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(54) PREVENTING TUMOR DEVELOPMENT AND METASTASIS

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(57) ABSTRACT

Treatment of tumors, especially breast cancer or glioblastoma tumors, by silencing RAI27A and/or TRAF3IP2, compositions and methods for same.

Specification includes a Sequence Listing.
<table>
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<tr>
<th>Name</th>
<th>Sequence</th>
<th>Variant 1</th>
<th>Variant 2</th>
<th>Variant 3</th>
<th>Variant 4</th>
<th>Variant 5</th>
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SEQ ID NO. 1: CCGGCTTCGAGCTGTTTTAATTTTCTGTTTTT
SEQ ID NO. 2: CCGGCTTCGAGCTGTTTTAATTTTCTGTTTTT
SEQ ID NO. 3: CCGGCTTCGAGCTGTTTTAATTTTCTGTTTTT
SEQ ID NO. 4: CCGGCTTCGAGCTGTTTTAATTTTCTGTTTTT
SEQ ID NO. 5: CCGGCTTCGAGCTGTTTTAATTTTCTGTTTTT
SEQ ID NO. 6: CCGGCTTCGAGCTGTTTTAATTTTCTGTTTTT
FIGURE 9

- PBS
- LV TRAF3IP2 II

Tumor volume $\text{mm}^3$

Day 0  Day 6  Day 9  Day 12  Day 15  Day 19  Day 22  Day 26  Day 29  Day 33

time after injection

FIGURE 10

U87
KD TRAF3IP2

FIGURE 11

U87
KD TRAF3IP2
FIGURE 12

![Graph showing absorbance over time for U87 wild type and U87 TRAF3IP2 KD.](image)

**Absorbance 450nm**

- U87 wild type
- U87 TRAF3IP2 KD

**Number of days after seeding the cells**

- p<0.01
- p>0.05
FIGURE 13

A

<table>
<thead>
<tr>
<th>Fold Change</th>
<th>SVP g12</th>
<th>U87</th>
<th>U87 TRAF3IP2 KD</th>
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<tbody>
<tr>
<td>p&lt;0.001</td>
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</tr>
</tbody>
</table>

B

U87 scrambled siRNA U87 KD TRAF3IP2

TRAF3IP2

GAPDH
FIGURE 14

G0/G1 Phase S Phase G2/M Phase

FIGURE 15

4780
35.6
1338.3
0
1000
2000
3000
4000
5000
6000

tumor volume in mm³

Week 8
Week 14

U87
U87 TRAF3IP2 KD
FIGURE 17B

ASO1 (SEQ ID NO. 13): mG*mG*mU*mG*mG*C*A*C*A*T*G*C*T*C*mC*mU*mU*mC*mU

ASO2 (SEQ ID NO. 14): mA*mG*mU*mG*mC*T*A*C*C*G*A*C*C*A*G*mC*mC*mU

ASO3 (SEQ ID NO. 15): mG*mG*mC*mC*mU*C*T*C*T*C*G*T*G*G*mU*mC*mC*mC*mA

ASO4 (SEQ ID NO. 16): mA*mU*mG*mC*mC*T*C*G*G*A*T*T*C*T*A*mU*mC*mC*mU*mC

ASO5 (SEQ ID NO. 17): mG*mU*mU*mG*mC*A*C*C*A*T*C*C*C*T*mG*mG*mC*mU*mA

ASO6 (SEQ ID NO. 18): mU*mG*mG*mU*mG*A*T*G*T*G*G*C*T*G*G*mU*mC*mC*mU*mG
FIGURE 18

TRAF3IP2

1239/Control (30mM)
1239/Control (10mM)
1239/Control (3mM)
1239/Control (0mM)

1241/Control (30mM)
1241/Control (10mM)
1241/Control (3mM)
1241/Control (0mM)

1316/Control (30mM)
1316/Control (10mM)
1316/Control (3mM)
1316/Control (0mM)

ASO2/Control (30mM)
ASO2/Control (10mM)
ASO2/Control (3mM)

ASO1/Control (30mM)
ASO1/Control (10mM)
ASO1/Control (3mM)

187 ACT1 /87 SCR

G E N E  E X P R E S S I O N  F O L D  C H A N G E

ASO
Exon 4
ASO
Exon 7
ASO
Exon 3
ASO
Exon 5
ASO
Exon 10
Land Virtue
Exons 9 & X10

0  1  2  3  4  5
FIGURE 19

A

Tumor Size mm

Days Post Tumor Induction

- TRAF3IP2 shRNA-LV Treated
- SCR shRNA-LV Treated

B

Luciferase

Day 14
Day 50

Day Post Tumor Induction

C
PREVENTING TUMOR DEVELOPMENT AND METASTASIS

CROSS-REFERENCES TO RELATED APPLICATIONS

[0001] This application is a continuation-in-part application of Ser. No. 14/814,130, filed Jul. 30, 2015, which claims priority to U.S. Ser. No. 62/031,021, filed Jul. 30, 2014, each of which is incorporated by reference herein in its entirety for all purposes.

STATEMENT REGARDING FEDERALLY SPONSORED RESEARCH

[0002] Not applicable.

FIELD OF THE INVENTION

[0003] The present invention relates to novel methods to prevent tumor metastasis and suppress tumor growth, especially of solid tumors, by interfering with tumor communication and its environment and by impacting the formation and development of the tumor microenvironment.

DESCRIPTION OF RELATED ART

[0004] Tumor development occurs following the accumulation of genetic and epigenetic alterations in tumor cells. It has been demonstrated that tumor growth is strongly influenced by non-malignant cells that together with the tumor cells form the tumor microenvironment. Numerous reports have revealed the complexity of the communication between tumor cells and the heterogeneous population of stromal cells within the tumor microenvironment.

