAs whose expression changed in a PAX3-FOXO1-dependent manner (46 downregulated and 38 upregulated) relative to the empty vector negative control (data not shown). A PubMed search of each of the individual 84 miRNA, using the miRNA name and the search term 'invasion', indicated that 10 of these miRNA promote cellular invasion (Table 2). Of these miRNA, 9/10 (90%) have an increased PAX3-FOXO1-dependent expression with the top two being increased > 20-fold. In a similar manner, 17 miRNA are important for inhibiting cellular invasion, of which 16/17 (94%) are

Table 1. Altered mRNA expression important for tumor cell invasion					
	· · · · · · · · · · · · · · · · · · ·				
Gene	Gene function	V vs PF	V vs P3		
mRNA that promote tumor invasion					
Capn6	Cytoskeletal organization	33.45			
Cdh6	Type II cadherin, development	23.23	9.17		
<sup>a</sup> Adamts1	Metalloprotease	21.98	5.65		
MsIn	Cell adhesion; overexpressed	20.93	48.60		
<sup>a</sup> Adamts5	in cancers rs5 Peptidase; aggrecanase to cleave				
<sup>a,b</sup> Cnr1	aggrecan	6.02			
Hoxb9	G-protein signaling Transcriptional activator increased in cancers	6.92 6.34			
a,b,cEafr1	FGF receptor	6.12			
<sup>a,b,c</sup> Fgfr4 <sup>a,b</sup> Igf2	Growth factor	5.37	- 13.18		
cPlxna2	Semaphorin co-receptors	3.32	13.10		
Erbb3	EGFR receptor tyrosine kinases	3.10	-4.04		
Klf5	Possible transcription factor	2.84	-4.04		
Pbx3					
	Transcriptional activator	2.59			
Stat3	Expression of genes in response to cell stimuli	2.58			
<sup>c</sup> Sulf2 <sup>c</sup> Lamc1	Remove 6-O-sulfate groups from heparan sulfate Mediate attachment, migration-	2.52			
	interacting extracellular matrix	2.43			
Pkp2	Linking cadherins to intermediate filaments  Activates the cyclin dependent kinese				
Cdc25b	Activates the cyclin-dependent kinase CDC2	2.20	2.05		
<sup>a</sup> Ncam1	Cell adhesion; cell-to-cell interactions	2.00	- 2.95		
Tpm3	Provide stability to actin filaments	- 2.08			
Fscn1	Cell migration, motility and adhesion	- 2.09			
Mdm2	E3 ubiquitin-protein ligase	- 2.12			
<sup>a</sup> Adam19	Matrix metalloproteinase	-2.15	-3.08		
<sup>a</sup> Jag2	Ligand that activates Notch	-2.17	<b>- 4.05</b>		
Cttn	Adherins and cytoskeleton	-2.23			
Arpc5 <sup>a,c</sup> Abi1	Control of actin polymerization Mediates signal transduction from	- 2.34 - 2.42			
<sup>a</sup> Jak1	Ras to Rac Cell signal transduction	- 2.45			
Mmp14	Metalloproteinase	- 2.58			
Ifitm1	Implicated in cell adhesion	- 2.68	- 2.88		
aElk3	Activated by signal-induced	- 2.74	2.00		
LING	phosphorylation	2.7 ¬			
Vim	Cytoskeletal protein	- 2.90			
ld1	Inhibits the DNA-binding transcription	- 3.00			
IUI	factors	- 3.00			
Pak1		-3.21			
<sup>a</sup> Cyr61	Cell motility and morphology Promotes the adhesion of endothelial	- 3.21 - 3.28			
Cyror	cells	- 3.20			
<sup>a</sup> Dusp1	Cellular response to environmental stress	-3.62			
<sup>a</sup> Lasp1	Binds to the actin cytoskeleton	- 3.91			
Ntn4	Protein related to laminins	- 4.93			
Etv4	Transcriptional activator	- 5.12			
<sup>a,c</sup> Flnb	Filamin; repair vascular injuries	- 5.82			
AxI	Transduces signals from the	-6.04			
7120	extracellular matrix	0.01			
<sup>a</sup> lgfbp2	Inhibits IGF-mediated growth	- 8.64			
Cxcl12	Chemotaxis; embryonic development		- 14.90		
b Ahr		- 19.00	11.13		
	Ligand-activated transcriptional activator				
Egfr	Receptor for members of EGF family		4.82		
Eps8	Functions as part of the EGFR pathway		3.59		
Sema3e	Axon guidance; Semaphorins		3.43		
Galnt2	Oligosaccharide biosynthesis		2.64		
Sparc	Involved in ECM synthesis		2.44		
Ghr	Transmembrane receptor for growth		2.28		
0.4.5	hormone				
Prdx1	Antioxidant protective		2.27		
Emp3	Involved in proliferation and cell-cell		2.04		
	interactions				
Rnf11	Transcriptional activator		- 2.01		
Муо5а	Cytoplasmic vesicle transport and		- 2.09		
	anchorage				

