miR-511-3p Modulates Genetic Programs of Tumor-Associated Macrophages

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SUMMARY

Expression of the mannose receptor (MRC1/CD206) identifies macrophage subtypes, such as alternatively activated macrophages (AAMs) and M2-polarized tumor-associated macrophages (TAMs), which are endowed with tissue-remodeling, proangiogenic, and protumoral activity. However, the significance of MRC1 expression for TAM’s protumoral activity is unclear. Here, we describe and characterize miR-511-3p, an intronic microRNA (miRNA) encoded by MRC1+ AAMs and TAMs. Unexpectedly, enforced expression of miR-511-3p tuned down the protumoral gene signature of MRC1+ TAMs and inhibited tumor growth. Our findings suggest that transcriptional activation of Mrc1 in TAMs evokes a genetic program orchestrated by miR-511-3p, which limits rather than enhances their protumoral functions. Besides uncovering a role for MRC1 as gatekeeper of TAM’s protumoral genetic programs, these observations suggest that endogenous miRNAs may operate to establish thresholds for inflammatory cell activation in tumors.

INTRODUCTION

Tumor-associated macrophages (TAMs) support tumor progression in mouse models of cancer (Qian and Pollard, 2010). The protumoral functions of TAMs are thought to depend, at least in part, on their production of growth, tissue-remodeling, and immunomodulatory factors. Together, these enhance tumor cell motility and invasion, activate fibroblasts to synthesize extracellular matrix (ECM) proteins, facilitate angiogenesis, and suppress antitumor immunity (Qian and Pollard, 2010; Squadrito and De Palma, 2011; Biswas and Mantovani, 2010; Sica and Bronte, 2007). However, TAMs comprise distinct subsets, which appear to contribute differentially to tumor progression (Qian and Pollard, 2010; Squadrito and De Palma, 2011). In the mouse, high expression of the mannose receptor (MRC1/CD206) and low expression of the integrin α X (CD11c) identify a TAM subset with enhanced proangiogenic, tissue-remodeling and protumoral activities (Pucci et al., 2009; Movahedi et al., 2010); a variable proportion of these MRC1+CD11c+ TAMs also express the angiopoietin receptor, TIE2, and have thus been termed TIE2-expressing macrophages (De Palma et al., 2005; Pucci et al., 2009; Mazzieri et al., 2011). Conversely, CD11c+MRC1low TAMs express a proinflammatory and angiostatic phenotype, and perhaps exert antitumoral functions (Pucci et al., 2009; Movahedi et al., 2010; Rolny et al., 2011). It is still unclear whether the diverse TAM subsets identified in mouse tumor models derive from distinct circulating monocyte precursors or are induced locally in the tumor from a common precursor/progenitor cell (PC) (Squadrito and De Palma, 2011). Yet, several tumor-derived factors, including cytokines produced by infiltrating immune cells, may instruct TAMs to acquire either pro- or antitumoral functions (DeNardo et al., 2010).

MRC1 is an endocytic receptor primarily expressed by subsets of macrophages and dendritic cells (DCs); it is primarily involved in the clearance of both host and microbe-derived glycoproteins (Taylor et al., 2005). MRC1 expression is strongly upregulated by IL-4 and IL-13, and downregulated by IFN-γ (Stein et al., 1992). Notably, these cytokines are pleiotropic modulators of macrophage activation; their context-dependent expression patterns may contribute to the remarkable heterogeneity of macrophage phenotypes observed throughout tissues and tumors. Whereas IFN-γ promotes a “classic” or proinflammatory macrophage activation program, IL-4 and IL-13 fuel an “alternative” macrophage activation program, which promotes ECM remodeling, angiogenesis, tissue growth, and repair (Gordon and Martinez, 2010; Martinez et al., 2009). Based on their gene expression signature and tissue-remodeling/proangiogenic activities, MRC1+ TAMs resemble IL-4-stimulated...
Figure 1. miR-511-3p Is the Active Strand of Mouse Pre-miR-511 and Is Coregulated with the Mrc1 Gene

(A) Genomic region comprising the mouse miR-511 locus and the surrounding Mrc1 gene on mouse chromosome 2, as retrieved by the UCSC (NCBI37/mm9) genome browser.

(B) Stem-loop structure of the mouse pre-miR-511. miR-511-5p and -3p sequences are shown in blue and red, respectively.

(C) Schematic of the proviral LV used to measure miR-511 activity (miRT-511 LV). The miRT sequences are cloned downstream to the GFP expression cassette, which is regulated by a bidirectional PGK promoter.

(D) Schematic of the proviral LV used to overexpress miR-511 (SFFV.miR-511 LV). The sequence of the primary miR-511 is cloned within the EF1α intron, downstream to a SFFV promoter.
alternatively activated macrophages (AAMs) (Biswas and Mantovani, 2010; Gordon and Martinez, 2010). Although IL-4, IFN-γ, and several other tumor-derived cytokines and growth factors have been identified that can modulate macrophage phenotypes in vitro and in vivo (Biswas and Mantovani, 2010; DeNardo et al., 2010; Qian and Pollard, 2010), the signals in tumors that regulate pro- versus antitumoral functions of the distinct TAM subsets are still poorly defined.

MicroRNAs (miRNAs) are small, single-stranded RNAs that are generated from endogenous hairpin-shaped transcripts (called primary miRNAs). It is now well established that the unique combination of miRNAs expressed in each cell type determines the fine tuning of hundreds of mRNAs, thus regulating gene expression and cell function (Bartel, 2009). Several miRNAs have been identified that are robustly expressed by human macrophages in vitro (Tserel et al., 2011). However, to our knowledge, no information is available on the miRNA expression profiles of the distinct TAM subsets. Here, we describe and characterize an intronic miRNA, miR-511-3p, which is embedded in and coexpressed with the Mrc1 gene. We show that the upregulation of MRC1, which is contextual with the differentiation (or alternative activation) of protumoral TAMs, triggers a negative-feedback response orchestrated by miR-511-3p that attenuates their protumoral genetic programs.

RESULTS

miR-511-3p Is the Active Strand of miR-511

We noted that the mouse Mrc1 gene, which is primarily expressed by protumoral TAMs (Pucci et al., 2009) and AAMs (Stein et al., 1992), contains a miRNA coding sequence, miR-511 (or mmu-miR-511), located in the fifth intron of the gene (Figure 1A). Processing of the precursor miRNA (termed pre-miR-511) should generate two mature miRNAs, miR-511-5p (located at the 5′ end of the pre-miRNA) and miR-511-3p (located at the 3′ end of the pre-miRNA) (Figure 1B).

To investigate whether miR-511-5p and -3p are expressed and biologically active in live cells, we used a lentiviral vector (LV) reporter system for miRNA activity (Brown et al., 2007). We incorporated four miRNA target (miRT) sequences with perfect complementarity to either miR-511-5p or -3p (termed miRT-511-5p and miRT-511-3p, respectively) into the 3′ untranslated region (UTR) of a green fluorescent protein (GFP) transgene expressed from a ubiquitously active bidirectional promoter, which also controls the expression of the reporter gene, ΔLNGFR (Figure 1C). We also generated a control LV expressing a GFP sequence not containing miRT sequences in its 3′ UTR (termed no-miRT). Following LV cell transduction the miRNA machinery will degrade the miRT-containing GFP transcript only in cells that express the cognate miRNA, in a manner that is dependent on miRNA abundance and/or activity. On the other hand, expression of ΔLNGFR is independent of miRNA activity and is used as an internal normalizer to calculate GFP repression by the miRNA of interest (Brown et al., 2007).

We initially studied miR-511 activity in 293T cells, which do not express miR-511 endogenously (data not shown). In order to artificially overexpress the pre-miR-511 (and thus both miR-511-5p and -3p mature miRNAs), we cloned a fragment of the Mrc1 intron encompassing the miR-511 locus, downstream to the spleen focus-forming virus (SFFV) promoter and upstream to an orange fluorescent protein (OFP) reporter gene (Figure 1D). We termed the resultant vector SFFV.miR-511 LV. We then transduced 293T cells with the miRT-511-5p, -3p, or no-miRT reporter LVs, and superinfected the transduced cells with the SFFV.miR-511 LV. As shown in Figure 1E, overexpression of pre-miR-511 repressed GFP much more efficiently in cells transduced with the miR-511-3p reporter LV, suggesting that the active strand of the pre-miR-511 is miR-511-3p.

The Mrc1 Gene and miR-511-3p Are Coregulated

Intronic miRNAs can be expressed from either host gene promoters or independent transcription regulatory elements (Baskerville and Bartel, 2005; Biasiolo et al., 2011). We then asked whether expression of the Mrc1 gene and miR-511-3p are transcriptionally coregulated. To address this question, we used mouse monocytic cell lines (RAW264.7 and P388D1; see Figure S1 available online) and bone marrow-derived macrophage (BMDM) cultures. qPCR analyses showed decreasing Mrc1 mRNA levels in the following cell cultures: IL-4-treated BMDMs; untreated BMDMs; P388D1 cells; LPS/IFN-γ-treated BMDMs; and RAW264.7 cells (Figure 1F). These data are consistent with previous reports showing that IL-4 and LPS/IFN-γ
robustly modulate MRC1 expression in cultured BMDMs (Stein et al., 1992). Interestingly, we found that the degree of GFP repression tightly correlated with the expression of the endogenous Mrc1 gene in cells transduced with the miRT-511-3p but not -5p reporter LV (Figures 1F and 1G; p < 0.02 by Spearman’s rank correlation test). As seen in overexpression experiments (Figure 1E), endogenous miR-511 repressed GFP much more efficiently in cells transduced with the miRT-511-3p than -5p reporter LV (Figure 1G). Finally, we found that GFP repression by miR-511-3p (but not -5p) also correlated with MRC1 protein levels (Figures 1H–1J). Together, these data corroborate the notion that miRT-511-3p is the active strand of miR-511, and strongly suggest that the Mrc1 gene and miR-511 are transcriptionally coregulated.