[0005] For example, the tumor-stromal cell interactions have a crucial role in tumor initiation and progression. These stromal cells, including fibroblasts, myofibroblasts, endothelial cells, mesothelial cells, adipocytes, tissue resident stem cells, and immune cells, are involved in tumor development via several mechanisms including:

(i) cell-cell and cell-matrix interactions influencing cancer cell sensitivity to apoptosis;
(ii) local release of soluble and genetically modifying factors promoting survival and tumor growth, growth of tumor blood vessels and resistance to attack by the patient’s immune system (crosstalk between stromal, immune cells and tumor cells);
(iii) direct cell-cell interactions with tumor cells (crosstalk or oncolytic trogocytosis);
(iv) generation of specific properties and niches within the tumor microenvironment that facilitate the acquisition of drug resistance; and
(v) conversion of cancer cells to cancer-initiating cells or cancer stem cells.

[0011] These interactions between malignant and non-malignant cells modify cellular compartments, leading to the co-evolution of tumor cells and their microenvironment.

[0012] Although the importance of microenvironmental alterations in tumor development is recognized, the molecular mechanisms underlying these changes are only now beginning to be understood. Detailed molecular characterization of various cell types from normal breast tissue, ductal carcinoma, and invasive breast tumors has revealed that gene expression changes occur in all cell types during breast tumor progression.

[0013] Recently, it has been shown that, in addition to bone marrow-derived MSCs, adipose tissue-derived MSCs display significant affinity to tumor microenvironment. The role of inflammation in the tumor microenvironment is crucial in the pathology of cancer because it regulates the directional movement of tissue resident immune cells and stem cells.

[0014] Although decades of research have yielded targeted therapies that are effective in eliminating or reducing some tumors, breast cancer remains the leading cause of morbidity and second-leading cause of death in women. Recent published reports suggest that reciprocal influences exist between breast tumor cells and the tumor microenvironment and that these interactions affect the growth and energetics of the tumor. These interactions reveal the contributions of individual cells within a tumor to the overall disease. In addition, a neurological tumor such as glioblastoma multiforme is even more malignant and the 5-year survival rate of patients diagnosed with such a tumor still is below 5%.

[0015] The present invention provides novel compositions and methods to affect the interactions between a tumor and its microenvironment to prevent, reverse, and/or reduce tumor growth and metastasis.

BRIEF SUMMARY OF THE DISCLOSURE

[0016] The present disclosure provides novel therapeutics for tumors, especially solid tumors, including breast cancers or glioblastomas, by interfering with tumor communication with the tumor environment and/or by regressing formation of the microenvironment, thereby preventing or reversing tumor metastasis and suppressing tumor growth. In a preferred embodiment, the present invention discloses a cancer therapy by silencing TRAF3IP2 and/or RAB27A expression.

[0017] Silencers can be delivered to a tumor in a number of ways, including at least:

1) Delivering silencing RNA by injecting an expression vector encoding the silencer to the tumor site, e.g., directly into a tumor site under visual, ultrasound, florescence, CT or MRI guidance or other imaging modalities, or indirectly through blood vessels or ducts that lead to the tumor.

2) Use of silencing RNA delivered by tumor targeting cells, such as migratory stem cells, e.g., MSCs, or any type of cells that due to their nature preferably migrate and engraft to the tumor site. Such cells would contain therein either an expression vector or a genomic copy of the sequence encoding the silencer.

3) Delivering encapsulated or otherwise protected silencing RNA to the tumor site. The silencing RNA is for example encapsulated into microspheres (i.e. exosomes) or micelles, liposomes and the like. The microspheres will be delivered by direct or indirect injection to the tumor site either through a transcutaneous injection or through a vessel or duct supplying the tumor site. Preferably, such RNAs will be RNase resistant, and if so, naked RNA may be used.

4) Silencing RNA linked to a specific tumor directed antibody or protamine coupled construct to increase the tumor specific concentration and to enhance the local effect of the silencing RNA within the tumor site.

5) Achieving a selective effect targeting the tumor cells and virtually avoiding an effect on non tumor cells by i) increasing the local concentration within the tumor by selective delivery means as described above, ii) by the fact
that the respective genes of TRAF3IP2 and of Rab27a are ten to thirty times (respectively) upregulated in tumor, especially in tumor stem cells, compared to normal stem cells, and (iii) the silencer is released in a (transactivator)-inducible manner (such as II.1B), thus expression is activated mainly in the tumor.

[0023] 6) Combinations and variations of the above.

[0024] Silencing TRAF3IP2 in tumor cells confines cytokine expression and ultimately limits the development of the tumor microenvironment. This eventually slows or prevents tumor growth and restrains tumor metastasis. The tumor cells exhibit significantly higher levels of exocytosis activities compared to non-malignant cells.

[0025] Two alternative transcripts of TRAF3IP2 encoding different proteins have been identified. A third transcript, which does not encode a protein and is transcribed in the opposite orientation, has also been identified. Overexpression of this transcript has been shown to reduce expression of at least one of the protein encoding transcripts, suggesting it has a regulatory role in the expression of this gene and indicating its use in the methods described herein.

[0026] For the actual silencer sequence used in our proof of concept studies, we used commercially available silencers (SIGMA ALDRICH®) RNA to target TRAF3IP2 and Rab27A either separately or in combination. However, any type of silencer for these genes could be used.

[0027] Basic design rules for the various types of silencers are available, and once designed the silencers can be tested for efficacy according to the methods discussed herein and in the literature.

[0028] For example, a short hairpin silencer (shRNA) generally has about 18-30 nucleotides (nt), preferably 21 nt, comprising a unique sense strand of target mRNA beginning with AA linked to a loop (3-9 nt) linked to an complement of the unique sense strand and finishing with polyT, thus forming a hairpin. An initiating G nt could also be used.

[0029] Another type of silencer is the siRNA of about 18-30 nt, preferably 21 nt, comprising a unique sense strand of the target mRNA beginning with AA and finishing with polyT.