Gene         Gene function         V vs PF         V vs PS           bc/gftr         Critical role in transformation events         -2.13           Hes6         Promotes cell differentiation         -2.17           Abl2         Non-receptor tyrosine protein kinases         -2.21           Peak1         Role in cell spreading and migration         -2.24           Zkscan3         Transcriptional regulator         -2.35           Tppp3         Tubulin and has microtubule-bundling activity         -2.25           Si3gal1         Transcription factor         -2.50           Ephna         Ephrin receptor subfamily         -2.53           Notch1         Developmental processes by controlling cell fact         -3.07           Cell Fabra         Ephrin receptor subfamily         -2.50           Vesproba         Histone demethylase; transcriptional correspensor         -3.07           Cell Surface receptor for cell-cell signaling         -3.46           Multifunctional kinase         -3.37           Nuak1         Multifunctional kinase         -3.37           Seppine         Multifunctional kinase         -3.73           Seppine         Alter the interaction of IGFs with receptor         -3.73           Seppine         Alter the interaction of IGFs with receptor <th>Table 1. (C</th> <th>Continued)</th> <th></th> <th></th>	Table 1. (C	Continued)		
Abs/Ea         Promotes cell differentiation         — 2.17           AblZ         Non-receptor tyrosine protein kinases         — 2.21           Peak1         Role in cell spreading and migration         — 2.24           Zkscan3         Transcriptional regulator         — 2.35           Tppp3         Tubulin and has microtubule-bundling activity         — 2.47           St3gal1         Transcription factor         — 2.50           Behal         Ephrin receptor subfamily         — 2.53           Notch1         Developmental processes by controlling cell fate         — 3.12           Jun         Transcriptional activator         — 3.12           Kdm5b         Hostone demethylase; transcriptional corepressor         — 3.46           Semana         Cell surface receptor for cell-cell signaling         — 3.46           bc-Met         Hepatocyte growth factor receptor         — 3.73           Romana         Multifunctional kinase         — 3.49           Serpine2         Alter the interaction of IGFs with receptor         8.28         — 5.68           Spy1         Alter the interaction of IGFs with receptor         8.28         — 5.68           Spy1         Alter the interaction of IGFs with receptor         8.28         — 5.68           Spy1         Antagonist of FGF	Gene	Gene function	V vs PF	V vs P3
Abi2         Non-receptor tyrosine protein kinases         -2.21           Peak1         Role in cell spreading and migration         -2.24           Zkscan3         Transcriptional regulator         -2.35           Tapp3         Tubulin and has microtubule-bundling activity         -2.47           Stagal1         Transcription factor         -2.50           Bepha2         Ephrin receptor subfamily         -2.53           Notch1         Developmental processes by controlling cell fate         -3.07           Jun         Transcriptional activator         -3.17           Kdm5b         Histone demethylase; transcriptional corepressor         -3.17           Cell surface receptor for cell-cell         -3.43           Nuah1         Multifunctional kinase         -3.87           MRNA that inhibit tumor invasion         -4.99           MBRNA that inhibit tumor invasion         8.28         -5.68           Serpinb1         Alter the interaction of IGFs with receptor         8.28         -5.68           CSpry1         Antagonist of FGF pathways         7.39         7.39           Spry2 inhibitor of angiogenesis and tumor growth         2.62         -2.37           Col4a2         Metastasis suppressor         2.62         -2.23           Spry2				
Peak1         Role in cell spreading and migration         -2.24           Zkscan3         Transcriptional regulator         -2.35           Tppp3         Tubulin and has microtubule-bundling activity         -2.47           St3gal1         Transcription factor         -2.50           Pepha2         Ephrin receptor subfamily         -2.53           Notch1         Developmental processes by controlling cell fate         -3.07           Jun         Transcriptional activator         -3.12           Kdm5b         Histone demethylase; transcriptional corepressor         -3.17           Corepressor         Cell surface receptor for cell-cell signaling         -3.46           bcrMet         Hepatocyte growth factor receptor         -3.33           MRNA that inhibit tumor invasion         Alter the interaction of IGFs with receptor         -3.87           Serpine1         Alter the interaction of IGFs with receptor         6.48           *Spry1         Antagonist of FGF pathways         7.39           *Proteinase inhibitor         6.48           *Pox         Bind microtubules         5.16           *Col4a2         Inhibitor of angiogenesis and tumor growth         2.62         -2.37           *Spry2         Inhibitor of angiogenesis and tumor growth         -2.20         -2.21 <td></td> <td></td> <td></td> <td></td>				
Zkscan3         Transcriptional regulator         − 2.35           Tppp3         Transfer of sialic acid to substrates         − 2.45           Bach1         Transcription factor         − 2.50           Pspha2         Ephrin receptor subfamily         − 2.53           Notch1         Developmental processes by controlling cell fate         − 3.17           Jun         Transcriptional activator         − 3.17           Kdm5b         Histone demethylase; transcriptional corepressor         − 3.17           **Sema6a         Cell surface receptor for cell-cell         − 3.46           **Sema6a         Cell surface receptor for cell-cell         − 3.47           **Nuak1         Multifunctional kinase         − 3.87           **Bind Multifunctional kinase         − 3.87           **Bind Multifunctional kinase         − 3.87           **Bind milibit tumor invasion         8.28         − 5.68           **Serpinal         Alter the interaction of IGFs with ecceptor         8.28         − 5.68           **Serpinal         Alter the interaction of IGFs with ecceptor         8.28         − 5.68           **Serpinal         **Nome inhibitor         6.48         8           **Serpinal         **Nome inhibitor         6.48         8           **Serpin				
Toppp3         Tubulin and has microtubule-bundling activity         -2.45           St3gal1         Transfer of sialic acid to substrates         -2.47           PEpha2         Ephrin receptor subfamily         -2.53           Notch1         Developmental processes by controlling cell fate         -3.07           Cell fate         -3.07           Sema6a         Transcriptional activator         -3.12           Kdm5b         Histone demethylase; transcriptional corepressor         -3.17           Cema6a         Cell surface receptor for cell-cell signaling         -3.46           signaling         Hepatocyte growth factor receptor         -3.73           Nuak1         Hepatocyte growth factor receptor inhibits resine proteases         -3.87           MRNA transcriptional kinase         -3.88           Inhibits transcriptional kinase         -3.89           Porpor         Antagonist of FGF pathways         7.39           Serpinb1         Antagonist of FGF pathways         7.39           Porpor         Antagonist of FGF pathways         7.39           Proteinase inhibitor         6.48           Bind microtubules         5.16           Collaz         Metastasis suppressor         2.62         -2.37           Spry2         Inhibitor of angiog				
St3gal1         activity           St3gal1         Transcription factor         - 2.50           Epha2         Ephrin receptor subfamily         - 2.50           Notch1         Developmental processes by controlling cell fate         - 3.07           Jun         Transcriptional activator         - 3.12           Kdm5b         Histone demethylase; transcriptional corepressor         - 3.17           Sema6a         Cell surface receptor for cell-cell signalling         - 3.46           bc.Met         Hepatocyte growth factor receptor         - 3.38           Serpine2         Inhibit serine proteases         - 4.99           mRNA taxt         Inhibit tumor invasion         8.28         - 5.68           Serpinb1         Alter the interaction of IGFs with receptor         8.28         - 5.68           Serpinb1         Proteinase inhibitor         6.48         - 5.68           Serpinb1         Inhibitor of angiogenesis and tumor growth         2.02         - 2.37           Colda2         Inhibitor of angiogenesis and tumor growth         2.62         - 2.37           Spry2         Inhibitor of angiogenesis and tumor growth         2.62         - 2.37           Spry2         Inhibitor of angiogenesis and tumor growth actor growth in growth         2.26         - 4.02     <				
Bach1 Transcription factor Ephrin receptor subfamily Notch1 Developmental processes by controlling cell fate  Jun Transcriptional activator Kdm5b Histone demethylase; transcriptional corepressor CSema6a Cell surface receptor for cell-cell signaling  bc.Met Hepatocyte growth factor receptor Nuak1 Multifunctional kinase 3.87 Nuak1 Multifunctional kinase 3.87 Serpine2 Inhibit tumor invasion Ablgfbp5 Alter the interaction of IGFs with receptor Sprp1 Antagonist of FGF pathways 3.9 Serpinb1 Proteinase inhibitor 6.48 Pack Bind microtubules 5.16 Inhibitor of angiogenesis and tumor growth 2.83 Spry2 Inhibitory effect on growth factor 2.10 Spry2 Inhibitory effect on growth factor 2.10 Spry2 Inhibitory effect on growth factor 2.10 signaling Ccited2 Inhibits transactivation of HIF1A 2.26 -4.02 induced genes Nonmuscle, cytoskeletal, alpha actinin 2.32 Inhibitors of the matrix 2.68 Remodeling the cytoskeleton 2.43 Lpp Involved in cell-cell adhesion and cell 2.62  "Timp2 Inhibitors of the matrix 2.68 Transmission of signals to the 2.68 "Timp2 Inhibitors of the matrix 2.88 Cdk1 Cell cycle regulatory kinase 3.01 Creb311 Transcriptional activator 2.88 Inhibits the DNA-binding transcription 3.13 Filip11 Regulator of the anti-angiogenic activity 5.68 2.21 Sox4 Regulation of embryonic development 4.18 4.30 Neg3 Inhibits signal transduction 3.78 4.93 Filip11 Modulate action of growth factors 8.56 2.21 Sox4 Regulation of embryonic development 4.18 4.30 Neg3 Inhibits rignal transduction 3.78 4.93 Filip11 Modulate action of growth factors 8.56 2.21 Sox6 Regulation of embryonic development 4.18 4.30 Neg3 Inhibits signal transduction 3.78 4.93 Filip11 Modulate action of growth factors 8.56 2.21 Sox6 Regulation of embryonic development 6.48 2.91 Scaffold protein in signal transduction 3.78 4.93 Filip11 Modulate action of growth factors 8.56 2.21 Sox6 Re	'''	activity		
Pepha2   Ephrin receptor subfamily   Automatical Developmental processes by controlling   Cell fate				
Developmental processes by controlling cell fate				
Kdm5b Histone demethylase; transcriptional corepressor  **Csema6a** Cell surface receptor for cell-cell signaling  **Br.Met** Nuak1** Nuak1** Multifunctional kinase		Developmental processes by controlling		- 3.07
Corepressor Cell surface receptor for cell-cell signaling b.CMet Hepatocyte growth factor receptor Nuak1 Multifunctional kinase Serpine2 Inhibits erine proteases - 3.87 Serpine2 Inhibits erine proteases - 4.99  mRNA that inhibit tumor invasion abligfbp5 Alter the interaction of IGFs with receptor  Spp1 Antagonist of FGF pathways 7.39 Serpinb1 Proteinase inhibitor 6.48 Bind microtubules 5.16 Col4a2 Inhibitor of angiogenesis and tumor 2.83 growth 2.82 Col4a2 Metastasis suppressor 2.62 - 2.37 Spry2 Inhibitory effect on growth factor - 2.10 signaling Deptor Negative regulator of the mTORC1 - 2.12 signaling CfCited2 Inhibits transactivation of HIF1A 2.26 - 4.02 induced genes Nonmuscle, cytoskeletal, alpha actinin - 2.32 isoform FIna Remodeling the cytoskeleton - 2.43 Lpp Involved in cell-cell adhesion and cell - 2.62 motility  CDIg5 Transmission of signals to the cytoskeleton   -2.68 wotility  Colf31 Transcriptional activator - 2.88 Tagh Actin crosslinking/gelling protein - 2.70 App Transcriptional activator - 3.04 Creb311 Transcriptional activator - 3.04 Phosphatase; negatively regulate (MAP) - 3.08 kinases  ald3 Inhibits tipe DNA-binding transcription - 3.13 factors  Wisp1 Downstream in the WNT1 signaling - 3.32 - 2.36 Sox4 Regulation of the anti-angiogenic activity - 5.68 Colf inhibits signal transduction - 7.98 Asp 16 Inhibits signal transduction - 7.98 Ags16 Inhibits signal transduction - 7.98 Ags16 Inhibits de DNA-binding transcription - 7.98 Ags16 Inhibits de DNA-binding transcription - 7.98 Ags16 Inhibits alignal transduction - 7.98 Ags16 Inhibits de DNA-binding transduction - 7.98 Ags16 Inhibits de DNA-binding transduction - 7.98 Ags16 Inhibits alignal transduction - 7.98 Ags16 Inhibits alignal transduction - 7.98 Ags16 Inhibits de DNA-binding transduction - 7.98 Ags16 Inhibits de DNA-binding transduction - 7.98 Ags17 Algentation of embryonic development - 6.48 Actin - 7.2.15 Adam9 Biological processes: cell-cell/matrix interactions Filip1 Repulator of the anti-angiogenic activity - 5.68 Actin - 7.2.15 A	Jun	Transcriptional activator		- 3.12
Cell surface receptor for cell-cell signaling   Signaling	Kdm5b			- 3.17
signaling Nuak1 Multifunctional kinase	SC			2.46
Nuck1 Serpine2 Inhibit serine proteases — 3.87 — 4.99  mRNA that inhibit tumor invasion  a-blgfbp5 Alter the interaction of IGFs with receptor  CSpy1 Antagonist of FGF pathways 7.39  Serpinb1 Proteinase inhibitor 6.48  Bind microtubules 5.16  Col4a2 Inhibitor of angiogenesis and tumor growth 1.00  GM2 Metastasis suppressor 2.62 — 2.37  Spy2 Inhibitory effect on growth factor signaling  Deptor Negative regulator of the mTORC1 — 2.12  signaling — 2.26 — 4.02  FCited2 Inhibits transactivation of HIF1A-induced genes  Actn1 Nonmuscle, cytoskeletal, alpha actinin isoform  Flna Remodeling the cytoskeleton — 2.43  Lpp Involved in cell-cell adhesion and cell motility  Col4b2 Transmission of signals to the cytoskeleton — 2.66  aTimp2 Inhibitors of the matrix — 2.68  metalloproteinases  Tagln Actin crosslinking/gelling protein — 2.70 — 3.56  Tapp Transcriptional activator — 2.88  Cell cycle regulatory kinase — 3.01  Transcriptional activator — 3.04  Phosphatase; negatively regulate (MAP) kinases  ald3 Inhibits the DNA-binding transcription — 3.13  factors  Wisp1 Downstream in the WNT1 signaling — 3.32 — 2.36  pathway  Rgs16 Elimigration and bone development — 4.18 — 4.30  Filip 11 Regulator of the anti-angiogenic activity — 5.68 — 2.21  Sox4 Regulation of embryonic development — 4.18 — 4.30  Filip 11 Regulator of the anti-angiogenic activity — 5.68 — 2.21  Sox4 Regulation of embryonic development — 6.48 — 2.91  "Akap12 Scaffold protein in signal transduction — 7.98 — 3.87  Fst11 Modulate action of growth factors — 8.56 — 2.16  "Mme Metallopeptidase — 2.93  Dusp6 Negatively regulate (MAP) kinases — 2.93  Biological processes: cell-cell/matrix interactions  Flnc Crosslink actin filaments — 2.58		signaling		- 3.46
Serpine2Inhibit serine proteases- 4.99mRNA that ablgfbp5Alter the interaction of IGFs with receptor8.28- 5.68CSpry1 Serpinb1Proteinase inhibitor and microtubules Col/a2 Inhibitor of angiogenesis and tumor growth7.39- 5.16 CA82 Metastasis suppressor Inhibitory effect on growth factor signaling2.62 - 2.37- 2.32Deptor Spry2 Inhibitory Inhibitory effect on growth factor signaling- 2.12 - 2.10 signaling- 2.26 - 4.02**Cited2 Inhibits transactivation of HIF1A- induced genes induced genes motility- 2.26 - 4.02**Fina Involved in cell-cell adhesion and cell motility- 2.66 - 2.62 - 2.62 motility**Dips **Timp2 **Inhibitors of the matrix metalloproteinases- 2.68 - 2.68 - 2.68**Tagln **CAb1 **Creb311 **Dusp4 **Phosphatase; negatively regulate (MAP) **Inhibits the DNA-binding transcription factors- 3.04 - 3.08 **Inhibits the DNA-binding transcription factors- 3.13 - 3.32 - 2.36**Wisp1 **Sp31 **Sp361 **Inhibits signal transduction **Filip11 **Soarfold protein in signal transduction **Fist11 **Modulate action of embryonic development **Fist11 **Modulate action of growth factors- 3.78 - 4.93 - 4.18 - 4.30 - 2.15 - 2.15 - 2.26 - 3.31 - 3.31 - 3.32 - 3.32 - 3.36 - 3.31 - 3.31 - 3.31 - 3.32 - 3.36 - 3.31 - 3.31 - 3.31 - 3.32 - 3.32 - 3.33 - 3.33 - 3.31 - 3.34 - 3.34 - 3.35 - 3.36 - 3.37 - 3.38 - 3.37 - 3.38 - 3.39 - 3.39 - 3.30 - 3.30 - 3.30 -				