Robust and Preferential Activity of miR-511-3p in MRC1+ TAMs

In order to analyze the expression pattern and activity of miR-511 in vivo, we implemented the aforementioned reporter system in a model of hematopoietic stem (HS)/PC transplantation. We transduced HS/PCs obtained from the BM of C57BL/6 mice with the miRT-511-5p, -3p, or no-miRT reporter LV, and transplanted the transduced cells into irradiated, syngenic mice. Lewis lung carcinoma (LLC) cells were injected subcutaneously 4 weeks after the transplant, and the tumors were grown for an additional 4 weeks. We then measured the degree of GFP repression in a variety of circulating and tumor-infiltrating myeloid cells, including circulating “resident” and “inflammatory” monocytes, circulating granulocytes, F4/80+MRC1+CD11clow and F4/80+CD11c+MRC1low TAMs, and tumor-infiltrating granulocytes/imature myeloid cells (iMCs). It should be noted that the majority of TAMs are MRC1+ in LLCs grown in C57BL/6 mice (Figure S2A).

We did not detect GFP repression in blood cells (granulocytes, inflammatory and resident monocytes) or tumor-infiltrating granulocytes/iMCs, indicating that neither miR-511-3p nor -5p are detectably active in these cells (Figure 2A). Conversely, we detected GFP repression in MRC1+ and, to a lesser extent, CD11c+ TAMs carrying miRT-511-3p but not -5p target sequences (Figure 2A). These in vivo data confirm that miR-511-3p is the active strand of the mouse pre-miR-511, and demonstrate that endogenous miR-511-3p is preferentially active in MRC1+ TAMs among tumor-infiltrating and circulating myeloid cells.
To rule out that our findings are mouse strain or tumor specific, we also analyzed miR-511-3p activity in FVB/n mice transplanted as above and challenged with N202 (Neo+) mammary carcinomas (Figure 2B). It should be noted that, contrary to LLCs, the majority of TAMs are CD11c+ in N202 tumors grown in FVB/n mice (Figure S2B). As seen in LLCs, we observed robust GFP repression (and thus miR-511-3p activity) in MRC1+ but not CD11c+ TAMs or infiltrating granulocytes/iMCs in N202 tumors analyzed at 4 weeks postinjection. These data were confirmed by immunofluorescence staining of tumor sections, showing lower GFP signal in MRC1+ than CD11c+ TAMs (Figure 2C). These findings indicate that preferential activity of miR-511-3p in MRC1+ TAMs is independent of the mouse strain, the tumor type, or the relative abundance of the distinct TAM subsets.

In addition to miRNA activity, we analyzed miRNA abundance by qPCR. We measured the expression of a panel of selected miRNAs, including miR-511-5p and -3p, in both MRC1+ and CD11c+ TAMs isolated from spontaneous MMTV-PyMT mammary tumors (Mazzieri et al., 2011) by fluorescence-activated cell sorting (FACS). Although both miR-511-5p and -3p were significantly upregulated in MRC1+ versus CD11c+ TAMs (>10-fold), miR-511-3p levels were much higher than -5p levels in either TAM subset (Figure 2D). Of note, expression of the Mrc1 gene was ~10-fold higher in MRC1+ than CD11c+ TAMs (Figure 2E), suggesting that in vivo as in vitro (Figures 1G and 1J) the host gene and the miRNA are transcriptionally coregulated.

miR-511-3p Is Preferentially Active in MRC1+ Tissue Macrophages

In addition to protumoral TAMs (Qian and Pollard, 2010; Squadrito and De Palma, 2011), certain tissue-resident macrophage populations express MRC1 (Gordon and Martinez, 2010; Martinez et al., 2009). We then asked whether miR-511-3p is also active in these cells. To this aim, we analyzed organs and tissues of FVB/n mice transplanted 8 weeks earlier with HS/PCs transduced with the miRT-511-5p or no-miRT reporter LVs (Figures 3A–3D). In agreement with the pattern of miR-511-3p activity in tumor-infiltrating myeloid cells, we detected GFP repression (and hence miR-511-3p activity) specifically in F4/80+Gr1− macrophages that express distinctly high MRC1 protein. These include MRC1+CD11c− adipose tissue macrophages (Chawla et al., 2011), MRC1+ lung/alveolar macrophages (Landsman and Jung, 2007), MRC1+CD11c− spleen red pulp macrophages, and MRC1−CD11c− liver Kupffer cells (Taylor et al., 2005). On the other hand, Gr1+ F4/80− granulocytes, CD11c+Gr1−/F4/80− DCs, and other MRC1-negative myeloid cells all displayed negligible miR-511-3p activity (Figures 3A–3D).

To corroborate these findings with miRNA expression data, we isolated macrophages from the adipose tissue by FACS, and measured the expression of a panel of selected miRNAs, including miR-511-5p and -3p. In agreement with the GFP repression data, we observed significantly higher miR-511-3p levels in MRC1+CD11c− than CD11c+MRC1− adipose tissue macrophages (Figure 3E). As seen in TAMs (Figure 2E), miR-511-3p levels were consistently higher than -5p levels in each macrophage subset (Figure 3E). Of note, abundance of the Mrc1 mRNA correlated with that of miR-511-3p (Figure 3F), further supporting the notion that the host gene and the miRNA are transcriptionally coregulated. Taken together, these data demonstrate robust miR-511-3p expression and activity in distinct MRC1+ tissue-macrophage subtypes.

**ROCK2 Is a Direct Target of miR-511-3p**

We then used TargetScan (Lewis et al., 2005) and DIANA microT (Maragkakis et al., 2009) to identify miR-511-3p predicted targets. The analysis retrieved a list of 145 genes (Table S1) that we analyzed by DAVID Bioinformatic resources 6.7 (Huang et al., 2009). We found that a significant proportion of these genes are involved in biological processes related to “cell morphogenesis” (Table S2).

To validate miR-511-3p predicted targets, we first generated a mutant miR-511-3p sequence by substituting four nucleotides in the pre-miR-511 sequence of the SFFV.miR-511 LV. Three out of four substitutions are located in the seed sequence of miR-511-3p, and were selected to not modify the complementary miR-511-5p sequence and to not perturb the stem-loop structure of the pre-miRNA (Figure S3A). We termed the resultant vector SFFV.miR-511-mut LV. To validate the mutant sequence, we transduced RAW264.7 monocytic cells with the miRT-511-5p or -3p reporter LV, and superinfected the transduced cells with the SFFV.miR-511 or -511-mut LV. As shown in Figure 4A, the four mutated nucleotides in the miR-511-3p sequence completely abrogated its activity.

We then performed dual-luciferase assays to test the 3′ UTRs of a small panel of miR-511-3p predicted targets, including Rho-dependent kinase-2 (Rock2), a serine/threonine kinase that regulates cell’s cytoskeleton contractility (Samuel et al., 2011). We first transduced RAW264.7 cells with either SFFV.miR-511 or SFFV.miR-511-mut LV and, 1 week later, transfected the dual-luciferase constructs in the transduced cells. We observed robust Rock2-UTR-dependent repression of luciferase activity in SFFV.miR-511-overexpressing cells, but not SFFV.miR-511-mut-overexpressing cells (Figure 4B). We further validated miR-511-3p/Rock2-UTR interaction by testing the Rock2 3′ UTR (as well as a mutated sequence; Figure S3B) in an in vitro GFP repression assay based on our LV reporter system (Figure 4C; Extended Experimental Procedures). By this approach, we confirmed direct interaction between miR-511-3p and the Rock2 3′ UTR.

Finally, we analyzed the expression of ROCK2 in RAW264.7 cells, P388D1 cells, and BMDMs engineered to overexpress either miR-511 or miR-511-mut. miR-511-3p downregulated ROCK2 both at the mRNA (Figures 4D and 4F) and protein (Figures 4E, 4G, and 4H) level. Taken together, these data demonstrate that ROCK2 is a direct target of mouse miR-511-3p.

**The Human MRC1 Gene Encodes for an Active miR-511-3p Sequence**

The human MRC1 gene contains a miR-511 sequence (hsa-miR-511) located in the fifth intron of the gene; of note, hsa-miR-511-3p is a miRNA, currently not annotated in miRBase (http://www.mirbase.org). The mature hsa-miR-511-3p but not -5p sequence is conserved in M. musculus and H. sapiens (Figure S4A). We then asked whether miR-511-3p activity is conserved in the two species. To identify the active strand of...
Figure 3. Preferential Expression and Activity of miR-511-3p in MRC1+ Tissue-Resident Macrophages

(A–D) Flow cytometry analysis of F4/80+ tissue-resident macrophages (further separated based on MRC1 and CD11c expression), Gr1+F4/80− granulocytes, and CD11c+F4/80− DCs, in the indicated tissues/organs of FVB/n mice. Flow cytometry dot plots on the left show the gating strategy. The scatterplot on the right shows GFP repression (versus no-miRT control) in the individual cell types. Each dot in the scatterplot corresponds to one mouse. Statistical analysis of fold-repression values was performed by unpaired Student’s t test.
the human miR-511, we generated both reporter and overexpressing LVs (Figure S4B), as described above for the mouse miR-511. We transduced human U937 monocytic cells with the reporter LVs and then superinfected the transduced cells with the overexpressing LVs. Analysis of GFP repression showed that, as in the mouse system, miR-511-3p is the active strand of the human pre-miR-511 (Figure SA).