[0030] Another type of silencer is the antisense sequence. These can be a unique antisense sequence from the target, or an RNAse resistant 18-30 nt antisense RNA sequence from the target. Effective antisense silencers may also be located in exons, but close to the acceptor splice site (SS).

[0031] miRNAs generally work when about 21-23 nt and have complementarity maintained in the first third of the small RNA and target mRNA, but mismatches arise in the remainder of the aligned sequence.

[0032] The above rules are guidelines only, however, and there is certainly variability in approaches. Therefore, it is typical to design 4-6 such silencers using the basic rules and then test each for activity, e.g., in an ex vivo system. Therefore, given the validity of the target, silencers can be readily designed using the target sequence.

[0033] In addition, validated silencers for several genes are already commercially available. LIFE TECHNOLOGIES® for example has 27 validated silencers (6 human) for TRAF3IP2, and 9 for Rab27A (3 human). SIGMA-ALDRICH® also provides several shRNAs and siRNAs for use, including the human TRAF3IP2 silencer MISSION® shRNA Lentiviral Transduction Particles (SHCLNV-NM 147200) and the human Rab27a silencer MISSION® shRNA Lentiviral Transduction Particles (SHCLNV-NM 004580). In addition, Sigma offers miRNA mimics, and esiRNA. Furthermore, the RNAi Consortium has built a library of shRNAs directed against 15,000 human and 15,000 mouse genes.

[0034] Furthermore, silencer RNAs can be stabilized against nucleases by incorporating modified bases therein, such as methylphosphonate, phosphorothioate, 2′-O-substituted RNA, phosphoramidite, morpholino and chimeras contain an internal core of unmodified phosphodiester RNA/RNA flanked by modified residues. These can be very useful where naked or encapsulated nucleic acid is directly delivered, as opposed to an expression vector encoding the silencer.

[0035] We have specifically targeted breast cancer and glioblastoma cell lines herein for proof of concept experiments, but we anticipate that the method can be used in many cancers or inflammatory conditions since TRAF3IP2 and/or Rab27A play a role therein. The TRAF3IP2 gene, for example, is implicated in several cancers, including but not limited to lung cancer, colon cancer, cervical cancer, endometrial cancer, liver cancer, ovarian cancer, prostate cancer, gastrointestinal cancer, testis cancer, thyroid cancer, carcinoid tumor, uterine carcer, pancreatic cancer, sarcoma, melanoma and the like. See e.g., proteinatlas.org/ENSG000000056972-TRAF3IP2/cancer. It is also implicated in inflammatory bowel disease, atopic dermatitis, psoriasis, Hodgkin disease, familial candidiasis, possibly ulcerative colitis, and the like. Any cancer or diseased tissue with at least 5 or 10 or higher levels of either of these transcripts can be addressed by in the methods herein.

[0036] Rab27A not only regulates exocytosis, and thus silencing Rab27A attenuates exocytosis. The lower exocytosis limits the release of oncocgenic molecules into the tumor microenvironment in both soluble and insoluble forms. This ultimately restricts the development of tumor microenvironment.

[0037] Rab27a is known to be highly expressed in some cancer as well, including pancreatic cancer, breast cancer, colorectal, lymphoma, prostate, melanoma, ovarian, thyroid, and the like.

[0038] While there are several methods of delivering silencers to tumors, one preferred method uses of mesenchymal stem cells or “MSCs”. Using their known preferred tumor homing capacity, MSCs are modified with a vector expressing the respective silencing sequence. Silencing vectors thus delivered to the tumor site foci using these MSCs that produce the respective silencing RNA against TRAF3IP2, Rab27a, or against both. In addition and as a means to increase the effect on tumor cells and minimize the effect on non-tumor cells, tumor-tropic subset of MSCs that are obtained and identified by their preference to migrate towards the tumor cells can be used. They can be created by prior exposure to exosomes that induce the needed epigenetic changes in the MSC or by selecting by FACS sorting MSCs expressing specific tumor surface markers, such CXCR4, or the PDGF b receptor.

[0039] The tumor-tropic MSCs carrying therapeutic vectors will home to the vicinity of tumor cells and then express the silencers in the tumor microenvironment where there is higher expression of II.1B, if we use an II.1B inducible promoter herein. The silencer is thus released and reduces tumor-related inflammation and tumor size with minimal off-target effects since healthy tissue won’t have high levels of II.1B. The MSCs containing silencing vector (5x10^7/
subject) are administered systemically, e.g., by injection into the bloodstream, into a local tumor supporting blood vessel or duct, or directly transcutaneous into the primary tumor or its metastasis.

[0040] Although we have used MSCs as delivery vehicles herein, this is an continually evolving area of research and another method may ultimately emerge as more preferred over the course of research. Other possible delivery vehicles include Rexin-G, an engineered retroviral nanoparticle that achieves targeting to cancerous lesions through the attachment of a collagen motif that binds to “newly exposed” extracellular matrix, which is typically associated with tumor tissue. Another possibility is to use a virus engineered to target a particular cancer cell, such as the parvo virus 11, or to link the silencer with tumor specific ligand or antibodies.

[0041] There are also non-viral methods of silencer delivery, including e.g., injecting naked DNA/RNA into a tumor, injected protected RNA into tumors, electroporation, the use of polymers, liposomes, and the like, to protect the nucleic acids, or to stabilize the silencer through linking it to Protamin.

[0042] Lentiviral vectors were used herein to encode the silencer sequences for TRAF3IP2 and RAB27A. Although data show that there is specificity for CD45+ cells transduction in vivo when administering lentiviral vectors, MDAMB231 and SW620 cells are highly transducible with lentiviral vectors. Thus, these vectors were useful for proof of concept studies. However, any suitable expression vector may be used herein, or the gene can be introduced into the genome of a homing cell (e.g., by homologous recombination), such as the MSCs discussed herein.