mRNA that inhibit tumor invasion  **ablgfbp5** Alter the interaction of IGFs with receptor  **Spry1** Antagonist of FGF pathways 7.39  **Serpinb1** Proteinase inhibitor 6.48  **Pocx** Bind microtubules 5.16  **Col4a2** Inhibitor of angiogenesis and tumor growth 1.50  **Gab2** Metastasis suppressor 2.62 -2.37  **Spry2** Inhibitory effect on growth factor signalling 1.50  **Deptor** Negative regulator of the mTORC1 5.10  **Signalling 1.50  **Cited2** Inhibits transactivation of HIF1A-1.50  **induced genes 1.50  **Actn1** Nonmuscle, cytoskeletal, alpha actinin isoform 1.50  **Fina Remodeling the cytoskeleton 1.50  **Inna Remodeling the cytoskeleto	ı			
a-blgfbp5 Alter the interaction of IGFs with receptor    **Spry1** Antagonist of FGF pathways   **Proteinase inhibitor   **Dox   **Bind microtubules   **Col422** Inhibitor of angiogenesis and tumor growth    **Cd82** Metastasis suppressor   **Spry2** Inhibitory effect on growth factor signalling    **Deptor Negative regulator of the mTORC1   **signalling    **Deptor Negative regulator of the mTORC1   **signalling    **Signalling    **Deptor Negative regulator of the mTORC1   **signalling    **Signalling    **Deptor Nonmuscle, cytoskeletal, alpha actinin isoform    **Flna Remodeling the cytoskeleton   **Inna Nonmuscle, cytoskeleton   **Inna Nonmuscle, cytoskeleton   **Inp   **Involved in cell-cell adhesion and cell inhibitors of the matrix metalloproteinases    **Tagln Actin crosslinking/gelling protein   **Papp    **Transcriptional activator   **Cell cycle regulatory kinase   **Calc   **Cell cycle regulatory kinase   **Calc   **Cell cycle regulatory kinase   **Alda    **Inhibits the DNA-binding transcription   **factors    **Wisp1    **Downstream in the WNT1 signalling   **pathway    **Rgs16   **Inhibits the DNA-binding transcription   **factors    **John Signal transduction   **Signal transduction   **John Signal transd	Serpine2	innibit serine proteases		- 4.99
receptor Seprinb1 Serpinb1 Proteinase inhibitor Serpinb1 Proteinase inhibitor Col4a2 Inhibitor of angiogenesis and tumor growth  Cd82 Metastasis suppressor Spry2 Inhibitory effect on growth factor signaling Deptor Negative regulator of the mTORC1 signaling  Cited2 Inhibits transactivation of HIF1A- induced genes  Actn1 Nonmuscle, cytoskeletal, alpha actinin isoform  Flna Remodeling the cytoskeleton Inhibitors of signals to the cytoskeleton  aTimp2 Inhibitors of the matrix metalloproteinases  Tagln Actin crosslinking/gelling protein Cell cycle regulatory kinase Cell cycle regulatory kinase Inhibits the DNA-binding transcription Factors  Wisp1 Downstream in the WNT1 signaling pathway Inhibits signal transduction Sovat Regulator of the anti-angiogenic activity Sox4 Regulation of embryonic development Filip Metallopeptidase Ciprosa Calcular calcular growth and differentiation Nefl Intracellular transport to axons and dendrites Dusp6 Regator Sorus Regulator processes: cell-cell/matrix interactions Finc Crosslink actin filaments Cell adhesion and growth factor Coll adhesion and growth factor Coll adhesion and growth factor Coll adhesion and growth factor Crosslink actin filaments Coll adhesion and growth factor Coll adhesion and gro	mRNA tha			
Spry1       Antagonist of FGF pathways       7.39         Sepinb1       Proteinase inhibitor       6.48         ³Dcx       Bind microtubules       5.16         Col4a2       Inhibitor of angiogenesis and tumor growth       2.83         Col4a2       Inhibitory effect on growth factor signaling       2.62       − 2.37         Spry2       Inhibitory effect on growth factor signaling       − 2.10       − 2.10       − 2.10         Signaling       Negative regulator of the mTORC1       − 2.12       − 2.12       − 2.26       − 4.02         Actn1       Negative regulator of the mTORC1       − 2.26       − 4.02         induced genes       Actn1       Nonmuscle, cytoskeletal, alpha actinin       − 2.26       − 4.02         Inhibit mack genes       Nonmuscle, cytoskeleton       − 2.43       − 2.62       − 2.62         Inpublic motility       Transcriptional of signals to the cytoskeleton in signals to the cytoskeleton       − 2.66       − 2.66       − 2.66       − 2.66       − 2.66       − 2.66       − 2.66       − 2.66       − 2.66       − 2.66       − 2.66       − 2.68       − 2.66       − 2.68       − 2.66       − 2.68       − 2.68       − 2.66       − 2.70       − 3.56       − 2.70       − 3.56       − 2.88       − 2.66       − 2	<sup>a,b</sup> lgfbp5		8.28	- 5.68
Serpinb1Proteinase inhibitor6.48aDCXBind microtubules5.16Col4a2Inhibitor of angiogenesis and tumor growth2.83Spry2Inhibitory effect on growth factor signalling2.62- 2.37DeptorNegative regulator of the mTORC1 signaling- 2.12- 2.12beta Cited2Inhibits transactivation of HIF1A- induced genes- 2.26- 4.02Actn1Inhibits transactivation of HIF1A- induced genes- 2.32- 2.32Actn1Remodeling the cytoskeleton motility- 2.43- 2.62Involved in cell-cell adhesion and cell cytoskeleton- 2.62- 2.62motilityTransmission of signals to the cytoskeleton- 2.66- 2.66a Timp2Actin crosslinking/gelling protein- 2.66- 2.56CApp CApp Transcriptional activator ranscriptional activator Phosphatase; negatively regulate (MAP) kinases- 3.01- 3.56Inhibits the DNA-binding transcription Inhibits signal transduction pathway- 3.32- 2.36Pilip11 SOX4 SOX4 Regulation of the anti-angiogenic activity Scaffold protein in signal transduction Pathaga Capptosa Development, cellular growth and differentiation- 3.78 Hodulate action of growth factors- 8.56 Hodulate action of growth factors- 8.56 Hodulate action of growth factors- 8.56 Hodulate action of growth and differentiation- 3.79 Hodulate action of growth and differentiation- 3.79 Hodulate action of growth factors- 8.56 Hodulate action of growth factors- 2.15 Hodulate actio	cSprv1		7.39	
a Dcx Col4a2Bind microtubules Inhibitor of angiogenesis and tumor growth5.16 2.83 2.83 3.83 3.83 3.84 3.84 3.87 3.87 3.84 3.85 3.85 3.86 3.86 3.86 3.87 3.86 3.86 3.87 3.87 3.86 3.87 3.				
Gd82 Metastasis suppressor 2.62 -2.37 Spry2 Inhibitory effect on growth factor 5pry2 signaling Deptor Negative regulator of the mTORC1 -2.12 signaling CCited2 Inhibits transactivation of HIF1A-induced genes Actn1 Nonmuscle, cytoskeletal, alpha actinin isoform Flna Remodeling the cytoskeleton -2.43 Lpp Involved in cell-cell adhesion and cell cytoskeleton Transmission of signals to the cytoskeleton cytoskeleton Transmission of signals to the cytoskeleton Transmission of signals to the cytoskeleton Actin crosslinking/gelling protein -2.68 Tagln Actin crosslinking/gelling protein -2.70 -3.56 CAPP Transcriptional activator -2.88 Cdk1 Cell cycle regulatory kinase -3.01 Creb311 Transcriptional activator -3.04 Dusp4 Phosphatase; negatively regulate (MAP) -3.08 kinases Ald3 Inhibits the DNA-binding transcription -3.13 factors Wisp1 Downstream in the WNT1 signaling -3.32 -2.36 pathway Rgs16 Inhibits signal transduction -3.78 -4.93 Cell migration and bone development -4.18 -4.30 Filip11 Regulator of the anti-angiogenic activity -5.68 -2.21 Sox4 Regulation of embryonic development -6.48 -2.91 Intracellular transport to axons and dendifferentiation Nefl Intracellular transport to axons and dendifferentiation Nefl Intracellular transport to axons and dendifferentiation Nefl Colladesion and growth factors -2.58				
Cd82 Spry2Metastasis suppressor Inhibitory effect on growth factor signaling2.62 -2.10-2.37Deptor SignalingNegative regulator of the mTORC1 signaling-2.12-4.02Cited2Inhibits transactivation of HIF1A- induced genes-2.26-4.02Actn1Nonmuscle, cytoskeletal, alpha actinin isoform-2.32-2.32FinaRemodeling the cytoskeleton motility-2.43-2.62LppInvolved in cell-cell adhesion and cell motility-2.62-2.66cytoskeleton a Transmission of signals to the cytoskeleton-2.66-2.66a Timp2Inhibitors of the matrix metalloproteinases-2.68-2.68Tagln CAb1 Cell cycle regulatory kinase Creb311 Transcriptional activator Phosphatase; negatively regulate (MAP) kinases-3.01-3.04Binhibits the DNA-binding transcription forms-3.13-4.93Rgs16 Sox4 Regulation of the anti-angiogenic activity Sox4 Regulation of embryonic development Budiate action of growth factors-3.78 -4.18 -4.30-4.93 -4.93Scaffold protein in signal transduction Fist1 Modulate action of growth factors-8.56 -2.216 -2.16-2.15 -2.15Metallopeptidase Caprc5aDevelopment, cellular growth and defirerentiation3.67 -2.15-2.15 -2.48Nefl Galn7GalNAc transferase 72.48Adam9 Biological processes: cell-cell/matrix interactions-2.15 -2.258Finc Crosslink actin filaments Cell adhesion and growth factor-2.15 -2.258	Col4a2	. 3 3	2.83	
Spry2   Inhibitory effect on growth factor signaling	6 100			
Signaling Negative regulator of the mTORC1 - 2.12 signaling  CCited2 Inhibits transactivation of HIF1A-induced genes Actn1 Nonmuscle, cytoskeletal, alpha actinin - 2.32 isoform  Flna Remodeling the cytoskeleton - 2.43 Lpp Involved in cell-cell adhesion and cell - 2.62 motility  CDIG5 Transmission of signals to the cytoskeleton  aTimp2 Inhibitors of the matrix - 2.68 metalloproteinases TagIn Actin crosslinking/gelling protein - 2.70 - 3.56  CApp Transcriptional activator - 2.88 Cdk1 Cell cycle regulatory kinase - 3.01 Creb3l1 Transcriptional activator - 3.04 Dusp4 Phosphatase; negatively regulate (MAP) - 3.08 kinases ald3 Inhibits the DNA-binding transcription - 3.13 factors  Wisp1 Downstream in the WNT1 signaling - 3.32 - 2.36 pathway Rgs16 Inhibits signal transduction - 3.78 - 4.93 CTns3 Cell migration and bone development - 4.18 - 4.30 Filip11 Regulator of the anti-angiogenic activity - 5.68 - 2.21 Sox4 Regulation of embryonic development - 6.48 - 2.91 Scaffold protein in signal transduction - 7.98 - 3.87 Fstl1 Modulate action of growth factors - 8.56 - 2.16 CMme Metallopeptidase - 8.56 - 2.16 CMme Negatively regulate (MAP) kinases - 2.93 Galnt7 GalNAc transferase 7 - 2.48  Adam9 Biological processes: cell-cell/matrix interactions Flnc Crosslink actin filaments - 2.58		Metastasis suppressor		- 2.37
DeptorNegative regulator of the mTORC1 signaling- 2.12 signalingCCited2Inhibits transactivation of HIF1A- induced genes- 2.26- 4.02Actn1Nonmuscle, cytoskeletal, alpha actinin isoform- 2.32- 2.32FlnaRemodeling the cytoskeleton Involved in cell-cell adhesion and cell cytoskeleton- 2.43LppInvolved in cell-cell adhesion and cell cytoskeleton- 2.66aTimp2Inhibitors of signals to the cytoskeleton- 2.66aTimp2Inhibitors of the matrix metalloproteinases- 2.68TaglnActin crosslinking/gelling protein call cell cycle regulatory kinase cell cycle regulatory kinase- 3.01Creb311Transcriptional activator Phosphatase; negatively regulate (MAP) kinases- 3.04ald3Inhibits the DNA-binding transcription factors- 3.13Wisp1Downstream in the WNT1 signaling pathway- 3.32- 2.36Rgs16Inhibits signal transduction cell migration and bone development and bone development cell migration and bone development and bone development cell migration of embryonic development cell migration of embryonic development and development, cellular growth and differentiation- 3.78 alist17 bodulate action of growth factors alist17 bodulate action of growth factors alist27 bevelopment, cellular growth and differentiation- 7.98 alist27 alist37 bodulate action of growth factors alist37 alist38 belopment, cellular growth and differentiation- 3.67 alist37 alist37 alist37 belopment alist38 alist37 alist38 alist37 alist38 <b< td=""><td>Spry2</td><td></td><td>-2.10</td><td></td></b<>	Spry2		-2.10	
Cited2   Inhibits transactivation of HIF1A- induced genes   Nonmuscle, cytoskeletal, alpha actinin isoform   -2.32   isoform   Flna   Remodeling the cytoskeleton   -2.43   Lpp   Involved in cell-cell adhesion and cell   -2.62   motility   -2.65   cytoskeleton   -2.66   cytoskeleton   -2.66   cytoskeleton   Inhibitors of the matrix   -2.68   metalloproteinases   -2.68   Call   Cell cycle regulatory kinase   -3.01   Creb3l1   Transcriptional activator   -3.04   Cell cycle regulatory kinase   -3.01   Creb3l1   Transcriptional activator   -3.04   Cell cycle regulatory kinase   -3.01   Creb3l1   Transcriptional activator   -3.04   Cell cycle regulatory kinase   -3.01   Creb3l1   Transcriptional activator   -3.04   Cell cycle regulatory kinase   -3.01   Creb3l1   Transcriptional activator   -3.04   Cell cycle regulatory kinase   -3.01   Creb3l1   Transcriptional activator   -3.04   Cell cycle regulatory kinase   -3.01   Creb3l1   Transcriptional activator   -3.04   Cell cycle regulator of the MVNT1 signaling   -3.32   -2.36   Cell migration and bone development   -4.18   -4.30   Filip1l   Regulator of the anti-angiogenic activity   -5.68   -2.21   Sox4   Regulation of embryonic development   -6.48   -2.91   Sox4   Regulation of embryonic development   -6.48   -2.91   Sox4   Regulation of growth factors   -8.56   -2.16   CMme   Metallopeptidase   15.17   Modulate action of growth factors   -8.56   -2.16   CMme   Metallopeptidase   15.17   Call cycle regulate (MAP) kinases   2.93   Call cycle regulate (MAP) kinases   2.93   Call cycle regulate (MAP) kinases   2.93   Call cycle regulate (MAP) kinases   2.94   Call cycle regulate (MAP) kinases   2.95   Call cycle regulate (MAP) kinases   2.94   Call cycle regulate (MAP) kinases   2.95   Call cycle regulate (MAP) kin	Deptor		- 2.12	
induced genes  Actn1 Nonmuscle, cytoskeletal, alpha actinin isoform  Remodeling the cytoskeleton  Involved in cell-cell adhesion and cell Ipp Involved in cell-cell adhesion and cell Involved in cell-cell in cell-cell Involved involved in cell-cell Involved in cell-cel	C C			
Actn1 Nonmuscle, cytoskeletal, alpha actinin isoform  Flna Remodeling the cytoskeleton — 2.43 Lpp Involved in cell-cell adhesion and cell — 2.62 motility  CDlg5 Transmission of signals to the — 2.66 cytoskeleton  Talmp2 Inhibitors of the matrix — 2.68 metalloproteinases  Tagln Actin crosslinking/gelling protein — 2.70 — 3.56  CApp Transcriptional activator — 2.88 Cdk1 Cell cycle regulatory kinase — 3.01 Creb3l1 Transcriptional activator — 3.04 Dusp4 Phosphatase; negatively regulate (MAP) — 3.08 kinases  Inhibits the DNA-binding transcription — 3.13 factors  Wisp1 Downstream in the WNT1 signaling — 3.32 — 2.36 pathway  Rgs16 Inhibits signal transduction — 3.78 — 4.93 Cell migration and bone development — 4.18 — 4.30 Filip11 Regulator of the anti-angiogenic activity — 5.68 — 2.21 Sox4 Regulation of embryonic development — 6.48 — 2.91 3Akap12 Scaffold protein in signal transduction — 7.98 — 3.87 Fstl1 Modulate action of growth factors — 8.56 — 2.16 CMme Metallopeptidase — 15.17 Caprc5a Development, cellular growth and differentiation Nefl Intracellular transport to axons and dendrites Dusp6 Negatively regulate (MAP) kinases — 2.93 Galnt7 GalNAc transferase 7 — 2.48  Adam9 Biological processes: cell-cell/matrix interactions Flnc Crosslink actin filaments — 2.15 Rhob Cell adhesion and growth factor — 2.58	*Cited2		- 2.26	<b>-4.02</b>
Fina Remodeling the cytoskeleton Lpp Involved in cell-cell adhesion and cell cytoskeleton Transmission of signals to the cytoskeleton Inhibitors of the matrix metalloproteinases Tagln Actin crosslinking/gelling protein Cell cycle regulatory kinase Cell cycle regulatory kinase Creb311 Transcriptional activator Dusp4 Phosphatase; negatively regulate (MAP) Kinases Inhibits the DNA-binding transcription Factors Wisp1 Downstream in the WNT1 signaling pathway Rgs16 Inhibits signal transduction Inhibits signal transduction Cell migration and bone development Cell migration and bone development Actin Cell migration of embryonic development Sox4 Regulation of embryonic development Actin Cell migration Metallopeptidase Cefprc5a Development, cellular growth and differentiation Nefl Intracellular transport to axons and dendrites Dusp6 Negatively regulate (MAP) kinases Galnt7 GalNAc transferase 7  Adam9 Biological processes: cell-cell/matrix interactions Finc Crosslink actin filaments Rhob Cell adhesion and growth factor	Actn1	Nonmuscle, cytoskeletal, alpha actinin	- 2.32	