As in the mouse system, predicted targets of human miR-511-3p (Table S3) comprise genes involved in biological processes related to “cell morphogenesis” (Table S4). Overexpression of human miR-511-3p decreased ROCK2 protein in U937 cells (Figure 5B), suggesting that ROCK2 is a direct target of human miR-511-3p, as predicted by TargetRank (Nielsen et al., 2007). We confirmed this finding by analyzing human miR-511-3p/ROCK2-UTR interaction in an in vitro GFP repression assay based on our reporter LV system (Figure 5C; Extended Experimental Procedures). Together, these data strongly suggest that miR-511-3p activity is conserved in mice and humans.

Overexpression of miR-511-3p in BM-Derived Cells Inhibits Tumor Growth and Alters Tumor Blood Vessel Morphology

To study the biological function of mouse miR-511-3p, we overexpressed it in BM-derived hematopoietic cells. To this aim, we transduced HS/PCs obtained from CD45.1/C57BL/6 mice with either SFFV.miR-511 or -511-mut LV, and transplanted the transduced cells into irradiated, congenic CD45.2/C57BL/6 mice, to obtain SFFV.miR-511 and SFFV.miR-511-mut mice, respectively. Four weeks after the transplant, we inoculated LLC cells subcutaneously in the transplanted mice and monitored tumor growth for 3–4 weeks.

Unsurprisingly, miR-511-3p overexpression in hematopoietic cells inhibited LLC growth (Figure 6A). We could reasonably exclude that tumor growth inhibition by miR-511-3p overexpression was due to defective hematopoiesis and/or altered recruitment of hematopoietic cells to the tumors. Indeed, miR-511-3p overexpression in hematopoietic cells did not affect the repopulating activity of the transduced HS/PCs, as shown by the similarly high frequency of CD45.1*OFP*, donor-transduced hematopoietic cells in the blood of both groups of mice (Figure S5A). Furthermore, miR-511-3p overexpression neither affected the recruitment of F4/80+ TAMs (which represent up to 60% of all tumor-infiltrating hematopoietic cells in this tumor model), Gr1+ neutrophils, NK, T and B cells to the tumors (Figure S5B), nor the relative frequency of MRC1+ and CD11c+ TAM subsets (Figure S5C).

We then asked whether miR-511-3p overexpression influenced tumor angiogenesis. Although we did not observe changes in vascular density by immunofluorescence staining of tumor sections (data not shown), we noted that miR-511-3p overexpression altered the architecture of the tumor microvascular network by augmenting blood vessel tortuosity and the occurrence of enlarged, glomerular-like structures (Figures 6B and 6C). Accordingly, morphometric analysis of thick tumor sections showed similar vascular area but significantly reduced total and mean length of blood vessels in SFFV.miR-511 compared with -511-mut mice (Figure 6D). Together, these data indicate that miR-511-3p overexpression in BM-derived cells inhibits tumor growth and dysregulates angiogenesis without affecting hematopoiesis detectably.

Overexpression of miR-511-3p in TAMs Globally Downregulates miR-511-3p Predicted Target Genes

Because the genetic programs of TAMs may influence tumor angiogenesis and growth (Qian and Pollard, 2010), we asked whether miR-511-3p overexpression had modulated TAM’s gene expression. To address this question, we sorted F4/80+OFP+ TAMs from LLCs grown in SFFV.miR-511 or -511-mut mice. qPCR analysis of selected miRNAs showed that the SFFV.miR-511 LV upregulated the expression of miR-511-3p by ~5-fold in the F4/80+OFP+ TAMs of SFFV.miR-511 compared to -511-mut mice, which only express the endogenous miR-511-3p sequence (Figure 7A). Of note, miR-511-3p-mut was only detected in the TAMs of SFFV.miR-511-mut mice, whereas miR-511-5p was expressed by both SFFV.miR-511 and SFFV.miR-511-mut TAMs. However, as seen in the TAMs of MMTV-PyMT mice (Figure 2D), expression of miR-511-5p was much lower than miR-511-3p, strongly suggesting that, even when overexpressed, it is rapidly degraded in vivo.

We then performed RNA-Seq analyses of the transcriptome of sorted F4/80+OFP+ TAMs. We used the Illumina HiSeq 2000 platform and retrieved 249 genes (out of 16,355; 1.5%; p < 0.05 adjusted for false discovery rate) that were differentially expressed in the TAMs of SFFV.miR-511 versus -511-mut mice (Table S5). Remarkably, it was found that the predicted targets of miR-511-3p (Table S1) were globally downregulated by miR-511-3p overexpression in TAMs (Figure 7B). We also used TargetRank to identify transcripts that contain in their 3’ UTR at least one sequence with perfect complementarity to the seed sequence of miR-511-3p (i.e., M8-A1 8-mer and M8 7-mer binding sites; Table S6), and found that such transcripts were globally downregulated by miR-511-3p overexpression (Figure 7C; see Extended Experimental Procedures). Conversely, genes containing M8-A1 8-mer or M8 7-mer binding sites for either miR-511-5p or -3p-mut were significantly less downregulated by miRNA overexpression (Figures 7D and 7E). These data demonstrate broad and robust miR-511-3p activity in TAMs by our overexpression platform.

Overexpression of miR-511-3p Tunes down the Protumoral Gene Signature of MRC1+ TAMs

Although the vast majority of the differentially expressed genes were downregulated by miR-511-3p overexpression in TAMs (Table S5), they could not be identified as miR-511-3p direct
Figure 4. ROCK2 Is a Direct Target of Mouse miR-511-3p

(A) GFP repression in RAW264.7 cells transduced with the miR-511-5p or -3p LVs and superinfected with the miR-511 or -511-mut overexpressing LVs (mean values ±SEM versus untransduced [UT] cells; n = 3 independent experiments). Statistical analysis of fold-repression values was performed by two-way ANOVA with Bonferroni posttest.

(B) Firefly luciferase activity in 293T cells untransduced or transduced with either miR-511 or -511-mut LV. The 3' UTRs of mouse podoplanin (Pdpn), semaphorin-3A (Sema3a), Rock2 (all miR-511-3p target genes), and CD163 were tested, together with a UTR-less plasmid (miRGLO). The Rock2 UTR was split into two fragments (Rock2(1) and Rock2(2)). The box-and-whisker graph shows luciferase activity (median ± minimum/maximum values versus miRGLO; n = 6–9 technical replicates from 3 independent experiments). Statistical analysis was performed by two-way ANOVA with Bonferroni posttest.

(C) GFP repression in P388D1 cells transduced with GFP reporter LVs containing either wild-type or mutant UTR sequences from the Rock2 gene. Cells were superinfected with either miR-511 or -511-mut overexpressing LV. Data show fold change of GFP repression (mean ± SEM; n = 3 independent experiments). Statistical analysis was performed by two-way ANOVA with Bonferroni posttest.

(D) qPCR of Rock2 expression in RAW264.7 cells overexpressing either miR-511 or -511-mut. The data show fold change (= 2\(^{-\Delta\Delta C_T}\); mean values ±SEM; n = 3 biological samples) versus untransduced cells (reference population). Normalization was performed by \(b_2m\). Statistical analysis of \(\Delta C_T\) values was performed by unpaired Student’s t test.

(E) Western blot analysis of ROCK2 in RAW264.7 cells overexpressing either miR-511 or -511-mut. A representative blot is shown on the right.

(F) qPCR of Rock2 expression in P388D1 cells overexpressing either miR-511 or -511-mut. The data show fold change (= 2\(^{-\Delta\Delta C_T}\); mean values ±SEM; n = 3 biological samples) versus untransduced cells (reference population). Normalization was performed by Hprt. Statistical analysis of \(\Delta C_T\) values was performed by unpaired Student’s t test.

(G and H) Western blot analysis of ROCK2 in P388D1 cells (G) or BMDMs (H) overexpressing either miR-511 or -511-mut. Analysis as in (E) (P388D1: nine technical replicates, three independent experiments; BMDMs: four technical replicates, two independent experiments).
targets by TargetScan, DIANA microT, and TargetRank, and possibly represent indirect targets of the miRNA. Interestingly, the downregulated genes are primarily involved in biological processes related to cell adhesion, morphogenesis, and ECM organization (Table S7). They comprise genes encoding for ECM proteins, such as collagens (e.g., type VI collagens), basal lamina proteins, and proteoglycans. Downregulated genes also include genes that control the synthesis and remodeling of the ECM, such as proteases (e.g., Adamst1, Adamst11, Mmp11, Mmp3), scavenger receptors (e.g., Sparc and Mrc2), and TGF-β family (e.g., Tgfbr3, Bmp1, Bmpr1a, Bmpr1b) or associated (Ltbp1) factors. Latent TGF-β binding protein-1 (LTBP1) is a secreted protein that has a role in the assembly, secretion, and activation of latent complexes of TGF-β in the ECM; by activating TGF-β, LTBP1 may stimulate ECM biosynthesis and enhance tumor growth (Saunier and Akhurst, 2006). Consistent with RNA-Seq analysis of TAMs (Table S5), miR-511-3p overexpression in P388D1 cells and BMDMs decreased expression of LTBP1 at the mRNA (Figure 7F) and protein (Figure 7G) level, respectively.