[0043] Common vectors are based on herpes simplex type 1 recombinant vector (HSV-1); adenovirus, adenovirus-associated viral vector (AAV); alpha virus; vaccinia virus; pox virus; Sendai virus; plasmid, retrovirus; ssDNA vectors; and the like. To date, adenovirus, retrovirus and naked plasmid DNA have made up more than half of the vectors tested in clinical trials of various gene therapies.

[0044] An IL1B transactivator-inducible system is a preferred promoter for use in our lentiviral vector. The IL1B promoter activates the expression of silencer RNA by binding the endogenous IL1B, which is highly produced by cells within tumor microenvironment. However, this promoter is exemplary only and there are many to choose from, including several antibiotic resistance or drug responsive promoters that can be safely used in humans (e.g., tamoxifen, tetracyclin, ampicillin and the like).

[0045] The disclosure provides one or more of the following embodiments, in any combinations thereof:

A pharmaceutical composition for the treatment of a tumor having increased expression of TRAF3IP2, wherein said composition comprises at least one silencing sequence for TRAF3IP2 in a pharmaceutically acceptable carrier in an amount effective for the therapeutic treatment of a tumor, wherein said silencing sequence reduces the expression of the TRAF3IP2 gene by at least two-fold as comparing to without the silencing sequence for TRAF3IP2, and wherein said silencing sequence is a modified portion of sense strand of NM_001164281.2 (SEQ ID NO. 7), NM_1472002.2 (SEQ ID NO. 8), XM_011535386.2 (SEQ ID NO. 9), NM_347686.3 (SEQ ID NO. 10), XM_000712336.4 (SEQ ID NO. 11), and NM_001164282.2 (SEQ ID NO. 12).

Any composition herein described, the composition comprises an expression vector encoding a TRAF3IP2 silencer operably coupled to an inducible promoter.

Any composition herein described, the silencing sequence is an siRNA, an miRNA, an shRNA, an antisense RNA, or an antisense oligonucleotide.

Any composition herein described, the silencing sequence is an expression vector encoded by an expression vector hosted in a mesenchymal stem cell (MSC) that targets said tumor.

Any composition herein described, the MSC having been previously exposed to exosomes from said tumor.

Any composition herein described, the silencing sequence is an antisense oligonucleotide that is 13-25 nucleotides in length.

Any composition herein described, wherein said silencer is an siRNA, an shRNA, an miRNA, or an antisense oligonucleotide.

Any composition herein described, wherein said silencer comprises any sequence herein referenced or described.

Any composition herein described, the antisense oligonucleotide is complementary to a portion of the sense strand of any one of SEQ ID NOs. 7-12.

Any composition herein described, the antisense oligonucleotide is selected from SEQ ID NOs. 13-18.

Any composition herein described, the pharmaceutically acceptable carrier is a nucleic acid carrier.

Any composition herein described, further comprising a silencing sequence for RAB27A.

Any composition herein described, the composition is formulated for parenteral administration, including direct injection into a tumor or its metastasis site by transcutaneous, intraarterial, intraductal, intravenous, intradermal, intramuscular, intraperitoneal, or subcutaneous administration.

Any composition herein described, the composition is used in treating glioblastomas or breast cancer, or for use in treating any cancer with at least 2-fold increased TRAF3IP2 and/or RAB27A expression.

A method of treating at least one tumor in a mammal comprising administering to the mammal an effective amount of any composition herein.

A method as herein described, wherein said tumor is a human breast cancer or a glioblastoma or any cancer with at least 2-fold increased TRAF3IP2 and/or RAB27A levels, or at least 10 fold, or at least 20 fold or at least 50 fold.

A method as herein described, wherein the composition is injected directly into said tumor and said injection is guided by ultrasound, fluoroscopy, imaging, CT, MRI, or just visually in order to enhance the local concentration of the silencer within the tumor.

A method as herein described, wherein the silencers are delivered to the tumor by an expression vector.
A method as herein described, wherein said siencers are encoded by expression vectors contained inside MSCs.
A method as herein described, wherein the siencers are delivered to the tumor by injection.
A method as herein described, wherein the siencers linked to an antibody targeting a breast tumor specific cell surface antigen.
A method to selectively treat a tumor and minimize side effects, by administering an effective amount of a siencer for TRAF3IP2 or Rab27a, or both, to a tumor that expresses at least 2 times the amount of TRAF3IP2 or Rab27a, or both, as compared to a non-tumor cell from the same tissue.
A method as herein described, further comprising enhancing the selective effect on tumor cells and avoiding effects on normal cells by increasing the local concentration of the siencer within the tumor by injecting said siencer(s) directly into said tumor.
A method as herein described, wherein said siencer(s) is encoded in an expression vector having an inducible promoter, thus enhancing the selective effect on tumor cells and avoiding effects on normal cells by means of selectively activating the production of the siencer by a switch that activates said inducible promoter.
A method as herein described, wherein said switch is preferentially or only found in said tumor.

[0046] As used herein, the term “expression vector” means a DNA or RNA into which a sequence of interest can be inserted that operably linked to a promoter such that the sequence will be transcribed or expressed from the promoter in the host cell/animal of interest. Thousands of such vectors are available. See e.g., Addgene.org which provides both a repository and a searchable database allowing vectors to be easily located and obtained from colleagues. See also Plasmid Information Database (PlasmID) and DNAU having over 191,000 plasmids. A collection of cloning vectors is also kept at the National Institute of Genetics as a resource for the biological research community. Furthermore, vectors (including particular ORFs therein) are usually available from colleagues.

[0047] As used herein, the term “silencing” refers to the down-regulation of gene expression. At least 65%, 70%, 75%, 80% reduction should be achieved, but preferably, this term refers to the ability of a cell to prevent the expression of a certain gene. Gene silencing can occur during either transcription or translation and is often used in research and gene therapies.

[0048] By “preventing” gene expression, we mean no detectable intact gene expression is detected when assayed by Northern blot using a radioactively end-labeled oligomer that is complementary to the gene being silenced. Nonetheless, there may be minute amounts of expression that could be detected by extremely sensitive methods.