Lpp Involved in cell-cell adhesion and cell motility  **CDIG5** Transmission of signals to the cytoskeleton  **Imp2* Inhibitors of the matrix metalloproteinases  **TagIn* Actin crosslinking/gelling protein	Elna		2.42	
motility  **CDIg5** Transmission of signals to the cytoskeleton  **Timp2* Inhibitors of the matrix metalloproteinases  **TagIn* Actin crosslinking/gelling protein	I			
cytoskeleton Inhibitors of the matrix metalloproteinases  TagIn Actin crosslinking/gelling protein -2.70 -3.56  CApp Transcriptional activator -2.88  Cdk1 Cell cycle regulatory kinase -3.01  Creb3l1 Transcriptional activator -3.04  Dusp4 Phosphatase; negatively regulate (MAP) -3.08  kinases  ald3 Inhibits the DNA-binding transcription -3.13  factors  Wisp1 Downstream in the WNT1 signaling -3.32 -2.36  pathway  Rgs16 Inhibits signal transduction -3.78 -4.93  Call migration and bone development -4.18 -4.30  Filip11 Regulator of the anti-angiogenic activity -5.68 -2.21  Sox4 Regulation of embryonic development -6.48 -2.91  and Akap12 Scaffold protein in signal transduction -7.98 -3.87  Fstl1 Modulate action of growth factors -8.56 -2.16  CMme Metallopeptidase 15.17  CGprc5a Development, cellular growth and differentiation  Nefl Intracellular transport to axons and dendrites  Dusp6 Negatively regulate (MAP) kinases  Galnt7 GalNAc transferase 7 2.48  Adam9 Biological processes: cell-cell/matrix interactions  Flnc Crosslink actin filaments  Flnc Crosslink actin filaments  Cell adhesion and growth factor -2.58	<del>-</del> PP		2.02	
a Timp2 Inhibitors of the matrix metalloproteinases  TagIn Actin crosslinking/gelling protein -2.70 -3.56  CApp Transcriptional activator -2.88  Cdk1 Cell cycle regulatory kinase -3.01  Creb3l1 Transcriptional activator -3.04  Dusp4 Phosphatase; negatively regulate (MAP) -3.08  kinases  ald3 Inhibits the DNA-binding transcription -3.13  factors  Wisp1 Downstream in the WNT1 signaling -3.32 -2.36  pathway  Rgs16 Inhibits signal transduction -3.78 -4.93  cTns3 Cell migration and bone development -4.18 -4.30  Filip1 Regulator of the anti-angiogenic activity -5.68 -2.21  Sox4 Regulation of embryonic development -6.48 -2.91  Sox4 Regulation of embryonic development -7.98 -3.87  Fstl1 Modulate action of growth factors -7.98 -3.87  Fstl1 Modulate action of growth factors -8.56 -2.16  CMme Metallopeptidase 15.17  CGprc5a Development, cellular growth and differentiation  Nefl Intracellular transport to axons and dendrites  Dusp6 Negatively regulate (MAP) kinases 2.93  Galnt7 GalNAc transferase 7 2.48  Adam9 Biological processes: cell-cell/matrix interactions  Flnc Crosslink actin filaments -2.15  Rhob Cell adhesion and growth factor -2.58	<sup>c</sup> Dlg5		- 2.66	
TagIn Actin crosslinking/gelling protein — 2.70 — 3.56  CApp Transcriptional activator — 2.88  Cdk1 Cell cycle regulatory kinase — 3.01  Creb3l1 Transcriptional activator — 3.04  Dusp4 Phosphatase; negatively regulate (MAP) — 3.08  kinases  ald3 Inhibits the DNA-binding transcription — 3.13  factors  Wisp1 Downstream in the WNT1 signaling — 3.32 — 2.36  pathway  Rgs16 Inhibits signal transduction — 3.78 — 4.93  cTns3 Cell migration and bone development — 4.18 — 4.30  Filip11 Regulator of the anti-angiogenic activity — 5.68 — 2.21  Sox4 Regulation of embryonic development — 6.48 — 2.91  alakap12 Scaffold protein in signal transduction — 7.98 — 3.87  Fist1 Modulate action of growth factors — 8.56 — 2.16  CMme Metallopeptidase — 15.17  CGprc5a Development, cellular growth and differentiation  Nefl Intracellular transport to axons and dendrites  Dusp6 Negatively regulate (MAP) kinases — 2.93  Galnt7 GalNAc transferase 7 — 2.48  Adam9 Biological processes: cell-cell/matrix interactions  Flnc Crosslink actin filaments — 2.15  Rhob Cell adhesion and growth factor — 2.58	aT:2		2.60	
TagInActin crosslinking/gelling protein- 2.70- 3.56CAppTranscriptional activator- 2.88- 3.01Cdk1Cell cycle regulatory kinase- 3.01- 3.04Creb3l1Transcriptional activator- 3.08- 3.08Dusp4Phosphatase; negatively regulate (MAP)- 3.08- 3.13kinasesInhibits the DNA-binding transcription- 3.13- 2.36Wisp1Downstream in the WNT1 signaling pathway- 3.78- 4.93Rgs16Inhibits signal transduction- 3.78- 4.93cTns3Cell migration and bone development- 4.18- 4.30Filip11Regulator of the anti-angiogenic activity- 5.68- 2.21Sox4Regulation of embryonic development- 6.48- 2.91aAkap12Scaffold protein in signal transduction- 7.98- 3.87Fstl1Modulate action of growth factors- 8.56- 2.16CMmeMetallopeptidase15.17cGprc5aDevelopment, cellular growth and differentiation3.673.67NeflIntracellular transport to axons and dendrites3.31Dusp6Negatively regulate (MAP) kinases2.93Galn7GalNAc transferase 72.48Adam9Biological processes: cell-cell/matrix interactions- 2.15FlncCrosslink actin filaments- 2.15RhobCell adhesion and growth factor- 2.58	"Timp2		- 2.68	
CApp Cdk1 Cell cycle regulatory kinase -3.01 Creb3l1 Transcriptional activator -3.04 Dusp4 Phosphatase; negatively regulate (MAP) -3.08 kinases ald3 Inhibits the DNA-binding transcription -3.13 factors Wisp1 Downstream in the WNT1 signaling -3.32 -2.36 pathway Rgs16 Inhibits signal transduction -3.78 -4.93 Cell migration and bone development -4.18 -4.30 Filip11 Regulator of the anti-angiogenic activity -5.68 -2.21 Sox4 Regulation of embryonic development -6.48 -2.91 a^Akap12 Scaffold protein in signal transduction -7.98 -3.87 Fstl1 Modulate action of growth factors -8.56 -2.16 fMme Metallopeptidase 15.17 CGprc5a Development, cellular growth and differentiation Nefl Intracellular transport to axons and dendrites Dusp6 Negatively regulate (MAP) kinases 2.93 Galnt7 GalNAc transferase 7 2.48  Adam9 Biological processes: cell-cell/matrix interactions Flnc Crosslink actin filaments -2.15 Rhob Cell adhesion and growth factor -3.04	Taaln		- 2.70	- 3.56
Creb3l1 Transcriptional activator — 3.04 Dusp4 Phosphatase; negatively regulate (MAP) — 3.08 kinases  ald3 Inhibits the DNA-binding transcription — 3.13 factors  Wisp1 Downstream in the WNT1 signaling — 3.32 — 2.36 pathway  Rgs16 Inhibits signal transduction — 3.78 — 4.93 Cell migration and bone development — 4.18 — 4.30 Filip11 Regulator of the anti-angiogenic activity — 5.68 — 2.21 30x4 Regulation of embryonic development — 6.48 — 2.91 alkap12 Scaffold protein in signal transduction — 7.98 — 3.87 Fstl1 Modulate action of growth factors — 8.56 — 2.16 CMme Metallopeptidase — 15.17 CGprc5a Development, cellular growth and differentiation Nefl Intracellular transport to axons and dendrites Dusp6 Negatively regulate (MAP) kinases — 2.93 Galnt7 GalNAc transferase 7 — 2.48  Adam9 Biological processes: cell-cell/matrix interactions Flnc Crosslink actin filaments — 2.15 Rhob Cell adhesion and growth factor — 3.08  Action — 3.13 Action — 4.93 Action — 4.18 Action — 4.93 Action — 4.18 Action — 4.93 Action — 4.18 Action — 4.93 Action — 4.18 Action — 4.93 Action — 4.93 Action — 4.93 Action — 4.93 Actio	1			
Dusp4Phosphatase; negatively regulate (MAP)- 3.08 kinasesald3Inhibits the DNA-binding transcription- 3.13 factorsWisp1Downstream in the WNT1 signaling pathway- 3.32 - 2.36 pathwayRgs16Inhibits signal transduction- 3.78 - 4.93cTns3Cell migration and bone development- 4.18 - 4.30Filip11Regulator of the anti-angiogenic activity- 5.68 - 2.21Sox4Regulation of embryonic development- 6.48 - 2.91a^3Akap12Scaffold protein in signal transduction- 7.98 - 3.87Fst11Modulate action of growth factors- 8.56 - 2.16cMmeMetallopeptidase15.17cGprc5aDevelopment, cellular growth and differentiation3.67NeflIntracellular transport to axons and dendrites3.31Dusp6Negatively regulate (MAP) kinases2.93Galnt7GalNAc transferase 72.48Adam9Biological processes: cell-cell/matrix interactions2.44FlncCrosslink actin filaments- 2.15RhobCell adhesion and growth factor- 2.58	Cdk1	Cell cycle regulatory kinase	-3.01	
kinases  Inhibits the DNA-binding transcription factors  Wisp1 Downstream in the WNT1 signaling pathway  Rgs16 Inhibits signal transduction -3.78 -4.93  CTns3 Cell migration and bone development -4.18 -4.30  Filip11 Regulator of the anti-angiogenic activity -5.68 -2.21  Sox4 Regulation of embryonic development -6.48 -2.91  Sakap12 Scaffold protein in signal transduction -7.98 -3.87  Fist11 Modulate action of growth factors -8.56 -2.16  Mme Metallopeptidase 15.17  CGprc5a Development, cellular growth and differentiation  Nefl Intracellular transport to axons and dendrites  Dusp6 Negatively regulate (MAP) kinases 2.93  Galnt7 GalNAc transferase 7 2.48  Adam9 Biological processes: cell-cell/matrix interactions  Flnc Crosslink actin filaments -2.15  Rhob Cell adhesion and growth factor -2.58	l _			
ald3Inhibits the DNA-binding transcription- 3.13Misp1Downstream in the WNT1 signaling pathway- 3.32- 2.36Rgs16Inhibits signal transduction- 3.78- 4.93cTns3Cell migration and bone development- 4.18- 4.30Filip11Regulator of the anti-angiogenic activity- 5.68- 2.21Sox4Regulation of embryonic development- 6.48- 2.91aAkap12Scaffold protein in signal transduction- 7.98- 3.87Fstl1Modulate action of growth factors- 8.56- 2.166MmeMetallopeptidase15.17cGprc5aDevelopment, cellular growth and differentiation3.67NeflIntracellular transport to axons and dendrites3.31Dusp6Negatively regulate (MAP) kinases2.93Galnt7GalNAc transferase 72.48Adam9Biological processes: cell-cell/matrix interactions2.44FlncCrosslink actin filaments- 2.15RhobCell adhesion and growth factor- 2.58	Dusp4		- 3.08	
Wisp1Downstream in the WNT1 signaling pathway-3.32-2.36Rgs16Inhibits signal transduction-3.78-4.93CTns3Cell migration and bone development Filip11-4.18-4.30Regulator of the anti-angiogenic activity Box4-5.68-2.21Begulation of embryonic development Box4-6.48-2.91Brst11Modulate action of growth factors CMme-8.56-2.16CMmeMetallopeptidase15.17CGprc5aDevelopment, cellular growth and differentiation3.67NeflIntracellular transport to axons and dendrites3.31Dusp6Negatively regulate (MAP) kinases Galnt72.93Galn47GalNAc transferase 72.48Adam9Biological processes: cell-cell/matrix interactions2.24FlncCrosslink actin filaments Cell adhesion and growth factor-2.15	ald3		- 3.13	
Rgs16 Inhibits signal transduction -3.78 -4.93  Fins3 Cell migration and bone development -4.18 -4.30  Filip11 Regulator of the anti-angiogenic activity -5.68 -2.21  Sox4 Regulation of embryonic development -6.48 -2.91  Akap12 Scaffold protein in signal transduction -7.98 -3.87  Fstl1 Modulate action of growth factors -8.56 -2.16  Mme Metallopeptidase 15.17  Gprc5a Development, cellular growth and differentiation  Nefl Intracellular transport to axons and dendrites  Dusp6 Negatively regulate (MAP) kinases 2.93  Galnt7 GalNAc transferase 7 2.48  Adam9 Biological processes: cell-cell/matrix interactions  Flnc Crosslink actin filaments -2.15  Rhob Cell adhesion and growth factor -2.58	Wisp1		- 3.32	- 2.36
Cell migration and bone development -4.18 -4.30  Filip11 Regulator of the anti-angiogenic activity -5.68 -2.21  Sox4 Regulation of embryonic development -6.48 -2.91  Akap12 Scaffold protein in signal transduction -7.98 -3.87  Fistl1 Modulate action of growth factors -8.56 -2.16  Metallopeptidase 15.17  Corpress Development, cellular growth and differentiation  Nefl Intracellular transport to axons and dendrites  Dusp6 Negatively regulate (MAP) kinases 2.93  Galnt7 GalNAc transferase 7 2.48  Adam9 Biological processes: cell-cell/matrix interactions  Flnc Crosslink actin filaments -2.15  Rhob Cell adhesion and growth factor -2.58	·	pathway		
Filip11 Regulator of the anti-angiogenic activity -5.68 -2.21  Sox4 Regulation of embryonic development -6.48 -2.91  a Akap12 Scaffold protein in signal transduction -7.98 -3.87  Fst11 Modulate action of growth factors -8.56 -2.16  a Mme Metallopeptidase 15.17  a Garc5a Development, cellular growth and differentiation 16.7  Nefl Intracellular transport to axons and dendrites 16.7  Dusp6 Negatively regulate (MAP) kinases 2.93  Galnt7 GalNAc transferase 7 2.48  Adam9 Biological processes: cell-cell/matrix interactions  Flnc Crosslink actin filaments -2.15  Rhob Cell adhesion and growth factor -5.68 -2.21  Action 17.88 -2.21  Action 2.58 -2.21  Action 2.				
Sox4Regulation of embryonic development-6.48-2.91aAkap12Scaffold protein in signal transduction-7.98-3.87Fst11Modulate action of growth factors-8.56-2.16CMmeMetallopeptidase15.17bevelopment, cellular growth and differentiation3.67NeflIntracellular transport to axons and dendrites2.93Dusp6Negatively regulate (MAP) kinases2.93Galnt7GalNAc transferase 72.48Adam9Biological processes: cell-cell/matrix interactions2.44FlncCrosslink actin filaments-2.15RhobCell adhesion and growth factor-2.58				
a Akap12Scaffold protein in signal transduction-7.98-3.87Fstl1Modulate action of growth factors-8.56-2.166 MmeMetallopeptidase15.175 Gprc5aDevelopment, cellular growth and differentiation3.67NeflIntracellular transport to axons and dendrites3.31Dusp6Negatively regulate (MAP) kinases2.93Galnt7GalNAc transferase 72.48Adam9Biological processes: cell-cell/matrix interactions2.44FlncCrosslink actin filaments-2.15RhobCell adhesion and growth factor-2.58				
Fstl1 Modulate action of growth factors  FMme Metallopeptidase 15.17  CGprc5a Development, cellular growth and differentiation  Nefl Intracellular transport to axons and dendrites  Dusp6 Negatively regulate (MAP) kinases  Galnt7 GalNAc transferase 7 2.48  Adam9 Biological processes: cell-cell/matrix interactions  Flnc Crosslink actin filaments -2.15  Rhob Cell adhesion and growth factor -2.58				
CMmeMetallopeptidase15.17CGprc5aDevelopment, cellular growth and differentiation3.67NeflIntracellular transport to axons and dendrites3.31Dusp6Negatively regulate (MAP) kinases2.93Galnt7GalNAc transferase 72.48Adam9Biological processes: cell-cell/matrix interactions2.44FlncCrosslink actin filaments-2.15RhobCell adhesion and growth factor-2.58				
cGprc5aDevelopment, cellular growth and differentiation3.67NeflIntracellular transport to axons and dendrites3.31Dusp6Negatively regulate (MAP) kinases2.93Galnt7GalNAc transferase 72.48Adam9Biological processes: cell-cell/matrix interactions2.44FlncCrosslink actin filaments-2.15RhobCell adhesion and growth factor-2.58			0.50	
NeflIntracellular transport to axons and dendrites3.31Dusp6Negatively regulate (MAP) kinases2.93Galnt7GalNAc transferase 72.48Adam9Biological processes: cell-cell/matrix interactions2.44FlncCrosslink actin filaments-2.15RhobCell adhesion and growth factor-2.58	l -	Development, cellular growth and		
dendrites  Dusp6 Negatively regulate (MAP) kinases 2.93 Galnt7 GalNAc transferase 7 2.48  Adam9 Biological processes: cell-cell/matrix interactions  Flnc Crosslink actin filaments -2.15 Rhob Cell adhesion and growth factor -2.58				
Dusp6 Galnt7Negatively regulate (MAP) kinases GalNAc transferase 72.93 2.48Adam9Biological processes: cell-cell/matrix interactions2.44Flnc RhobCrosslink actin filaments Cell adhesion and growth factor-2.15 -2.58	Nefl			3.31
Gaint7GainNAc transferase 72.48Adam9Biological processes: cell-cell/matrix interactions2.44FIncCrosslink actin filaments-2.15RhobCell adhesion and growth factor-2.58	Duché			2.02
Adam9Biological processes: cell-cell/matrix interactions2.44FlncCrosslink actin filaments- 2.15RhobCell adhesion and growth factor- 2.58				
interactions  Flnc Crosslink actin filaments – 2.15  Rhob Cell adhesion and growth factor – 2.58				
Flnc Crosslink actin filaments -2.15 Rhob Cell adhesion and growth factor -2.58	Adam9			2.44
3 · · · · · · · · · · · · · · · · · · ·	Flnc			- 2.15
signaling	Rhob			- 2.58
		signaling		