We and others previously showed that MRC1+ TAMs express a distinguishing gene signature and are protumoral in mouse models of cancer; genes upregulated in MRC1+ TAMs may thus identify the protumoral gene signature of TAMs (Pucci et al., 2009; Movahedi et al., 2010). We then hypothesized that miR-511-3p overexpression inhibited tumor growth by attenuating the protumoral genetic programs of MRC1+ TAMs. To test this hypothesis, we first sorted MRC1+ and CD11c+ TAMs from LLC tumors grown for 4 weeks in wild-type, nontransplanted C57BL/6 mice, and subjected the isolated cells to RNA-Seq analysis. About 14% of the identified genes were differentially expressed between MRC1+ and CD11c+ TAMs (p < 0.05 adjusted for false discovery rate; Table S8), corroborating the notion that MRC1+ and CD11c+ TAMs represent distinct cell subsets (Pucci et al., 2009). Of note, many of the genes upregulated in MRC1+ versus CD11c+ TAMs encode for molecules with previously established protumoral bioeffectors.
strategy to stably overexpress miRNAs in BM-derived cells, we found that miR-511-3p broadly and specifically attenuates the expression of genes that define the protumoral signature of MRC1+ TAMs (Pucci et al., 2009; Movahedi et al., 2010). Consistent with this finding, miR-511-3p overexpression inhibited tumor growth. On the other hand, miR-511-3p overexpression did not alter the proinflammatory gene signature of CD11c+ macrophages (Pucci et al., 2009; Movahedi et al., 2010), suggesting specific activity of the miRNA in a TAM subtype. Interestingly, miR-511-3p was most biologically active in tissue-resident macrophages bearing features of AAMs (Chawla et al., 2011; Gordon and Martinez, 2010; Landsman and Jung, 2007; Martinez et al., 2009). These cells are known to participate in both pathological and physiological processes, including host defense from parasites, stimulation of angiogenesis and tissue repair, promotion of tissue fibrosis, and regulation of organ metabolism (Gordon and Martinez, 2010). Future studies are now needed to address the significance of miR-511-3p in the regulation of alternative activation of macrophages.

Interestingly, miR-511-3p downregulated TAM expression of multiple genes involved in ECM synthesis and remodeling; these include collagens and other fibrous proteins, proteases, and scavenger receptors. Of note, the composition and biophysical properties of the ECM influence tumor growth and progression. Increased collagen deposition/crosslinking and ECM stiffening stimulate tumor cell proliferation, invasion, and malignancy (Egeblad et al., 2010; Levental et al., 2009). Furthermore, the composition and biophysical properties of the ECM regulate vascular morphogenesis in tumors (Bauer et al., 2009). Indeed, ECM density controls the extension speed of vascular sprouts, and a high matrix-fiber anisotropy (i.e., directional tension) provides strong contact guidance cues for endothelial cells and stimulates sprout branching (Bauer et al., 2009). Although ECM fibrous proteins are mainly produced by fibroblasts and epithelial cells (Egeblad et al., 2010; Kalluri and Zeisberg, 2006), there is also evidence that some collagens and other ECM proteins may be robustly expressed by in vitro-cultured macrophages (Schnoor et al., 2008). Yet, the significance of TAM-produced ECM fibrous proteins for tumor growth and vascularization has remained largely unexplored. Our deep sequencing analyses indicate that MRC1+ TAMs express several ECM genes (including genes encoding for...
Figure 7. miR-511-3p Overexpression in TAMs Tunes Down Their Protumoral Gene Signature

(A) qPCR of selected miRNAs in F4/80+OFP+ TAMs isolated from LLCs grown in mice overexpressing either miR-511 or -511-mut in hematopoietic cells. The data show the relative abundance of each miRNA (mean values ± SEM versus Let7a; n = 4 biological samples). Statistical analysis of ΔCt values was performed by unpaired Student’s t test.

(B) Cumulative distribution of fold changes in the whole transcriptome (13,747 genes; transcripts with less than 10 reads, and miR-511-3p predicted targets were excluded from the analysis) of TAMs overexpressing miR-511 (versus -511-mut; red line). The green line shows the cumulative distribution of fold changes in transcripts that are miR-511-3p predicted targets (145 genes). Note the global repression of miR-511-3p target genes. Statistical analysis was performed by one-sided Kolmogorov-Smirnov test.

(C–E) Cumulative distribution of fold changes in the whole transcriptome (13,747 genes; transcripts with less than 10 reads, and miR-511-3p predicted targets were excluded) of TAMs overexpressing miR-511 (versus -511-mut; red line). The green and blue lines show the cumulative distribution of fold changes in transcripts that contain M8-A1 8-mer and M8 7-mer target sites, respectively, for miR-511-3p (C), -5p (D), or -3p-mut (E). Statistical analysis was performed by one-sided Kolmogorov-Smirnov test.

(F) P388D1 Ltb1 mRNA

(G) BMDMs LTBP1 protein

(H) MRC1+ TAMs

(I) CD11c+ TAM gene signature

(J) LLC Rock2 mRNA

(K) BMDMs Rock2 mRNA
collagens and other fibrous proteins), which were globally and significantly downregulated by miR-511-3p overexpression in TAMs. Because MRC1+ TAMs represent a major component of the perivascular tumor stroma and support vascular morphogenesis in tumors (Mazzieri et al., 2011; Squadrito and De Palma, 2011), modulation of ECM-protein synthesis/remodeling by miR-511-3p in MRC1+ TAMs may have the potential to influence ECM dynamics in the perivascular microenvironment. This would be consistent with our finding that miR-511-3p overexpression in TAMs altered the morphology of intratumoral blood vessels, possibly as a consequence of changes in the biophysical properties of the perivascular ECM (and/or in the levels of TAM-derived angiogenic factors). It is also possible that miR-511-3p is horizontally transferred from MRC1+ TAMs to other tumor-associated stromal cells via microvesicles or exosomes (Yang et al., 2011).

We identified ROCK2 as a direct target of miR-511-3p. Our data indicate that MRC1+ TAMs as well as IL-4-polarized BMDMs express higher ROCK2 mRNA levels than CD11c+MRC1− TAMs or unstimulated BMDMs. Of note, constitutive ROCK activation in epithelial cells increases collagen synthesis and tissue stiffness (Samuel et al., 2011). It is tempting to speculate that increased ROCK activity in MRC1+ TAMs might enhance their expression and secretion of ECM proteins as part of their protumoral genetic program. Because miR-511-3p downregulated ROCK2 both at the mRNA and protein level, our data suggest that miR-511-3p might negatively regulate ROCK activity in MRC1+ TAMs and downregulate their expression of ECM genes relevant to tumor progression.

Although several miRNAs have been identified that regulate immune cell functions (O’Connell et al., 2010), little is known of their roles in the modulation of TAM heterogeneity and functions. A recent report used Illumina miRNA Chips to analyze the miRNA expression signature of human monocytes and monocyte-derived DCs/macrophages (Tserel et al., 2011). Several miRNAs were found to be differentially expressed between DCs/macrophages and freshly isolated monocytes. Among these, human miR-511 (representing the hsa-miR-511-5p sequence described in our study) was highly upregulated in DCs/macrophages (Tserel et al., 2011). Although the report of Tserel and colleagues may appear consistent with our findings, we did not detect significant activity of either human or mouse miR-511-5p in several independent cell assays, both in vitro and in vivo. Conversely, we identified miR-511-3p as the bioactive strand of both human and mouse pre-miR-511. It should be noted, however, that the human miR-511-3p sequence is not annotated in miRBase and was not assayed in the Illumina miRNA Chips employed by Tserel and colleagues, so its differential expression could not be assessed (Tserel et al., 2011). Although the mechanisms that regulate miRNA strand selection are still unclear, it is likely that the thermodynamic stability of the two ends of the pre-miRNA determines which strand is to be selected for loading into the RISC complex, and which is to be degraded (Khvorova et al., 2003). Recent studies have also illustrated that miRNA strand selection may be cell type, context, and species specific (Biasiolo et al., 2011; Kuchenbauer et al., 2011). Nevertheless, our analyses indicate that miR-511-3p but not -5p is highly conserved across mammalian species, pointing to a preeminent role of this miRNA strand. Further studies are, therefore, needed to clarify the significance of miR-511-5p expression and activity in macrophages.

In summary our findings reveal an unexpected layer of gene expression control in TAMs, which relies on an endogenous molecular switch that is activated in a tumor-promoting (MRC1+) subset of these cells. Enhancing miR-511-3p activity in TAMs (e.g., via delivery of macrophage-targeted miRNA carriers) may represent a therapeutic strategy to reprogram them from a protumoral to an antitumoral phenotype.

**EXPERIMENTAL PROCEDURES**

Detailed experimental procedures are presented as Extended Experimental Procedures.

**Mice**

C57BL/6, CD45.1/C57BL/6, and FVB/n mice were purchased by Charles River Laboratory (Calco, Milan). FVB/MVMT-PyMT mice were obtained from the NCI-Frederick Mouse Repository (Frederick, MD) and established as a colony at the San Raffaele animal facility. All procedures were performed according to protocols approved by the Animal Care and Use Committee of the Fondazione San Raffaele del Monte Tabor (IACUC 324, 335, and 447).
and communicated to the Ministry of Health and local authorities according to the Italian law.

LV Transduction
For experiments in vitro, cells were transduced with LV doses ranging from $10^4$ to $10^8$ transducing units (TU)/ml. When required, sequential transduction was performed by (i) transducing the cells with the first LV; (ii) washing and replating the cells; and (iii) transducing the cells with the second LV (superinfection) on day 5–7 after the first transduction. For HS/PC transplantation, $10^8$ HS/PCs/ml were prestimulated for 6 hr in serum-free medium containing a cocktail of cytokines, and then transduced with miRT reporter or miR-511-overexpressing LVs with a dose equivalent to $10^6$ TU/ml. After transduction, $10^6$ cells were infused into the tail vein of lethally irradiated mice.