[0049] The term “silencer” as used herein refers to a exogenous sequence that can be introduced into cells and used to silence gene expression in that cell. There are several different types of silencers, including at least antisense oligonucleotides, ribozymes, RNA interference, and the like. Genes can be silenced by e.g., dsRNA that decomposes mRNA, siRNA molecules that cause the endonucleic cleavage of the target mRNA molecules or by miRNA molecules that suppress translation of the mRNA molecule or by shRNA, as well as by endoribonuclease-prepared siRNAs (esiRNAs), which are a mixture of siRNA oligos resulting from cleavage of long double-stranded RNA (dsRNA) with an endoribonuclease such as *Escherichia coli* RNase III or dicer. The term “silencer” is not limited to any one particular methodology, unless so specified.

[0050] “Exosomes” what is meant herein are cell-derived vesicles that are present in many and perhaps all biological fluids, including blood, urine, and cultured medium of cell cultures.

[0051] As used herein, the expressions “cell”, “cell line” and “cell culture” are used interchangeably and all such designations include progeny. Thus, the words “cells” and similar designations include the primary subject cell and cultures derived therefrom without regard for the number of transfers. It is also understood that all progeny may not be precisely identical in DNA content, due to deliberate or inadvertent mutations that arise after genetic engineering is concluded. Mutant progeny that have the same function or biological activity as screened for in the originally transformed cell are included. Where distinct designations are intended, it will be clear from the context.

[0052] The terms “operably associated” or “operably linked,” as used herein, refer to functionally coupled nucleic acid sequences.

[0053] As used herein “recombinant” is relating to, derived from, or containing genetically “engineered” material. In other words, the genome was intentionally manipulated by the hand of man in some way.

[0054] “Reduced activity” or “inactivation” is defined herein to be at least a 75% reduction in protein/gene activity, as compared with an appropriate control species.

[0055] Preferably, at least 80, 85, 90, 95% reduction in activity is attained, and in the most preferred embodiment, the activity is eliminated (100%). Proteins can be inactivated with inhibitors, by mutation, or by suppression of expression or translation, and the like. A negative superscript, as in ACT, indicates reduced activity.

[0056] As used herein, “pharmaceutically acceptable carrier” refers to any carrier that is capable of delivering oligonucleotide to target cells. Examples of the pharmaceutically acceptable carrier include, but not limited to, nucleic acid carrier, cationic lipids, peptide-mediated carrier such as cell-penetrating peptides, mannogal carrier, liposomes, small molecule tags (including cholesterol-modification, membrane-permeant peptides, folate, antibiotics, VITE, and VITA), and cationic polymers.

[0057] As used herein the specification, “a” or “an” may mean one or more. As used herein in the claim(s), when used in conjunction with the word “comprising”, the words “a” or “an” may mean one or more than one. As used herein “another” may mean at least a second or more.

[0058] The term “about” means the stated value plus or minus the margin of error of measurement or plus or minus 10% if no method of measurement is indicated.
The use of the term “or” in the claims is used to mean “and/or” unless explicitly indicated to refer to alternatives only or if the alternatives are mutually exclusive.

The terms “comprise,” “have,” “include” and “contain” (and their variants) are open-ended linking verbs and allow the addition of other elements when used in a claim.

Wherever any of the phrases “for example,” “such as,” “including” and the like are used herein, the phrase “and without limitation” is understood to follow unless explicitly stated otherwise. Similarly “an example,” “exemplary” and the like are understood to be non-limiting.

The term “substantially” allows for deviations from the description that do not negatively impact the intended purpose. Descriptive terms are understood to be modified by the term “substantially” even if the word “substantially” is not explicitly recited. Therefore, for example, the phrase “wherein the lever extends vertically” means “wherein the lever extends substantially vertically” so long as a precise vertical arrangement is not necessary for the lever to perform its function.

The disclosure may use one or more of the following abbreviations:

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>ASO</td>
<td>Anti-sense Oligonucleotides</td>
</tr>
<tr>
<td>ASO</td>
<td>Adipose tissue derived stem cells</td>
</tr>
<tr>
<td>bi-siRNA</td>
<td>bifunctional siRNA</td>
</tr>
<tr>
<td>esiRNA</td>
<td>Endonuclease-prepared siRNAs</td>
</tr>
<tr>
<td>GFP</td>
<td>Green fluorescent protein</td>
</tr>
<tr>
<td>KD</td>
<td>Knock down (refers to silencers herein)</td>
</tr>
<tr>
<td>miRNA</td>
<td>microRNA</td>
</tr>
<tr>
<td>MSC</td>
<td>Mesenchymal stem cells</td>
</tr>
<tr>
<td>RAB27A</td>
<td>RAS-ASSOCIATED PROTEIN 27A (UniProt PS1159)</td>
</tr>
<tr>
<td>RFP</td>
<td>Red fluorescent protein</td>
</tr>
<tr>
<td>RNAi</td>
<td>RNA interference</td>
</tr>
<tr>
<td>shRNA</td>
<td>Small hairpin RNA</td>
</tr>
<tr>
<td>siRNA</td>
<td>Small interfering RNA</td>
</tr>
<tr>
<td>TRAF3IP2</td>
<td>TRAF3-INTERACTING PROTEIN 2 aka NUCLEAR FACTOR KAPPA-B ACTIVATOR 1 or ACT1 (UniProt P043734)</td>
</tr>
</tbody>
</table>

BRIEF DESCRIPTION OF THE DRAWINGS

Fig. 1 shows the localization of MSCs in tumor location. MDA-MB231 of genetically modified GFP expressing cells were injected intra mammary in 4-6 week old NIHIII immune-deficient female mice (n=5). 5x10^5 MSC cells were injected into the tail vein of these animals, which were euthanized 7 weeks post injection. The tumor tissues were extracted, fixed, and subjected to immunohistochemistry using HLA antibody to detect the human cells and DAPI for staining DNA. The samples were imaged with Leica confocal microscope (10X).