Table 1.	(Continued)	
Gene	Gene function	V vs PF V vs P3
Mtss1 Igfbp3	Actin bundling Bind and inhibit IGF (affect growth)	- 2.77 - 3.33
°Dyrk2	Cellular growth and/or development	- 3.36

Abbreviations: FGF, fibroblast growth factors; EGFR, epidermal growth factor receptor; IGF, insulin-like growth factor. Total RNA was isolated using the miRNeasy mini kit (Qiagen), allowing for the isolation of RNA < 30 bp in length, according to the manufacturer's specifications. Poly-A+ mRNA was isolated from 4  $\mu g$  total RNA, to generate the cDNA libraries, using the Illumina sample preparation kit according to the manufacturer's specifications (Illumina, San Diego, CA, USA). The cDNA libraries were provided a unique index identifier, allowing the clustering of several samples into a single sequencing lane, and deep-sequencing analyses were performed in triplicate from three independent cell growth, RNA isolation and cDNA library constructions. The raw data were groomed and trimmed for quality of the read using online Galaxy analysis (https://usegalaxy.org), resulting in 40-41 high-quality base pair reads for each sequence with between 4-6 million independent reads for each sample. The sequences were mapped to the mouse genome using Tophat analysis, transcripts were assembled using the Cufflinks program, and individual replicates were merged into a single file using Cuffmerge. The resulting transcript reads were normalized using Fragments Per Kilobase of transcript per Million mapped reads analysis, which normalizes each identified sequence for the length of the identified transcript and the volume of the total read yield from each run. Differential expression was determined from these normalized values comparing vector versus Pax3-FOXO1 (V vs PF) or vector versus Pax3 (V vs P3) using the Cuffdiff program, which not only compares differential expression of the merged files between sets but also utilizes the sequence results from the three independent determinations within each set to assign statistical significance to the differential expression. alndicates genes with similar trends in expression changes in human tumor samples.<sup>22–25</sup> blndicates genes demonstrated in the literature to be direct targets of PAX3 or PAX3-FOXO1.<sup>15,16,20,21</sup> clndicates genes with known PAX3-FOXO1 binding sites in their promoter.<sup>20</sup>