Tumor Experiments
LLC/3LL cells (5 $\times$ $10^6$) were injected subcutaneously in syngenic C57BL/6 mice, and tumors were grown for 3–4 weeks; tumor size was determined by caliper measurements. N202 mammary carcinoma cells (5 $\times$ $10^6$) were injected subcutaneously in syngenic FVB/n mice, and tumors were grown for 4 weeks.

For miRT reporter studies we performed two independent experiments. In the first experiment, transduced HS/PCs were transplanted in irradiated C57BL/6 mice, which were subsequently challenged with LLC cells. In the second experiment, transduced HS/PCs were transplanted in irradiated FVB/n mice, subsequently challenged with N202 cells. For miR overexpression studies we performed three independent experiments. In each experiment, transduced HS/PCs were transplanted in irradiated C57BL/6 mice, subsequently challenged with LLC cells; tumor growth was analyzed for 3–4 weeks in the first two experiments. In the first experiment, tumors were also harvested for sorting of TAMs and RNA-Seq analysis. In the second experiment, mice were randomly selected for Microfill perfusion and analysis of the tumor-associated vasculature. In the third experiment, tumors were harvested for sorting of TAMs and qPCR of miRNAs.

Calculation of miRNA Activity
In most of the experiments, we calculated miR-511-mediated GFP repression (indicated as “fold-repression”) in live cells by using the following equation:

$$ \frac{[\text{MFI}_{\text{LNGFR}_{\text{miRT}}} \times (\text{MFI}_{\text{GFP}_{\text{miRT}}}^{-1})]}{[\text{MFI}_{\text{LNGFR}_{\text{no-miRT}}} \times (\text{MFI}_{\text{GFP}_{\text{no-miRT}}}^{-1})]} $$

where MFI is the mean fluorescence activity of either GFP or LNGFR measured by flow cytometry.

Statistical Analysis
Statistical analysis of the data is described in the figure legends and Extended Experimental Procedures. Statistical significance of the data is indicated as follows: *: $p < 0.05$; **: $p < 0.01$; ***: $p < 0.001$.

ACCESSION NUMBERS
RNA sequencing data (12 TAM samples) have been deposited in the GEO repository at NCBI under accession number GSE34903.

SUPPLEMENTAL INFORMATION
Supplemental Information includes five figures, 13 tables, and Extended Experimental Procedures and can be found with this article online at doi:10.1016/j.celrep.2011.12.005.

LICENSING INFORMATION
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EXTENDED EXPERIMENTAL PROCEDURES

Lentiviral Vector (LV) Construction and Production

Human and mouse miR-511 target (miRT) sequences were designed based on miRNA sequences obtained from the miRNA Registry (http://microrna.sanger.ac.uk/). Oligonucleotides used for generating miRT sequences are shown in Table S10. Briefly, to generate the miRT LVs, the Sense 1 (S1), Sense 2 (S2), Antisense 1 (AS1), and Antisense 2 (AS2) oligonucleotides were annealed and ligated into the 3'-UTR of the GFP gene contained in a LV co-expressing ΔLNGFR and GFP from a bidirectional PGK promoter (Amendola et al., 2005), as shown in Figure 1C.

In order to overexpress mouse and human miR-511, or their mutated forms, we designed (GeneArt Invitrogen) DNA sequences encompassing the miR-511 intronic sequence of the MRC1 gene (Table S11). Briefly, we cloned the DNA fragment into the multicloning site present in the EF1a intron of a LV containing the SFFV promoter, exon 1 and intron 1 of the EF1α gene, and the OFP reporter gene, as shown in Figure 1D.

Vesicular stomatitis virus (VSV)-pseudotyped, third-generation LVs were produced by transient four-plasmid cotransfection into 293T cells and concentrated by ultracentrifugation, as described (De Palma and Naldini, 2002). Expression titers of OFP- or ΔLNGFR-expressing LVs were determined on HeLa cells by limiting dilution. Vector particle content was measured by HIV-1 Gag p24 antigen immunocapture (NEN Life Science Products; Waltham, MA). Vector infectivity was calculated as the ratio between titer and particle content. Titer of 293T conditioned medium (before ultracentrifugation) ranged from 10⁶ to 10⁷ transducing units/ml and infectivity from 10⁴ to 10⁵ transducing units/ng of p24.

LV Transduction

293T, RAW264.7, P388D1, U937, and bone marrow derived macrophages (BMDMs) were transduced with LV doses ranging from 10⁴ to 10⁵ transducing units/ml. The fraction of ΔLNGFR⁺ (miRT reporter LVs) or OFP⁺ (overexpressing LVs) cells was greater than 80% in each experiment. When required, sequential transduction was performed by (i) transducing the cells with the first LV for 12 hr; (ii) washing and replating the cells; (iii) transducing the cells with the second LV (superinfection) on day 5-7 after the first transduction, for 12 hr in standard conditions.

Calculation of miRNA Activity

In most of the experiments, we calculated miR-511-mediated GFP repression (indicated as “fold-repression”) in live cells by using the following equation:

\[
\frac{\text{MFI}_{\text{ΔLNGFR} \text{miRT}} \times (\text{MFI}_{\text{GFP} \text{miRT}})^{-1}}{\text{MFI}_{\text{ΔLNGFR} \text{no-miRT}} \times (\text{MFI}_{\text{GFP} \text{no-miRT}})^{-1}},
\]

where MFI is the mean fluorescence activity of either GFP or ΔLNGFR measured by flow cytometry.

In some experiments, we calculated GFP repression by using the following equation:

\[
\frac{\text{MFI}_{\text{ΔLNGFR} \text{miRT}} \times (\text{MFI}_{\text{GFP} \text{miRT}})^{-1}}{\text{MFI}_{\text{ΔLNGFR} \text{miRT/UT}} \times (\text{MFI}_{\text{GFP} \text{miRT/UT}})^{-1}},
\]

where miRT/UT is the GFP or ΔLNGFR MFI of cells that contain the reporter but not the overexpressing LV.

Cell Lines

Human 293T, mouse RAW264.7, mouse P388D1 and mouse LLC cells were maintained in Iscove’s modified Dulbecco’s medium (IMDM; Sigma) supplemented with 10% fetal bovine serum (FBS; GIBCO) and a combination of penicillin-streptomycin and glutamine. Human U937 cells were maintained in RPMI supplemented as above.

Bone Marrow Derived Macrophages (BMDMs)

BM cells were obtained by flushing the femurs of 8-week old female FVB/n mice. Cells were plated in RPMI complete medium (10% FBS; penicillin-streptomycin; glutamine) supplemented with M-CSF (50 ng/ml), and cultured for one week to allow macrophage differentiation. In most experiments, bone marrow cells were transduced with reporter or overexpressing LVs on day 1 post-plating. BMDMs were then polarized by culturing them in the presence of IL4 (20 ng/ml, Peprotech) for 24-72 hr, or LPS (100 ng/ml, Sigma) + IFN-γ (200 U/ml, Peprotech) for 24-48 hr in RPMI medium supplemented with M-CSF (50 ng/ml). BMDMs for flow cytometry analysis were cultured on Petri dishes (non-tissue culture treated, bacterial grade); BMDMs for protein or mRNA extraction were cultured on regular tissue culture dishes.

For analysis of intracellular MRC1 protein, BMDMs were permeabilized using the Cytofix/Cytoperm fixation and permeabilization kit (BD Biosciences), following the manufacturer’s instructions. For immunofluorescence analysis, BM cells were transduced, differentiated into BMDMs and polarized in Chamber Slides (Lab-Tek). Cells were then fixed for 15 min in 4% paraformaldehyde and then blocked with 5% fetal bovine serum in PBS containing 1% bovine serum albumin (BSA) and 0.1% Triton X-100 (PBS-T). For immunofluorescence staining of MRC1, we used goat anti-mouse MRC1 antibodies (R&D) followed by donkey anti-goat alexa647
Actin was labeled by phalloidin-alexa546 (Invitrogen). Cell nuclei were labeled by DAPI (Molecular Probes). GFP was acquired as direct fluorescence.

**Hematopoietic Stem/Progenitor Cell (HS/PC) Isolation, Transduction and Transplantation**

Six-week old female CD45.1/C57BL/6, C57BL/6 or FVB/n mice were killed with CO2 and their BM was harvested by flushing the femurs and the tibias. Lineage-negative cells (BM-lin– cells) enriched in HS/PCs were isolated from BM using a cell purification kit (StemCell Technologies) and transduced by concentrated LVs, as described (De Palma et al., 2008). Briefly, 10^6 cells/ml were pre-stimulated for 6 hr in serum-free StemSpan medium (StemCell Technologies) containing a cocktail of cytokines (IL-3 (20 ng/ml), SCF (100 ng/ml), TPO (100 ng/ml) and FLT-3L (100 ng/ml), all from Peprotech) and then transduced with miRT reporter or miR-511-overexpressing LVs with a dose equivalent to 10^6 LV Transducing Units/ml, for 12 hr in medium containing cytokines, as described (De Palma et al., 2008). After transduction, 10^6 cells were infused into the tail vein of lethally irradiated, 6-week–old, female C57BL/6 or FVB/n mice (radiation doses: 1150 cGy split in 2 doses for C57BL/6 mice; 950 cGy split in 2 doses for FVB/n mice).

**Tumor Growth Experiments**

LLC/3LL cells (5 × 10^6) were injected subcutaneously in syngenic C57BL/6 mice, and tumors grown for 3-4 weeks. Tumor size was determined by caliper measurements, and tumor volume calculated by a rational ellipse formula (m1 × m1 × m2 × 0.5236, where m1 is the shorter axis and m2 is the longer axis), as described (De Palma et al., 2005). N202 mammary carcinoma cells (5 × 10^6) were injected subcutaneously in syngenic FVB/n mice, and tumors grown for 4 weeks.