Fig. 2 shows the effect of exosomes on the gene expression of MSCs. To study the effect of exosomes on MSCs’ gene expression, MSCs were incubated with purified exosomes derived from MDA-MB231 cells (MDAexo) for 14 h in 37°C C. and 5% CO2. The changes in gene expression in MSCs were assessed using PCR array. The perturbed genes that displayed greater than two fold changed expression were grouped based on their function of cell adhesion (A), extracellular matrix proteins (B), cell growth and proliferation (C), and cell cycle (D). The graphs are representatives of triplicate experiments (P<0.05).

Fig. 3A shows intra-mammary engraftment of MSCs. The MSCs were exposed to purified MDAexo for 14 hours and then 5x10^5 cells (in Matrigel) were engrafted into mammary glands of NIHIII nude mice (female, 6-8 weeks old). The animals were observed for tumor growth weekly and euthanized after 12 weeks. Panels (from left to right) show the animals injected with PBS, Matrigel, and unexposed MSCs as controls. The MSC-exposed animals develop tumors at the site of injection versus no visible tumors growing on the controls. The euthanized animals were dissected at week 12 post-engraftment. Histology on the tumor tissue show positive immuno-reaction to penecytokeratin E-cadherin antibodies (Leica, 20X).

Fig. 3B shows methylation levels increased in exposed MSCs. MSCs exposed to MDA-MB231 culture condition media and MDAexo show enhanced levels of methylation; the methylation levels are reversible when exposed MSCs are treated with 5-Aza-2’-deoxycytidine (n=5, P<0.05).

Fig. 3C shows methylated genes in MSCs. Using PCR array, the methylated genes were identified in MSCs exposed to MDA-MB231 culture condition media and MDAexo (n=3, P<0.05).

Fig. 4A shows the expression of TRAF3IP2 and RAB27A in cultures of MDA-MB231, 184A1 and MSCs. The expression of both TRAF3IP2 and RAB27A are significantly higher in MDA-MB231 cells than in 184A1 and MSCs. The co-cultures of 184A1 and MSCs with MDA-MB231 cells enhanced the expression of TRAF3IP2 and RAB27A in both 184A1 and MSCs.

Fig. 4B Using silencing RNA, the expression of TRAF3IP2 and RAB27A were silenced in MDA-MB231, 184A1 and MSCs. The doubling time was calculated and compared to wild type cells. The silencing of RAB27A and TRAF3IP2 decreases the proliferation of MDA-MB231 cells, while having no effect on MSC replication capacity.

Fig. 4C Using a protein array technique, the cytokines released in culture media (CM) of MDA-MB231 and MDAexo cells were assessed. Cytokine array analysis shows that the level of cytokines mostly involved in breast cancer progression and metastasis are significantly reduced in MDAexo cells (n=3; P<0.05).

Fig. 5 shows the effect of silencing TRAF3IP2 and RAB27A on tumor cells. MDA-MB231 cells were silenced for the expression of TRAF3IP2 and RAB27A, then the selected gene expression was assessed using PCR array and compared to wild type cells set as zero in the graphs. The perturbed genes that displayed greater than two fold changed
expression were grouped based on their function of cell adhesion (A), transcription factors (B), cell growth and proliferation (C), and extracellular matrix proteins (D). The graphs are representatives of triplicate experiments (P<0.05, *P<0.001). Panel (E) shows an electron micrograph of MDA-MB231, MDAKDRAP2F2M and MDAKDRAP2F4 cells. The cells were negatively stained using uranyl acetate and viewed by electron microscopy. The scale bar represents 200 nm.

[F0073] FIG. 6 demonstrates altering tumor microenvironment formation in vivo. A. 1×10⁶ MDAKDRAP2F2M and MDAKDRAP2F4 cells in PBS and Martigel were injected into mammary in NIH/III female mice (4-6 weeks old). As controls, a group of animals were injected with 1×10⁶ MDA-MB231 cells in PBS and Martigel, another group was injected with Martigel, and another group was injected with PBS (n=15/group). Tumor growth was measured, and control animals injected with MDA-MB231 cells were euthanized 8 weeks post-injection. Animals injected with MDAKDRAP2F2M cells were euthanized 30 weeks post-injection for further analysis. MDAKDRAP2F2M-injected animals showed minimal tumor growth and were euthanized on week 52 of injection for further analysis. B shows a graph illustrating the survival of animals injected with MDA-MB231, MDAKDRAP2F2M and MDAKDRAP2F4 cells (P<0.05).

[F0074] FIG. 7 shows graphs of xenograft tumor weight and volume. Animals injected with MDA-MB231, MDAKDRAP2F2M and MDAKDRAP2F4 cells were sacrificed and tumors were isolated and weighted. A illustrates tumor weight and B displays tumor volume in injected animals at different time points.

[F0075] FIG. 8 shows the sequence of several silencer sequences, or provides an accession number for some variation.

[F0076] FIG. 9 shows tumor volume in treated animals with lentiviral vector carrying silencing sequence for TRAF1P2 (LentiKDRAP2F2).

[F0077] FIG. 10 is a photograph of U87 cells (a glioblastoma cell line) transduced with lentiviral vector carrying a silencing sequence for TRAF1P2 and GFP (green).

[F0078] FIG. 11 shows a scanning electron micrograph showing morphological changes in U87KDRAP2F2 compared to wild type U87.

[F0079] FIG. 12 is a cell proliferation assay showing a slight decrease in U87KDRAP2F2 cell proliferation, as compared to the U87 wild type cell.

[F0080] FIG. 13 shows TRAF1P2 gene (A) and Protein (B) expression levels. Wild type U87 were transduced with scrambled silencer RNA (SVG12) and used as control in these experiments. Scrambled shRNA is a non-target silencer RNA, which is used as a control in these experiments.