downregulated, with the top four being downregulated > 12-fold. In conjunction with the results of our PubMed search, which also described the target genes responsible for the invasive effect of the miRNA, we used miRTarBase<sup>29</sup> to identify known target genes whose biological function may contribute to an invasive phenotype, with validation on miRTarBase by at least two independent experimental methods. Interestingly, only a small number of the altered miRNA have the expected inverse correlation to our observed changes in mRNA expression (miR-222/miR-221 and TIMP2, and miR-362 and CD82).

A similar analysis determined a total of 58 miRNA whose expression changed upon the expression of PAX3 (25 downregulated and 33 upregulated) relative to the empty vector negative control (data not shown). Of these genes, a PubMed search determined that 7 are important for promoting while 16 inhibit cellular invasion. Of those miRNA, 5/7 (71%) that promote invasion are decreased whereas 10/16 (63%) that inhibit invasion are increased, with the top inhibitory miRNA being increased >15-fold. Finally, for both PAX3-FOXO1- and PAX3-dependent miRNA changes, the literature provides direct evidence for the genes they target in order to exert their effects on invasion (Table 2). As seen with PAX3-FOXO1 changes, only one of the PAX3-altered miRNA has the expected inverse correlation to mRNA expression (miR-206 and MET, Tables 1 and 2). Interestingly, three sets of miRNA are present as clusters in the mouse genome and have similar changes in expression. These include miRNA 222 and 221, which are upregulated to a similar extent by PAX3-FOXO1 while being downregulated to a similar extent by PAX3 (Table 2, a); miRNA 362 and 532, which are downregulated to a similar extent by PAX3-FOXO1 but are unaffected by PAX3 (Table 2, b); and miRNA 133b and 206, which are unaffected by

miRNA	Gene target	Gene function	V vs PF	V vs P3
miRNA that promo	ote tumor cell invasion			
515-3p	te tamer cen invasion		+30.45	+5.01
196a-5p	HOXA5	Developmental transcription factor	+24.39	
	ING5	Suppresses growth and invasion	,	
30d-3p	GALNT7	Glycopeptide transferase	+3.39	_
301a-3p	SMAD4	Signal transduction activator	+3.07	
501a-5p	TXNIP	Suppressor of tumor cell growth	T3.07	_
	BBC3	Pro-apoptotic protein		
	PTEN	Tumor suppressor protein		
	COL2A1	Collagen 2 alpha 1		
	RUNX3	Transcriptional tumor suppressor		
	TGFBR2	TGF beta receptor		
	SOCS6	Suppressor of cytokine signaling		
222-3p	TIMP2	Metallopeptidase inhibitor	+2.78	- 2.52
	TIMP3	Metallopeptidase inhibitor		
221-5p	RECK	Negatively regulates metalloproteinases	+2.63	_
	TIMP2	Metallopeptidase inhibitor		
	MMP3	Matrix metalloproteinase		
	MMP9	Matrix metalloproteinase		
	PTEN	Tumor suppressor		
	TIMP3	Metallopeptidase inhibitor		
55-5p	<u>-</u>		+2.18	- 2.53
221-3p	MMP3	Matrix metalloproteinase	+2.10	- 3.31
221 JP	MMP9	Matrix metalloproteinase	12.10	5.51
	PTEN			
		Tumor suppressor		
00.5	TIMP3	Metallopeptidase inhibitor	2.25	
83-5p	ITGB1	Integrin—cell adhesion receptor	+2.05	
	SOCS6	Cytokine signal transduction regulator		
	PDCD4	Inhibit translation—tumor suppressor		
362-3p	CD82	Metastasis suppressor protein	− 3.67	_
et-7g-5p	GAB2	Signaling adaptor protein	_	- 2.18
<b>.</b>	FN1	Cell surface adhesion molecule		
28a-5p	CCND1	Cyclin D1	_	- 2.10
.04.56	HOXB3	Developmental transcription factor		20
3b-3p	PTEN	Tumor suppressor	<u></u>	+2.19
.JD-JP	ATG12	Regulates autophagy		12.13
	HMGB2	Architectural transcription factor		
กRNA that inhibit	tumor formation			
a-3p	TAGLN2	Unknown function	<b>– 18.50</b>	_
45a-5p	HIF-2 alpha	Hypoxia-induced transcription factor	<b>– 16.76</b>	_
·	EGFR	Growth factor receptor		
	OCT4	Developmental transcription factor		
	MUC1	Cell adhesion molecule		
	MYC	Growth-related transcription factor		
	D52			
22- 5-		Unknown—overexpressed in cancer cells	16.76	
33a-5p	TAGLN2	Unknown function	– 16.76	_
	LASP1	Actin-binding protein		
	FSCN	Actin-binding protein		
	MMP14	Matrix metalloproteinase		
35-5p	SP1	Transcriptional regulator	<b>– 12.50</b>	+2.95
532-5p	CXCL2	Regulatory chemokine	<i>–</i> 4.67	_
48a-3p	S1PR1	Receptor to regulate adhesion	- 4.41	- 2.98
33a-3p	TAGLN2	Unknown function	- 3.72	_
	LASP1	Actin-binding protein		
	FSCN	Actin-binding protein		
	MMP14	Matrix metalloproteinase		
40h 2			2.44	
48b-3p	WNT	Developmental ligand	- 3.44	_
	NRP1	Membrane-bound signaling protein		
9a-3p	FRA1	FOS family member	<b>– 2.87</b>	_
.9a-3p	HSP47	Serine proteinase inhibitor	<b>– 2.65</b>	_
•	LAMC2	Extracellular matrix glycoprotein		
	ITGA6	Integrin—cell adhesion receptor		
4b-5p		•	- 2.60	_
49-3p	FOXM1	Transcription factor	- 2.34	- 7.04
143-2h	RAP1a	Adhesion signaling protein	2.5 .	7.04
	RAP1b	Adhesion signaling protein		
122h F			2.26	
133b-5p	MMP14	Matrix metalloproteinase	- 2.26	_
0d-5p	CCNE2	Cyclin E2	- 2.22	_
74-3p	RAC1	GTPase-signaling molecule	- 2.07	_
	EGFR	Growth factor receptor		
	EP300	Histone acetyltransferase—chromatin		
39-5p	NACC1	Transcriptional corepressor	- 1.93	+2.19
JP	BCL6	Transcriptional corepressor	1.23	12.19
		Regulator of p53 stability		
	MDM2			

miRNA	Gene target	Gene function	V vs PF	V vs P3
338-3p	SMO	G-protein coupled receptor	+2.09	+15.27
	MMP9	Matrix metalloproteinase		
	PREX2a	Guanine nucleotide exchange factor		
	ZEB2	Transcriptional repressor		
	MACC1	Transcriptional activator		
<sup>c</sup> 133b-3p	FSCN1	Actin-binding protein	_	+5.74
	MMP9	Matrix metalloproteinase		
<sup>c</sup> 206-3p	MET	Growth factor receptor	<del>_</del>	+4.49
	Cdc42	Regulates actin polymerization		
	NOTCH3	Developmental receptor		
582-5p	RAB27a	Membrane-bound GTPase	_	+4.41
	PGGT1B	Geranylgeranyl transferase enzyme		
	LRRK2	Leucine-rich repeat kinase		
	DIXDC1	Positive regulator of Wnt signaling		
345-5p	BAG3	Inhibits HSP chaperone activity	<del>_</del>	+3.09
<sup>c</sup> 206-5p	MET	Growth factor receptor	<del>_</del>	+2.31
	Cdc42	Regulates actin polymerization		
	NOTCH3	Developmental receptor		
486-5p	ARHGAP5	Rho GTPase-activating protein	<del>_</del>	+2.09
	PIK3R1	PI3K regulatory subunit		
	OLFM4	Extracellular matrix glycoprotein		
31-3p			+4.19	+2.05
34c-3p	PAC1	Adenylate cyclase-activating receptor	+3.38	+3.33
·	MARCKS	F-actin crosslinking protein		
	elF4E	Translation elongation factor		
615-5p	AKT2	Ser/Thr protein kinase	+34.90	- 2.53
	IGF2	Growth factor ligand		
193-3p	ERBB4	Receptor tyrosine growth factor receptor	+3.26	_
- 1	S6K2	Ribosomal kinase		
181c-3p	SMAD7	Negatively regulates TGF beta signaling	_	- 3.77
30a-5p	ITGB3	Integrin—cell-adhesion receptor	_	- 2.50
	NCAM	Cellular-adhesion molecule		
	SEPT7	Cytoskeletal GTPase—actin organization		
	MTDH	Activates NFkB		
30c-2-3p	TRADD	Mediates apoptosis and NFkB signaling	_	-2.32
•	CCNE1	Cyclin E1		