For miRT reporter studies (Figures 2A–2C), we performed 2 independent experiments. In the first experiment, HS/PCs were transplanted in irradiated C57BL/6 mice (3 groups of mice: no-miRT, miRT-511-3p, and miRT-511-5p; n = 4-6 mice/group), which were subsequently challenged with LLC cells subcutaneously. In the second experiment, HS/PCs were transplanted in irradiated FVB/n mice (2 groups: no-miRT and miRT-511-3p; n = 5-6 mice/group), which were subsequently challenged with N202 cells subcutaneously. Tumors were allowed to grow for 4 weeks.

For miR overexpression studies, we performed 3 independent experiments. In each experiment, HS/PCs were transplanted in irradiated C57BL/6 mice (2 groups: SFFV.miR-511 and SFFV.miR-511-mut), which were subsequently challenged with LLC cells subcutaneously. Tumor growth was monitored for 3-4 weeks in two out of three experiments. In the first experiment (n = 4-8 mice/group), tumors were harvested at the end of the experiment for cell sorting of TAMs and RNA-Seq analysis. In the second experiment (n = 11 mice/group), 5 mice/group were randomly selected for Microfill perfusion and analysis of the tumor-associated vasculature. Tumors injected in mice from the third experiment (n = 4 mice/group) were harvested at 3 weeks post-injection and processed for cell sorting and qPCR of miRNAs.

**Flow Cytometry of Blood Leukocytes, Tumor- and Organ-Derived Cells**

All cell suspensions were incubated with rat anti-mouse FcγIII/II receptor (CD16/CD32) blocking antibodies (4 μg/ml; BD Inc.) together with the antibodies listed in Table S12. After antibody staining, the cells were washed, stained with fluorochrome-labeled streptavidin (if required) and re-suspended in 7-amino-actinomycin D (7-AAD)-containing buffer, to exclude nonviable cells from further analyses. GFP was acquired as direct fluorescence in the FL1 channel; OFP was acquired as direct fluorescence in the FL2 channel. ΔLNGFR was analyzed using an anti-ΔLNGFR antibody (see Table S12). Flow cytometry used a BD FACSCanto II (BD Bioscience) apparatus.

**Peripheral Blood Cells.** Peripheral blood was collected from the tail vein. After red blood cell lysis and 7-AAD vital staining, cells were immunostained with the appropriate antibodies. The different cell types/subsets were identified as follows:

- **Resident monocytes:** 7-AAD-CD11b+CD115+GR1– cells;
- **Inflammatory monocytes:** 7-AAD-CD11b+CD115+GR1+ cells;
- **Granulocytes:** 7-AAD-CD11b+CD115–GR1+ cells;
- **Donor-derived hematopoietic cells:** 7-AAD-CD45.1+ or 7-AAD-CD45.1+OFP+ cells.

**Tumors.** Tumors were excised and made into single cell suspensions by collagenase IV (0.2 mg/ml, Worthington), dispase (2 mg/ml, GIBCO) and DNaseI (0.1 mg/ml, Roche) treatment in IMDM medium for 30 min at 37°C. The different cell types/subsets were identified using different combinations of antibodies, as detailed below:

- **Lewis lung carcinoma**
  - MRC1+ TAMs: 7AAD-/CD11b+/F4/80+/MRC1+/CD11c– cells;
  - CD11c+ TAMs: 7AAD-/CD11b+/F4/80+/MRC1–/CD11c+ cells;
  - Granulocytes/iMCs: 7AAD-/CD45+/CD11b+/F4/80–/GR1+ cells;
  - B cells: 7AAD-/CD45+/CD19+;
  - T cells: 7AAD-/CD45+/CD3+;
  - NK-cells: 7AAD+/CD45+/NK1.1+.

- **N202 mammary carcinoma**
  - MRC1+ TAMs: 7AAD+/GR1+/F4/80+/MRC1+/CD11c– cells;
CD11c+ TAMs: 7AAD−/GR1+/F4/80+/MRC1−/CD11c+ cells;  
Granulocytes/iMCs: 7AAD−/GR1+/F4/80− cells;  

Tissues and Organs. Tissues and organs were harvested and processed as described above (see “Tumors”). Adipose tissue was isolated from the retroperitoneal cavity.

Adipose tissue  
MRC1+ adipose tissue macrophages: 7AAD−/GR1−/F4/80+/MRC1+/CD11c− cells;  
CD11c+ adipose tissue macrophages: 7AAD−/GR1−/F4/80+/CD11c+/MRC1− cells;  
Double-negative macrophages: 7AAD−/GR1−/F4/80+/CD11c−/MRC1− cells;  
Granulocytes: 7AAD−/GR1+/F4/80− cells;  

Spleen  
MRC1+ red pulp macrophages: 7AAD−/GR1−/F4/80+/MRC1+/CD11c− cells;  
CD11c+ DCs/macrophages: 7AAD−/GR1−/F4/80+/CD11c+/MRC1− cells;  
Double-negative macrophages: 7AAD−/GR1−/F4/80+/CD11c−/MRC1− cells;  
Granulocytes: 7AAD−/GR1+/F4/80− cells;  

Lung  
MRC1+CD11c− lung macrophages: 7AAD−/GR1−/F4/80+/MRC1+/CD11c− cells;  
MRC1+CD11c+ alveolar macrophages: 7AAD−/GR1−/F4/80+/CD11c+/MRC1+ cells;  
Double-negative macrophages: 7AAD−/GR1−/F4/80+/CD11c−/MRC1− cells;  
Granulocytes: 7AAD−/GR1+/F4/80− cells;  

Liver  
MRC1+CD11c− Kupffer cells: 7AAD−/GR1−/F4/80+/MRC1+/CD11c− cells;  
CD11c+F4/80− DCs: 7AAD−/GR1−/F4/80−/CD11c+ cells;  
Granulocytes: 7AAD−/GR1+/F4/80− cells;  

Note that cell sorting used different gating strategies (see below).

Sorting of TAMs and Adipose-Tissue Macrophages for Gene Expression—RNA-Seq or qPCR—Studies  
Tumors and retroperitoneal adipose tissue were excised, made into single-cell suspensions and stained with the antibodies listed in Table S12. To sort cells, we used a MoFlo apparatus (Dako). After sorting, purity of the cells was always > 90%. Five-50 × 10⁴ cells were obtained from each sorting session.

MRC1+ and CD11c+ TAMs of MMTV-PyMT mammary tumors were isolated as 7-AAD−/CD45+/F4/80+/MRC1+/CD11c− cells (n = 3) and 7-AAD−/CD45+/F4/80+/MRC1−/CD11c+ cells (n = 3), respectively. For each mouse, we obtained 4-5 small biopsies from as many tumors and pooled them together before tissue processing. Sorted cells were used for qPCR analysis of selected miRNAs and mRNAs.

TAMs of LLCs were isolated as follows:

1) 7-AAD−/CD11b+/Gr1−/F4/80+/OFP+ cells (n = 3/group) for RNA-Seq analysis of mRNAs; both miR-511-3p and miR-511-3p-mut everexpressing TAMs were isolated;  
2) 7-AAD−/CD11b+/Gr1−/CD31−/MRC1+/CD11c− cells (n = 3) or 7-AAD−/CD11b+/Gr1−/CD31−/MRC1+/CD11c+ cells (n = 3) for RNA-Seq analysis of mRNAs.  
3) 7-AAD−/F4/80+/OFP+ cells (n = 4/group) for qPCR analysis of miRNAs; both miR-511-3p and miR-511-3p-mut everexpressing TAMs were isolated.

MRC1+ and CD11c+ adipose tissue macrophages were isolated as 7AAD−/CD11b+/Gr1−/F4/80+/MRC1+/CD11c− cells (n = 3) and 7AAD−/CD11b+/Gr1−/F4/80+/MRC1−/CD11c+ cells (n = 3), respectively. Sorted cells were used for qPCR analysis of selected miRNAs and mRNAs.

Sorted cells were washed and lysed in QiaZol or RLT buffer (QIAGEN) for either RNA-Seq analysis (see below) or qPCR. For qPCR of miRNAs, we retrotranscribed small RNAs using the High Capacity cDNA Reverse Transcription Kit (Applied Biosystems) and small RNA reverse transcription primers specific for the mature form of Let7a, miR-16, miR-142-3p, miR-155, miR-511-5p, miR-511-3p, miR-146a, miR-15a, or miR-511-3p-mut. We then performed qPCR on retrotranscribed small RNAs, using either custom (for miR-511-3p and miR-511-3p-mut) or inventoried (other miRNAs) TaqMan small RNA assays (Applied Biosystems). For qPCR of mRNAs, we retrotranscribed RNA with SuperScript III (Vilo kit, Invitrogen). All qPCR analyses used TaqMan probes from Applied Biosystems.

qPCR (miRNA and mRNA) was run for 40 cycles in standard mode using an ABI7900HT or ViiA7 apparatus (Applied Biosystems). The SDS 2.2.1 software was used to extract raw data (Ct) and to perform gene expression analysis. To determine gene expression, the difference (ΔCt) between the threshold cycle (Ct) of each mRNA/miRNA and that of the reference gene was calculated by applying an equal threshold (0.02).
Immunofluorescence Staining and Confocal Microscopy

N202 tumors were cut into 6–10 μm cryostatic sections for immunofluorescence staining and confocal microscopy. Briefly, tumors were fixed for 2 hr in 4% paraformaldehyde, equilibrated for 12 hr in PBS containing 15% sucrose, 24 hr in PBS/20% sucrose, and eventually 48 hr in PBS/30% sucrose. The samples were then embedded in O.C.T. compound for quick freezing in liquid nitrogen. Cryostatic sections were laid on slides and immediately stained. Sections were then blocked with 5% fetal bovine serum in PBS containing 1% bovine serum albumin (BSA) and 0.1% Triton X-100 (PBS-T). For immunofluorescence staining of MRC1, we used goat anti-mouse MRC1 antibodies (R&D) followed by donkey anti-goat alexa546 secondary antibodies (Molecular Probes). For immunofluorescence staining of CD11c, we used APC-conjugated, hamster anti-mouse CD11c antibodies (BD Pharmigen). Concentrations of antibodies and detailed information on immunofluorescence staining protocols will be available upon request. Cell nuclei were labeled by TO-PRO-3 (Molecular Probes). GFP was acquired as direct fluorescence using the 488 nm channel.