[F0081] FIG. 14 is a cell cycle analysis of U87KDRAP2F2 and wild type U87.

[F0082] FIG. 15 shows the in vivo tumorigenesis of U87KDRAP2F2 cells. Tumor size was measured using caliper and volume calculated and plotted here against time.


[F0084] FIG. 17A. TRAF1P2 mRNA variants comparison.

[F0085] FIG. 17B. Antisense oligonucleotide design for TRAF1P2 silencing.

[F0086] FIG. 18. Selection and optimization of ASOs in targeting TRAF1P2 in glioblastoma cells.

[F0087] FIG. 19. Effect of silencing TRAF1P2 in a flank xenograft model. A. Suppression of glioblastoma tumors by TRAF1P2 shRNA-LV injected subcutaneously onto tumors compared to scrambled shRNA-LV injected tumors. Frequency of administration is shown in the graph. B. Tumor size was measured biweekly ("P<0.05; *P<0.001). C. Animals images for luciferase weekly. Immunohistochemical localization of TRAF1P2, caspase 8, Ki67, IL-8, and VEGF in tumors treated with TRAF1P2 shRNA-LV or scrambled shRNA-LV. Scale: 100 µm.

**Detailed Description of the Disclosure**

[F0088] Detailed descriptions of one or more preferred embodiments are provided herein. It is to be understood, however, that the present invention may be embodied in various forms. Therefore, specific details disclosed herein are not to be interpreted as limiting, but rather as a basis for the claims and as a representative basis for teaching one skilled in the art to employ the present invention in any appropriate manner.

[F0089] Furthermore, while the invention is exemplified in breast cancer cell lines injected into mice with particular vectors and silencers, this is for proof of concept only, and the methods are expected to work in many different tumors with a variety of silencer delivery methods and with a variety of silencer sequences.

[F0090] The following materials were used herein:

<table>
<thead>
<tr>
<th>Material</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>MDA-MB231</td>
<td>Human breast cancer cell line, available from Sigm-Aldrich.</td>
</tr>
<tr>
<td>MDAKDRAP2F2M</td>
<td>MDA-MB231 cells transfected with a lentiviral vector encoding a TRAF1P2 silencer.</td>
</tr>
<tr>
<td>MDAKDRAP2F4 cells</td>
<td>MDA-MB231 cells transfected with a lentiviral vector encoding RAB2A silencer.</td>
</tr>
<tr>
<td>184A1 cells</td>
<td>Human mammary gland cell lines, established by chemical transformation (ATCC® CRL-8798).</td>
</tr>
<tr>
<td>Lentiviral vector</td>
<td>A lentiviral-based vector (e.g., pl.KO.1-puro or pl.KO.1-puro-CMV-TurboGFP™), preferably having a transactivator inducible promoter, such as IL1B promoter which will be activated in presence of excessive amounts of IL1B within tumor microenvironment.</td>
</tr>
<tr>
<td>U87</td>
<td>U87 is a human primary glioblastoma cell line formally known as U-87 MG. It has epithelial morphology, and was obtained from a stage four 44 year-old cancer patient, and can be obtained from ATCC (HTB-14).</td>
</tr>
<tr>
<td>SVG12</td>
<td>SVG12 is scrambled silencer RNA construct in a lentiviral vector that functions as a control for transduction in these experiments.</td>
</tr>
</tbody>
</table>
Exosomes

[0091] Exosomes are the main insoluble components of the tumor microenvironment. Exosomes are small membrane
ous extracellular vesicles (40-140 nm in diameter) that are
released in extracellular space. In addition to production by
tumor cells, exosome-like vesicles are produced by various
non-malignant cell types. Structurally, these vesicles consist of
a lipid bi-layer membrane similar to the cellular membr
brane, proteins, including host specific proteins, miRNA,
microRNA (miRNA) and transcription factors.

[0092] Exosomes can affect various cell types by trans
ferring their content to various cells. The growing interest
in the characterization of exosome-like vesicles in cancer
research arises from their potential role in carrying a large
array of oncoenic elements released by malignant cells,
such as oncoenic proteins and miRNAs. Such oncoenic
proteins and miRNAs can be transported in the tumor micro
vironment and can be taken up by recipient non-malignant cells;
this can result in the transfer of oncoenic activity.

[0093] It has been shown that the release of exosomes into
extracellular spaces is through exocytosis. RAB27A is one
of the exocytosis regulators. RAB27A, a membrane-bound
protein, is thought to be important for directing secretory
lysosomes to the immunological synapse and for their release
from mastocytes. At the membrane, RAB27A is activated
by exchange of bound nucleotide GDP for GTP. Active
RAB-GTP then recruits effector proteins from the cytosol to
the membrane. These are a diverse group of proteins that
include lipid kinases and phosphatases, molecular motors,
and tethering factors, which are involved in protein transport
and small GTPase mediated signal transduction.

[0094] A tumor can neither grow nor metastasize without
the development of supporting stroma. In solid tumors, the
associated stroma consists of a mixture of several cell types,
cytokines, chemokines, and extracellular exosome-like vesicles. These accumulations change the function and com
position of tissue surrounding the cancer cells and form the
tumor microenvironment. As noted above, the tumor
microenvironment contains both cellular and acellular frac
tions. The acellular fraction, consisting mainly of soluble
inflammatory cytokines and insoluble extracellular exo
somes-like vesicles, is involved in tumor-related inflamma
tion and growth. Tumor cells take part in releasing both
cytokines and exosomes into the tumor microenvironment
via exocytosis. Exocytosis is a cellular process that directs
the contents of secretory vesicles out of the cell membrane
and into the extracellular space. This process is regulated
mainly by the function of the RAB27A gene.

[0095] Mesenchymal stem cells (MSCs) are a type of
stromal cells abundant in the tumor microenvironment.
MSCs have been identified in several tissues. Adipose tissue
and bone marrow have been described among the major
sources of MSCs in adults. MSCs resemble fibroblasts in
terms of shape and markers; they are capable of self-renewal
and contribute to tissue regeneration by differentiation into
osteoblasts, chondrocytes, adipocytes, myocytes, macro
phage-like cells and myofibroblasts, depending upon the
requirements of the site to which they are recruited.