Abbreviation: HSP, heat shock proteins. Total RNA was isolated using the miRNeasy mini kit (Qiagen), allowing for the isolation of RNA < 30 bp in length, according to the manufacturer's specifications. miRNA was isolated from 4  $\mu$ g total RNA to generate the cDNA libraries, using the Illumina sample preparation kits according to the manufacturer's specifications. The cDNA libraries were provided a unique index identifier, allowing the clustering of several samples into a single sequencing lane, and deep-sequencing analyses were performed in triplicate from three independent cell growth, RNA isolation and cDNA library constructions. Raw fast q sequences were obtained from the Illumina Genome Analyzer II (Illumina, San Diego, CA, USA) using the 'Demultiplex' algorithm in the CASAVA 1.8.2 software (Illumina) that allows the identification of individual samples by 'index sequences' contained within the adapters and introduced during the adapter ligation and amplification of the samples. miRNAKey was used for data analysis at default settings. The pipeline clips the Illumina 3' adapter sequences from the reads, maps the clipped reads to miRBase and uses the Seq-EM algorithm to estimate the distribution of multiply mapped reads across the observed miRNAs. Sequences < 16 bases after adaptor clipping were removed. The read counts obtained were then used for differential expression analysis comparing vector versus Pax3-FOXO1 (V vs PF) or vector versus Pax3 (V vs P3) between control and experimental samples using EBSeq from the R package with a false discovery rate of 5%. We used the default 'Median Normalization' in EBSeq to make the counts comparable across samples. Target genes for each miRNA were identified either as a result of the indicated PubMed search or using miRTarBase, <sup>29</sup> which lists experimentally validated direct targets. Several miRNA are expressed in clusters and show similar changes in expression. <sup>a</sup>Upregulated by PAX3-FOXO1 and downregulated by PAX3. <sup>b</sup>Downregulated by PAX3-FOXO1 and upregulate

PAX3-FOXO1 are upregulated to a similar extent by PAX3 (Table 2 c)

To validate the observed changes, we performed a quantitative RT–PCR analysis on a subset of mRNA and miRNA. We tested IGF2, which was reported to promote cellular invasion in a variety of tumors, <sup>30–33</sup> and CDK1, which has alternative roles in inhibiting cellular invasion in different tumor models. <sup>34–38</sup> We also examined two miRNA (miR-196a-5p and miR301a-3p), both demonstrated to promote cellular invasion. <sup>39–44</sup> We observed quantitative and significant changes in expression for all of the mRNA and miRNA examined that are consistent with our mRNA deep-sequencing results (Figure 1).

Our comparative transcriptomic results suggest that the mRNA and miRNA changes induced by the oncogenic fusion protein would be predicted to promote cellular invasion, whereas those changes that occur in a PAX3-dependent manner should inhibit the invasive capacity of cells. Therefore, we used a standard invasion assay to determine whether our observed mRNA and

miRNA changes translate into experimental differences in the invasive potential of primary myoblasts. Consistent with our mRNA and miRNA changes, we observed a nearly 2-fold increase in primary myoblasts expressing PAX3-FOXO1, consistent with the previous reports, <sup>16</sup> whereas cells expressing PAX3 had a nearly 4-fold decrease in invasive potential (Figure 2).

Taken together, our results build off of previous work, in which a single gene was examined, <sup>16</sup> and show that the sole expression of PAX3-FOXO1 in the absence of any complimentary genetic mutations is capable of globally altering mRNA and miRNA levels to promote the invasive capacity of primary myoblasts. Further, this is the first study to examine how the oncogenic fusion protein alters miRNA levels, which combined with our global mRNA results allow us to develop a more expansive picture of the underlying regulatory mechanisms by which the expression of PAX3-FOXO1 promotes invasion.

In this regulatory mechanism, the somatic and random acquisition of the chromosomal translocation creates the fusion

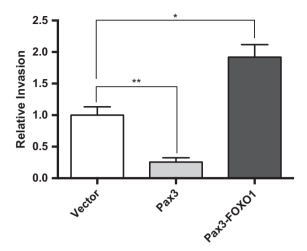


Figure 2. Pax3-FOXO1 promotes whereas Pax3 inhibits primary myoblast invasive capacity. Invasive capacity was determined using stably transduced proliferating primary myoblasts (empty vector (white bars), PAX3 (gray bars) or PAX3-FOXO1 (black bars)) using the BD Biocoat Tumor Invasion System (Becton Dickinson, Franklin Lakes, NJ, USA). About 50 000 cells suspended in proliferation media were added to the insert plate with proliferation media supplemented with hepatocyte growth factor (hHGF, PeproTech, Rocky Hill, NJ, USA) at 25 ng/ml being used as the chemoattractant. After 24 h of incubation, the insert system was transferred to a second 24-well plate containing calcein AM in Hank's balanced salt solution (HBSS) that enabled the fluorescent labeling of cells that invaded through the Matrigel matrix. Fluorescence of the invaded cells was read at wavelengths of 494/517 (Ex/Em) using a Synergy HT multi-well microplate reader (BioTek, Winooski, VT, USA). *P*-values were computed using non-parametric one-way ANOVA analysis comparing all samples with results obtained, with cells expressing empty vector (\*P = 0.03, \*\*P = 0.001). ANOVA, analysis of variance.

protein, which alters, either directly or indirectly, the expression of mRNA important for enhancing cellular invasion. PAX3-FOXO1 achieves this by both decreasing the expression of genes that inhibit invasion (80% of downregulated genes) while also greatly increasing the expression of nearly half of the altered genes that promote invasion (Table 1). However, our results demonstrate that miRNAs, which post-transcriptionally 'fine tune' gene expression, also have a significant role, as nearly all of the increased miRNAs promote invasion and a majority of the decreased miRNAs inhibit invasion. Further, a closer inspection of the data reveals that although some of the miRNA changes have an inverse correlation to target genes present in our results, there are only three miRNA (miR-221, miR-222 and miR-362) that have such a correlation with their target genes (TIMP2 and CD82, respectively). Therefore, instead of post-transcriptionally contributing to our observed mRNA changes, the altered miRNA target a different set of genes important for invasive capacity, thereby greatly increasing the number of affected genes.

The global nature of the mRNA and miRNA expression changes that result upon the sole expression of PAX3-FOXO1 provide a basis for how it may be necessary to rethink approaches to the development of therapies for ARMS. At present, many developmental ARMS therapies focus on attacking a single gene or pathway mechanistically located downstream of the fusion protein. However, given that the expression of PAX3-FOXO1 alters the expression of 70 different genes and 27 different miRNAs to affect the invasive potential of cells, it is not too surprising that such focused and targeted therapies are not proving effective in Phase I or Phase II clinical trials for ARMS.<sup>45–48</sup> It is conceivable that the loss of a single gene through these targeted and focused therapies could easily be compensated through the changes in

nearly 100 other affected genes, thereby negating the effects of the treatment.

Work in the past few years identified multiple aspects of the invasive process as potential targets for therapy development. Along these lines we propose developing a multi-faceted regimen that targets several of these processes, targets that include tumorpromoting genes we found to be the most highly upregulated in our study (Table 1). These processes include cytoskeletal remodeling, which is mediated in part by the intracellular signaling cysteine proteases calpains<sup>49</sup> (Capn6 is upregulated 33-fold in our study), cellular adhesion mediated by such molecules as mesothelin<sup>50</sup> (Msln is upregulated 21-fold in our study) and matrix metalloproteases,<sup>51</sup> in particular the Adamts family of proteases<sup>52</sup> (Adamts1 is upregulated 22-fold and Adamts5 is upregulated 7.5-fold in our study). A regimen that minimally targets these three processes would inhibit the necessary biological events required for invasion and metastasis. Alternatively, inhibiting one of these events (for example, matrix metalloproteases) could serve as one arm of a novel multi-faceted regimen for the treatment of ARMS, a regiment that also targets other ARMS molecular processes such as inhibiting phosphorylation of PAX3-FOXO1, 17 attacking aneuploid cells and preventing enhanced proliferation.

## **CONFLICT OF INTEREST**

The authors declare no conflict of interest.

### **ACKNOWLEDGEMENTS**

Funding for this project is from the National Cancer Institute grant R01CA138656, the Louisiana State University School of Medicine Research Enhancement Bridge Grant and the Louisiana Cancer Research Consortium (LCRC) (ADH). JZ has been partially supported by grants from the National Institute of General Medicine Sciences (NIGMS) grants P20GM103501, subproject #2, P30GM114732 and U54GM104940-01, and the National Institute on Minority Health and Health Disparities (NIMHD) grants P20MD004817 and U54MD006176-01. All deep sequencing was performed in the LCRC Translational Genomics Core facility.

## **REFERENCES**

- 1 Horn RC Jr., Enterline HT. Rhabdomyosarcoma: a clinicopathological study and classification of 39 cases. *Cancer* 1958; **11**: 181–199.
- 2 Punyko JA, Mertens AC, Baker KS, Ness KK, Robison LL, Gurney JG. Long-term survival probabilities for childhood rhabdomyosarcoma. A population-based evaluation. *Cancer* 2005; **103**: 1475–1483.
- 3 Galili N, Davis RJ, Fredericks WJ, Mukhopadhyay S, Rauscher FJ 3rd, Emanuel BS et al. Fusion of a fork head domain gene to PAX3 in the solid tumour alveolar rhabdomyosarcoma. *Nat Genet* 1993; **5**: 230–235.
- 4 Shapiro DN, Sublett JE, Li B, Downing JR, Naeve CW. Fusion of PAX3 to a member of the forkhead family of transcription factors in human alveolar rhabdomyosarcoma. *Cancer Res* 1993; **53**: 5108–5112.
- 5 Fredericks WJ, Galili N, Mukhopadhyay S, Rovera G, Bennicelli J, Barr FG et al. The PAX3-FKHR fusion protein created by the t(2;13) translocation in alveolar rhabdomyosarcomas is a more potent transcriptional activator than PAX3. Mol Cell Biol 1995; 15: 1522–1535.
- 6 Hollenbach AD, Sublett JE, McPherson CJ, Grosveld G. The Pax3-FKHR oncoprotein is unresponsive to the Pax3-associated repressor hDaxx. Embo J 1999; 18: 3702–3711
- 7 Miller PJ, Hollenbach AD. The oncogenic fusion protein Pax3-FKHR has a greater post-translational stability relative to Pax3 during early myogenesis. *Biochim Biophys Acta* 2007; **1770**: 1450–1458.
- 8 Epstein JA, Song B, Lakkis M, Wang C. Tumor-specific PAX3-FKHR transcription factor, but not PAX3, activates the platelet-derived growth factor alpha receptor. *Mol Cell Biol* 1998; **18**: 4118–4130.
- 9 Ayalon D, Glaser T, Werner H. Transcriptional regulation of IGF-I receptor gene expression by the PAX3-FKHR oncoprotein. Growth Horm IGF Res 2001; 11: 289–297.
- 10 Ginsberg JP, Davis RJ, Bennicelli JL, Nauta LE, Barr FG. Up-regulation of MET but not neural cell adhesion molecule expression by the PAX3-FKHR fusion protein in alveolar rhabdomyosarcoma. Cancer Res 1998; 58: 3542–3546.