Confocal microscopy used an Axioskop 2 plus direct microscope (Zeiss) equipped with a Radiance 2100 (BioRad) three-laser confocal device, or an Axiovert 200 microscope (Zeiss) equipped with a 5-laser Perkin Elmer UltraView Spinning disk confocal device. Fluorescent signals from the individual fluorophores were sequentially acquired from single optical sections and analyzed and pseudocolored by Paint Shop Pro X (Corel).

Analysis of Tumor-Associated Vasculature

Mice were thoracotomized under deep anesthesia. The entire vasculature of the animal was thoroughly rinsed by perfusing with saline (20–30 ml) via left ventricle injection. When perfusion was complete, 20 ml of yellow silicone rubber Microfill MV 122 (Flow Tech Inc., Carver, Massachusetts) were infused through the left ventricle at 1 mL/min flow rate. When filling was complete, all organs displayed a rich, yellow coloration. The heart was then clamped and the animal placed under refrigeration at 4°C overnight, to allow polymerization. On the following day, tumors were taken by careful dissection, and placed in a 50% mixture of water and glycerin. At successive 24 hr intervals, the glycerin concentration was raised to 75%, then 85%, and finally pure glycerin.

The analysis of the tumor-associated vasculature was performed on 200 μm-thick slices obtained from the tumor periphery (4 slices) and inner tumor mass (4 slices) of each tumor. The whole tumor slice was photographed at low magnification (4x). Pictures for morphometric analysis were taken using a Zeiss Axio Imager connected to an Axiocam MRc5 camera (Zeiss) and analysis was performed using NeuronJ application of ImageJ software.

Target Gene Prediction

We used two distinct bioinformatics tools, TargetScan (Lewis et al., 2005) and DIANA microT (Maragkakis et al., 2009) to search for miR-511-3p target genes. The analysis retrieved a list of putative target genes that we analyzed by using DAVID Bioinformatic resources 6.7 (Huang et al., 2009).

We also investigated whether genes containing in their 3'-UTR at least one sequence with perfect complementarity to the seed sequence of the miRNA (i.e., to positions 2-8; Bartel, 2009) were modulated by miR-511-3p overexpression in vivo. miRNA seed/3'-UTR interactions include M8-A1 8mers (the mRNA sequence binds to the miRNA from position 2 to 8 and contains an Adenosine in position 1) and M8 7mers (the mRNA sequence matches the miRNA from position 2 to 8). We then used TargetRank (Nielsen et al., 2007) to retrieve the cellular transcripts that contain in their 3'-UTR either M8-A1 8-mer or M8 7-mer binding sites for miR-511-3p, 3p-mut or ~5p (see Table S6).

Luciferase Assays for UTR/miRNA Interactions

In order to validate miR-511-3p predicted targets, we cloned the 3'-UTR of Sema3A, Pdpn, CD163 and Rock2 downstream to the firefly luciferase of the pmir-GLO construct (Promega). The Rock2 UTR was split into 2 fragments (Rock2(1): 1537 bp; Rock2(2): 1771 bp). The Rock2(1) fragment contains four closely spaced, predicted miRNA sites; the Rock2(2) fragment contains only one predicted miRNA site (by RNAhybrid bioinformatic tool). To amplify the selected 3'-UTRs we used the following primers:

*Sema3A:*

Fw primer: TGC GCCACCTCCCCAAAACCTC;
Rv primer: TCC GTACTCTGGTCTCAGGCT;

*Pdpn:*

Fw primer: ACAGGTGTTCCTCCCAACACATCTG;
Rv primer: TGG CCTCATTTCTGGACACAATCAGG;

*CD163:*

Fw primer: GCC TTGACAGGACAGCGACAGCT;
Rv primer: TCCCCAACGTGTTTCACCTCC;

*Rock2(1):*

Fw primer: CGGCGCATGCTTGCCCTACCTG;
Rv primer: CCG ACCAGAGCAGCAGCTG;

*Rock2(2):*

Fw primer: ACC TTGAGAAGGCCCAGTTG;
Rv primer: ACCAAAGTGAATCGGAGGCGG;
In order to clone the PCR products downstream to the firefly luciferase expression cassette within the pmir-GLO construct, we incorporated a restriction site for NheI at the 5’-end of all Fw-primer sequences, and a restriction site for Sall at the 3’-end of all Rv primers.

Untransduced RAW264.7 cells, or RAW264.7 cells expressing exogenous miR-511 sequences (by SFFV.miR-511 or –511-mut LVs), were transfected using Lipofectamine 2000 (Invitrogen) with 50 ng of the pmir-GLO-based plasmids. Cells were lysed after 24 hr using the Dual-Luciferase Reporter Assay protocol (Promega). Renilla luciferase was used to normalize firefly luciferase activity.

**GFP Repression Assays for ROCK2-UTR/miR-511-3p Interactions**

In order to validate the interaction between miR-511-3p and its predicted binding sites in the ROCK2 3’-UTR, we employed the GFP/ΔLNGFR bidirectional LV. We generated by DNA synthesis (GeneArt Invitrogen) a 500 nt-long DNA fragment of either mouse or human ROCK2 3’-UTR encompassing a “core” UTR region containing 3 or 4 putative target sites for miR-511-3p (Table S13). We also generated mutant ROCK2 3’-UTR fragments, as illustrated in Figure S3B (mouse UTR) and S4C (human UTR); point mutations were selected that should abrogate miRNA seed/target mRNA interactions (see Figures S3B and S4C). In order to clone the synthetic DNA fragment downstream to the GFP expression cassette, we incorporated a restriction site for XbaI at the 5’-end, and a restriction site for Sall at the 3’-end of the fragment.

We then transduced mouse P388D1 (for testing the mouse ROCK2-UTR) or human U937 cells (for testing the human ROCK2-UTR) with the reporter LVs carrying wild-type, mutant, or no ROCK2 3’-UTR, and superinfected the transduced cells 5 days later with miR-511-overexpressing vectors (SFFV.miR-511 and –511-mut LVs). We then analyzed GFP and ΔLNGFR expression to measure GFP repression.

**Western Blot**

Mouse (P388D1, RAW264.7) and human (U937) monocytic cell lines were transduced with the SFFV-miR-511 or –511-mut LVs, expanded in culture for at least 2 weeks, collected and directly stored at –80°C. BM cells were transduced with the SFFV-miR-511 or –511-mut LVs on day 1 post-plating, and BMDMs obtained by culturing the cells in the presence of M-CSF for 1 week. BMDMs were then washed with PBS, collected and directly lysed. Each cell type was homogenized in 10x volume of RIPA lysis buffer (10mM Tris-Cl, pH 7.2, 150mM NaCl, 1mM EDTA pH 8) with 1% Triton X-100, 1% deoxycholate, 0.1% SDS, protease and phosphatase inhibitor mixture (Roche). Samples were then diluted in Laemmli’s SDS-sample buffer. Proteins (~60 μg) were separated by electrophoresis on 8% polyacrilamide (Sigma) gels and transferred onto trans-blot nitrocellulose membranes (Amersham). Ponceau staining (Sigma) was performed to confirm that the samples were loaded equally. The membranes were blocked in 5% nonfat dry milk in TBS-T and then incubated for 1 hr at room temperature. Primary antibodies were diluted in 3% bovine serum albumin (BSA, Sigma) or 5% nonfat dry milk in TBS-T, and the membranes were incubated overnight at 4°C. The primary antibody was removed, and the blots washed in TBS-T and then incubated for 1 hr at room temperature in horseradish peroxidase-conjugated secondary antibodies (Amersham).

The primary antibodies used were: mouse anti-ROCK2 (CLONE: 21; BD Biosciences); rabbit polyclonal anti-LTBP1 (Abcam); mouse anti-GAPDH (Sigma); rabbit anti-calnexin (GenTex). Reactive proteins were visualized using LiteBlot (Euroclone, Celbio) or secondary antibodies (Amersham).

**RNA-Seq**

RNA was extracted from sorted TAMs (see above) using either Qiazol reagent (QIAGEN) for F4/80^+^OPF^+^ TAMs (miR overexpressing TAMs) or RNeasy columns (QIAGEN) for MRC1^+^ and CD11c^+^ TAMs (wild-type TAMs), following the manufacturer’s instructions.

RNA isolated from MRC1^+^ and CD11c^+^ TAMs was depleted of rRNA using the Ribo-Zero RNA removal kit (Epicerin Biotechnologies), followed by concentration using RNA Clean & Concentrator –5 columns (Zymo Research), with in-tube DNase (Epicerin Biotechnologies) digestion. Illumina sequencing libraries were prepared according to the TruSeq RNA Sample Preparation Guide (Revision A), starting at the RNA fragmentation step. RNA fragmentation with “Elute, Prime, Fragment Mix” was performed for 4 min at 94°C. Sequencing was performed on a HiSeq 2000 (Illumina) using paired-end cBot v2 and TruSeq SBS reagents. Libraries were sequenced using 2 × 50 bp paired-end reads, with two indexed samples per lane, yielding 89-115 million reads (4.4-5.8 Gb) per sample. The sequencing was carried out using HiSeq Control Software (HCS) version 1.1.37.19. Image analysis and base calling was performed using Illumina’s real time analysis (RTA) software version 1.7.48. Reads were filtered to remove those with low base call quality using Illumina’s default chastity criteria. The results were then demultiplexed and converted to fastq format files by CASAVA version 1.7.

RNA isolated from sorted F4/80^+^OPF^+^ TAMs did not undergo rRNA depletion, but was processed using the poly-T oligonucleotide coated magnetic beads provided with the Illumina TruSeq RNA Sample Preparation kit as directed by the manufacturer. RNA fragmentation with “Elute, Prime, Fragment Mix” was again performed for 4 min at 94°C. Sequencing was performed as above, except that 100 bp paired-end reads were generated, yielding 69-84 million reads (6.9–8.4 Gb) per sample.
RNA-Seq Analysis

Paired-end sequence reads were aligned to the mouse genome (mm9; http://www.ensembl.org) using Bowtie (doi:10.1186/gb-2009-10-3-r25) and TopHat (doi:10.1093/bioinformatics/btp120). Reads were mapped to known genes and splice junctions by providing TopHat with an annotation file (Mus_musculus.NCBIM37.62.gtf; http://www.ensembl.org). Samtools (doi: 10.1093/bioinformatics/btp352) was then used to remove PCR-generated duplicate reads. Count data for each exon was generated using htseq-count from the HTseq package (http://www-huber.embl.de/users/anders/HTSeq/).

Differential expression between sample groups was identified from the sequence count data using the R package DEseq (doi:10.1186/gb-2010-11-10-r106). Expression differences were considered significant at a false discovery rate (FDR) of 5%.

Cumulative distribution analyses of fold-changes in the whole transcriptome and transcripts that contain M8-A1 8-mer and M8 7-mer target sites were generated in R (http://www.r-project.org), as described (Grimson et al., 2007). Statistical analysis was performed by one-sided Kolmogorov-Smirnov test.

SUPPLEMENTAL REFERENCES

Figure S1. Immunophenotyping of P388D1 and RAW264.7 Monocytic Cells, Related to Figure 1

P388D1 cells were stained with the indicated antibodies to analyze the expression of myeloid (CD11b, CD11c), monocyte/macrophage (F4/80), B cell (CD19) and T cell (CD3) markers (top row). Unstained cells (fluorescence-minus one, FMO) are shown in the bottom row. RAW264.7 cells were stained with the indicated antibodies to analyze the expression of myeloid (CD11b, CD11c), monocyte/macrophage (F4/80) and T cell (CD3) markers (top row). Unstained cells (FMO) are shown in the bottom row. Note that both P388D1 and RAW264.7 cells express the monocyte/macrophage-specific marker, F4/80.
Figure S2. Identification of Tumor-Infiltrating Hematopoietic Cells in Lewis Lung Carcinoma (LLC) and N202 Mammary Carcinoma, Related to Figure 2
(A) LLCs grown for 4 weeks in C57BL/6 mice. Left panel: Flow cytometry analysis of the following circulating leukocyte subsets: CD115*Gr1− resident monocytes (1); CD115*Gr1+ inflammatory monocytes (also termed myeloid-derived suppressor cells in tumor-bearing mice) (2); and CD115−Gr1+ granulocytes (3). Middle and right panels: Flow cytometry analysis of tumor-derived leukocytes; tumors were harvested and made into single-cell suspensions. Middle panels: 7-AAD−CD11b+F4/80+MRC1−CD11clow TAMs (1); and 7-AAD−CD11b+F4/80+CD11c+MRC1low TAMs (2). Right panels: 7-AAD−CD45+CD11b+F4/80−Gr1+ granulocytes/iMCs (1).

(B) N202 mammary carcinomas grown for 4 weeks in FVB/n mice. Tumors were harvested and made into single cell suspensions, and the following cell types analyzed: 7-AAD−Gr1−F4/80− granulocytes/iMCs (1); 7-AAD−F4/80−Gr1−MRC1−CD11clow TAMs (2); and 7-AAD−F4/80−Gr1−CD11c+MRC1low TAMs (3).
Figure S3. Stem-Loop Structures of the Mouse Pre-miR-511 and Pre-miR-511-mut, and Annotated Sequences of the Wild-Type and Mutated Mouse Rock2 3' UTR, Related to Figure 4

(A) Stem-loop structures of the mouse pre-miR-511 and pre-miR-511-mut. miR-511-5p and 3p sequences are shown in blue and red, respectively; Mutated nucleotides are shown in green. Note that both pre-miR-511 and pre-miR-511-mut generate a wild-type miR-511-5p sequence upon processing of the pre-miRNA.

(B) Core sequence of the mouse Rock2 3'-UTR. Wild-type sequence: The genomic sequence of the Rock2 3'-UTR encompassing the miR-511-3p target sites is illustrated. Four target sites (8-mer-A1; 7-mer-M8; 7-mer-M8; 6-mer) are indicated by red letters, and the binding of the miRNA illustrated for each site. Interactions were retrieved by RNA-Hybrid. Mutated sequence: Each of the four miR-511-3p target sites was mutated by substituting 3 nucleotides in each site. Mutated nucleotides are underlined and shown in gray boxes.
Figure S4. Stem-Loop Structures of the Human Pre-miR-511 and Pre-miR-511-mut, and Annotated Sequences of the Wild-Type and Mutated Human ROCK2 3' UTR, Related to Figure 5

(A) Conservation of miR-511-5p and miR-511-3p among species. The yellow shading indicates 100% identity.

(B) Stem-loop structures of the human pre-miR-511 and pre-miR-511-mut. miR-511-5p and miR-511-3p sequences are shown in blue and red, respectively; Mutated nucleotides are shown in green. Note that both pre-miR-511 and pre-miR-511-mut generate a wild-type miR-511-5p sequence upon processing of the pre-miRNA.

(C) Core sequence of the human ROCK2 3'-UTR. Wild-type sequence: The genomic sequence of the ROCK2 3'-UTR encompassing the miR-511-3p target sites is illustrated. Three target sites (8-mer-A1; 7-mer-M8; 7-mer-M8) are indicated by red letters, and the binding of the miRNA illustrated for each site. Interactions were retrieved by RNA-Hybrid. Mutated sequence: Each of the three miR-511-3p target sites was mutated by substituting 3 nucleotides in each site. Mutated nucleotides are underlined and shown in gray boxes.

Wild-type sequence:

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<th>miR-511-3p (7mer-M8)</th>
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Mutated sequence:

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<th>miR-511-3p (7mer-M8)</th>
<th>miR-511-3p (7mer-A1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>GAAACA</td>
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<td>5’...UCACUUUUUUAUAACACAU</td>
<td>5’...AUAAUAUAAACACACACACAU</td>
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</table>

Figure S4. Stem-Loop Structures of the Human Pre-miR-511 and Pre-miR-511-mut, and Annotated Sequences of the Wild-Type and Mutated Human ROCK2 3' UTR, Related to Figure 5

(A) Conservation of miR-511-5p and miR-511-3p among species. The yellow shading indicates 100% identity.
(B) Stem-loop structures of the human pre-miR-511 and pre-miR-511-mut. miR-511-5p and miR-511-3p sequences are shown in blue and red, respectively; Mutated nucleotides are shown in green. Note that both pre-miR-511 and pre-miR-511-mut generate a wild-type miR-511-5p sequence upon processing of the pre-miRNA.
(C) Core sequence of the human ROCK2 3'-UTR. Wild-type sequence: The genomic sequence of the ROCK2 3'-UTR encompassing the miR-511-3p target sites is illustrated. Three target sites (8-mer-A1; 7-mer-M8; 7-mer-M8) are indicated by red letters, and the binding of the miRNA illustrated for each site. Interactions were retrieved by RNA-Hybrid. Mutated sequence: Each of the three miR-511-3p target sites was mutated by substituting 3 nucleotides in each site. Mutated nucleotides are underlined and shown in gray boxes.
Figure S5. Hematopoietic Chimerism and Identification of Tumor-Infiltrating Hematopoietic Cells in LLCs of miR-511-Overexpressing Mice, Related to Figure 6

(A) Proportions of CD45.1\(^+\) and CD45.1\(^+\)OFP\(^+\) leukocytes in the blood of mice either overexpressing miR-511-3p (SFFV.miR-511) or miR-511-3p-mut (SFFV.miR-511-mut) in hematopoietic cells. Blood leukocytes were analyzed at the end of the experiment (8 weeks post-HS/PC transplant). Data show the percentage of marker-positive cells (mean values ± SEM; n = 4 mice/group).

(B) Proportions of B cells (CD19\(^+\)), NK cells (NK1.1\(^+\)), CD8\(^+\) T cells, total TAMs (F4/80\(^+\)), and granulocytes (GR1\(^+\)) among the total LLC-infiltrating CD45\(^+\) hematopoietic cells. Tumors were analyzed at the end of the experiment (8 weeks post-HS/PC transplant; 4 weeks post-tumor injection). Data show the percentage of marker-positive cells (mean values ± SEM; n = 4 mice/group).

(C) Proportion of MRC1\(^+\) and CD11c\(^+\) TAMs among the total F4/80\(^+\) TAMs in LLCs grown in mice either overexpressing miR-511 or miR-511-mut in hematopoietic cells. Tumors were analyzed at the end of the experiment (8 weeks post-HS/PC transplant; 4 weeks post-tumor injection). Data show the percentage of marker-positive cells (mean values ± SEM; n = 4 mice/group).