- 11 Linardic CM. PAX3-FOXO1 fusion gene in rhabdomyosarcoma. *Cancer Lett* 2008; **270**: 10–18.
- 12 Sorensen PH, Lynch JC, Qualman SJ, Tirabosco R, Lim JF, Maurer HM et al. PAX3-FKHR and PAX7-FKHR gene fusions are prognostic indicators in alveolar rhabdomyosarcoma: a report from the children's oncology group. J Clin Oncol 2002; 20: 2672–2679.
- 13 Anderson J, Ramsay A, Gould S, Pritchard-Jones K. PAX3-FKHR induces morphological change and enhances cellular proliferation and invasion in rhabdomyosarcoma. Am J Pathol 2001; 159: 1089–1096.
- 14 Lagha M, Kormish JD, Rocancourt D, Manceau M, Epstein JA, Zaret KS et al. Pax3 regulation of FGF signaling affects the progression of embryonic progenitor cells into the myogenic program. Genes Dev 2008; 22: 1828–1837.
- 15 Khan J, Bittner ML, Saal LH, Teichmann U, Azorsa DO, Gooden GC et al. cDNA microarrays detect activation of a myogenic transcription program by the PAX3-FKHR fusion oncogene. Proc Natl Acad Sci USA 1999; 96: 13264–13269.
- 16 Marshall AD, Lagutina I, Grosveld GC. PAX3-FOXO1 induces cannabinoid receptor 1 to enhance cell invasion and metastasis. Cancer Res 2011; 71: 7471–7480.
- 17 Loupe JM, Miller PJ, Ruffin DR, Stark MW, Hollenbach AD. Inhibiting phosphorylation of the oncogenic PAX3-FOXO1 reduces alveolar rhabdomyosarcoma phenotypes identifying novel therapy options. *Oncogenesis* 2015; **4**: e145.
- 18 Dietz KN, Miller PJ, Hollenbach AD. Phosphorylation of serine 205 by the protein kinase CK2 persists on Pax3-FOXO1, but not Pax3, throughout early myogenic differentiation. *Biochemistry* 2009; 48: 11786–11795.
- 19 Dietz KN, Miller PJ, Iyengar AS, Loupe JM, Hollenbach AD. Identification of serines 201 and 209 as sites of Pax3 phosphorylation and the altered phosphorylation status of Pax3-FOXO1 during early myogenic differentiation. *Int J Biochem Cell Biol* 2011: 43: 936–945.
- 20 Cao L, Yu Y, Bilke S, Walker RL, Mayeenuddin LH, Azorsa DO et al. Genome-wide identification of PAX3-FKHR binding sites in rhabdomyosarcoma reveals candidate target genes important for development and cancer. Cancer Res 2010; 70: 6497–6508.
- 21 Khan J, Simon R, Bittner M, Chen Y, Leighton SB, Pohida T et al. Gene expression profiling of alveolar rhabdomyosarcoma with cDNA microarrays. Cancer Res 1998; 58: 5009–5013.
- 22 Davicioni E, Finckenstein FG, Shahbazian V, Buckley JD, Triche TJ, Anderson MJ. Identification of a PAX-FKHR gene expression signature that defines molecular classes and determines the prognosis of alveolar rhabdomyosarcomas. Cancer Res 2006; 66: 6936–6946.
- 23 De Pitta C, Tombolan L, Albiero G, Sartori F, Romualdi C, Jurman G et al. Gene expression profiling identifies potential relevant genes in alveolar rhabdomyosarcoma pathogenesis and discriminates PAX3-FKHR positive and negative tumors. Int J Cancer 2006; 118: 2772–2781.
- 24 Ebauer M, Wachtel M, Niggli FK, Schafer BW. Comparative expression profiling identifies an in vivo target gene signature with TFAP2B as a mediator of the survival function of PAX3/FKHR. Oncogene 2007; 26: 7267–7281.
- 25 Lae M, Ahn EH, Mercado GE, Chuai S, Edgar M, Pawel BR et al. Global gene expression profiling of PAX-FKHR fusion-positive alveolar and PAX-FKHR fusionnegative embryonal rhabdomyosarcomas. J Pathol 2007; 212: 143–151.
- 26 Zalc A, Rattenbach R, Aurade F, Cadot B, Relaix F. Pax3 and Pax7 play essential safeguard functions against environmental stress-induced birth defects. *Dev Cell* 2015; 33: 56–66.
- 27 Begum S, Emami N, Cheung A, Wilkins O, Der S, Hamel PA. Cell-type-specific regulation of distinct sets of gene targets by Pax3 and Pax3/FKHR. *Oncogene* 2005; 24: 1860–1872.
- 28 Epstein JA, Shapiro DN, Cheng J, Lam PY, Maas RL. Pax3 modulates expression of the c-Met receptor during limb muscle development. *Proc Natl Acad Sci USA* 1996;
- 29 Hsu SD, Tseng YT, Shrestha S, Lin YL, Khaleel A, Chou CH et al. miRTarBase update 2014: an information resource for experimentally validated miRNA-target interactions. Nucleic Acids Res 2014: 42: D78–D85.
- 30 Chen YW, Boyartchuk V, Lewis BC. Differential roles of insulin-like growth factor receptor- and insulin receptor-mediated signaling in the phenotypes of hepatocellular carcinoma cells. Neoplasia 2009; 11: 835–845.
- 31 Mu Q, Wang L, Yu F, Gao H, Lei T, Li P et al. Imp2 regulates GBM progression by activating IGF2/PI3K/Akt pathway. Cancer Biol Ther 2015; 16: 623–633.
- 32 Pivonello C, Negri M, De Martino MC, Napolitano M, de Angelis C, Provvisiero DP et al. The dual targeting of insulin and insulin-like growth factor 1 receptor enhances the mTOR inhibitor-mediated antitumor efficacy in hepatocellular carcinoma. Oncotarget 2016; 7: 9718–9731.
- 33 Sciacca L, Mineo R, Pandini G, Murabito A, Vigneri R, Belfiore A. In IGF-I receptor-deficient leiomyosarcoma cells autocrine IGF-II induces cell invasion and protection from apoptosis via the insulin receptor isoform A. Oncogene 2002; 21: 8240–8250.
- 34 Chang WL, Yu CC, Chen CS, Guh JH. Tubulin-binding agents down-regulate matrix metalloproteinase-2 and -9 in human hormone-refractory prostate cancer cells a critical role of Cdk1 in mitotic entry. *Biochem Pharmacol* 2015; **94**: 12–21.

- 35 Vanan I, Dong Z, Tosti E, Warshaw G, Symons M, Ruggieri R. Role of a DNA damage checkpoint pathway in ionizing radiation-induced glioblastoma cell migration and invasion. Cell Mol Neurobiol 2012; 32: 1199–1208.
- 36 Wei Y, Chen YH, Li LY, Lang J, Yeh SP, Shi B et al. CDK1-dependent phosphorylation of EZH2 suppresses methylation of H3K27 and promotes osteogenic differentiation of human mesenchymal stem cells. Nat Cell Biol 2011; 13: 87–94.
- 37 Zhang C, Elkahloun AG, Robertson M, Gills JJ, Tsurutani J, Shih JH et al. Loss of cytoplasmic CDK1 predicts poor survival in human lung cancer and confers chemotherapeutic resistance. PLoS ONE 2011; 6: e23849.
- 38 Zhang L, Chen X, Stauffer S, Yang S, Chen Y, Dong J. CDK1 phosphorylation of TAZ in mitosis inhibits its oncogenic activity. *Oncotarget* 2015; **6**: 31399–31412.
- 39 Egawa H, Jingushi K, Hirono T, Ueda Y, Kitae K, Nakata W *et al.* The miR-130 family promotes cell migration and invasion in bladder cancer through FAK and Akt phosphorylation by regulating PTEN. *Sci Rep* 2016; **6**: 20574.
- 40 Fang Y, Sun B, Xiang J, Chen Z. MiR-301a promotes colorectal cancer cell growth and invasion by directly targeting SOCS6. Cell Physiol Biochem 2015; 35: 227–236.
- 41 Liu M, Du Y, Gao J, Liu J, Kong X, Gong Y et al. Aberrant expression miR-196a is associated with abnormal apoptosis, invasion, and proliferation of pancreatic cancer cells. *Pancreas* 2013; **42**: 1169–1181.
- 42 Liu XH, Lu KH, Wang KM, Sun M, Zhang EB, Yang JS et al. MicroRNA-196a promotes non-small cell lung cancer cell proliferation and invasion through targeting HOXA5. BMC Cancer 2012; 12: 348.
- 43 Wang M, Li C, Yu B, Su L, Li J, Ju J et al. Overexpressed miR-301a promotes cell proliferation and invasion by targeting RUNX3 in gastric cancer. J Gastroenterol 2013; 48: 1023–1033.
- 44 Zhang W, Zhang T, Jin R, Zhao H, Hu J, Feng B *et al.* MicroRNA-301a promotes migration and invasion by targeting TGFBR2 in human colorectal cancer. *J Exp Clin Cancer Res* 2014; **33**: 113.
- 45 Bagatell R, Norris R, Ingle AM, Ahern C, Voss S, Fox E *et al.* Phase 1 trial of temsirolimus in combination with irinotecan and temozolomide in children, adolescents and young adults with relapsed or refractory solid tumors: a Children's Oncology Group Study. *Pediatr Blood Cancer* 2014; **61**: 833–839.
- 46 Geoerger B, Kieran MW, Grupp S, Perek D, Clancy J, Krygowski M et al. Phase II trial of temsirolimus in children with high-grade glioma, neuroblastoma and rhabdomyosarcoma. Eur J Cancer 2012; 48: 253–262.
- 47 Pappo AS, Vassal G, Crowley JJ, Bolejack V, Hogendoorn PC, Chugh R et al. A phase 2 trial of R1507, a monoclonal antibody to the insulin-like growth factor-1 receptor (IGF-1R), in patients with recurrent or refractory rhabdomyosarcoma, osteosarcoma, synovial sarcoma, and other soft tissue sarcomas: results of a Sarcoma Alliance for Research through Collaboration Study. Cancer 2014; 120: 2448–2456.
- 48 Weigel B, Malempati S, Reid JM, Voss SD, Cho SY, Chen HX et al. Phase 2 trial of cixutumumab in children, adolescents, and young adults with refractory solid tumors: a report from the Children's Oncology Group. Pediatr Blood Cancer 2014; 61: 452–456.
- 49 Leloup L, Wells A. Calpains as potential anti-cancer targets. *Expert Opin Ther Targets* 2011; **15**: 309–323.
- 50 Pastan I, Hassan R. Discovery of mesothelin and exploiting it as a target for immunotherapy. *Cancer Res* 2014; **74**: 2907–2912.
- 51 Cathcart J, Pulkoski-Gross A, Cao J. Targeting matrix metalloproteinases in cancer: bringing new life to old ideas. *Genes Dis* 2015; **2**: 26–34.
- 52 Viapiano MS, Hockfield S, Matthews RT. BEHAB/brevican requires ADAMTS-mediated proteolytic cleavage to promote glioma invasion. J Neurooncol 2008; 88: 261–272.
- 53 Miller PJ, Dietz KN, Hollenbach AD. Identification of serine 205 as a site of phosphorylation on Pax3 in proliferating but not differentiating primary myoblasts. *Protein Sci* 2008; 17: 1979–1986.
- 54 Lam PY, Sublett JE, Hollenbach AD, Roussel MF. The oncogenic potential of the Pax3-FKHR fusion protein requires the Pax3 homeodomain recognition helix but not the Pax3 paired-box DNA binding domain. *Mol Cell Biol* 1999; **19**: 594–601.



Oncogenesis is an open-access journal published by Nature Publishing Group. This work is licensed under a Creative Commons Attribution 4.0

International License. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/