[0096] MSCs have been found to be incorporated into
tumors as well as in inflammatory milieu, such as healing
wounds. In tumor biology, the homing of MSCs to tumors is
the most significant hallmark of these cells. Several reports
have indicated that MSCs are capable of homing to the
tumor site, but results of current studies investigating the
signals that recruit MSCs to developing tumor sites are
controversial. During the normal homing process, which is
common to both hematopoietic stem cells (HSCs) and
MSCs, the cells migrate from their locations via proteolysis
and are directed to a particular injury site. Reports indicate
that MSCs are recruited to tumor sites in the same fashion.
This tropism of MSCs has been exploited for gene therapy
and delivering drugs in a targeted way to the tumor site, and
we have also used this tropism herein.

[0097] A recently published report described the effect
cytokines exert in recruitment of MSCs to the tumor site in
breast cancer (Mühlberg et al. 2009, Gehmert et al. 2010,
Senst et al. 2013, Ilner et al. 2014). It also showed that
co-culturing MSCs and MDA-MB231 cells (MSC+MDA-
MB231) enhances the expression of cytokines from tumor
cells. GRO-α, IL-6, IL-8, CXCL1 and MCP1 are chemoca
tractant proteins. As these chemotactants are released at a
high level when MSCs and cancer cells are in proximity they
have a significant effect in MSC homing towards tumor cells
(Id.).

[0098] To study the homing capability of MSCs into a
tumor site in vivo, genetically modified GFP-expressing
MDA-MB231 cells were injected intra-mammary into 4-6
weeks old NIHIII immune-deficient female mice (n=5).
5x10^5 MSC were injected into the tail vein of these ani
mals. The tumor tissues were extracted seven days following
MSCs injections, and the tumor tissue was harvested, fixed,
and subjected to immunohistochemistry using HNA anti
body to detect the human cells and DAPI for staining DNA.

[0099] FIG. 1 shows the homing of MSCs in tumor site.
These experiments confirm that MSCs are highly suitable to
be used as delivery agents to deliver silencers to the tumor site
as they preferably engraft to the tumor because they are
attracted by respective cytokines produced and released by
the tumor cells.

MSC Effect on Tumors

[0100] MSCs contribute to tumor growth in a number of
ways, including their roles in expressing growth factors and
enhancing vessel formation. Data has shown that the tumor
microenvironment modifies MSCs’ properties toward pro
moting breast cancer and metastasis, especially for MSCs
residing in breast adipose tissue, called adipose derived stem
cells or “ASCs”.

[0101] To study the effect of insoluble factors on stromal
cells, including MSCs, exosomes were purified from cul
tures of MDA-MB231 cells. MSCs were incubated with
purified exosomes from MDA-MB231 cells (MDAexo) for
14 hours in 37° C. and 5% CO₂. The changes in the gene
expression in MSCs were assessed, and the graphs illus
trated in FIG. 2A-2D shows the genes modified following
MSCs exposure to MDAexo.

[0102] The exosome exposed MSCs (5x10^5) are called
MDAexo, herein and were injected intra-mammary into
NIHIII immune-deficient mice. The animals developed a
growing tumor-like mass at the site of injection within 12
weeks, as shown in FIG. 3A. Exposure to either MDA-
MB231 culture condition media or to MDAexo enhances the
methylation in MSCs, as seen in FIG. 3B. The methylation
level was reversible when MDAexo-exposed MSCs were
N
treated with 5-Aza-2’-deoxycytidine. The gene expression
analysis showed several genes, including BRCA1, PAXS,
and APC, were highly methylated, as shown by FIG. 3C.
Silencing TRAF3IP2 and/or RAB27A

Tumor microenvironment components that are initially released from breast cancer cells activate the key transcription factors in inflammatory and stromal cells, similar to those described in breast cancer cells. This leads to the production and release of inflammatory mediators, which proceed to trigger cancer-related inflammation. The IKK/NF-κB signaling pathway has been shown to transcriptionally regulate inflammatory cytokine expression, and both IKK and NF-κB have been targeted to reduce cancer-related inflammation in the tumor microenvironment. However, these approaches were unsuccessful due to the activation of alternative pathways such as Toll-like receptors (TLRs).

TRAF3IP2 encodes ACT1, a signaling adaptor involved in the regulation of adaptive immunity. Other possible pathways involving TRAF3IP2 are shown in FIG. 16. Studies of TRAF3IP2-deficient mice suggest that TRAF3IP2 is a negative regulator of humoral immunity through its inhibitory effect on CD40- and BAFFR-mediated signaling. TRAF3IP2 operates as a positive signaling adaptor in IL-17-mediated cellular immune responses. IL-17 is a dominant ‘signature’ cytokine of TH-17 cells and up-regulates neutrophil-mobilizing cytokines, chemokines, and tissue-degrading matrix metalloproteases.

IL-17-dependent receptor ligation induces TRAF3IP2 recruitment to the cytoplasmic tail of the IL-17R. This in turn allows the incorporation of the TNF receptor-associated factors TRAF3 and TRAF6 into the signaling complex and the subsequent downstream activation of the MAPK and NF-κB pathway. Accordingly, TRAF3IP2 is not only involved in pathways balancing humoral and cellular immunity, but also represents a chief link between IL-17-mediated adaptive immune responses and NF-κB as the master regulator of innate immunity controlling the inducible transcription of various pro-inflammatory cytokines.

The data presented herein indicates that TRAF3IP2 mediates IKK dependent NF-κB activation as well as TLR4 signaling. It has been shown that IL-17 signals exclusively via TRAF3IP2, and TRAF3IP2 gene deletion abrogates IL-17-dependent inflammatory signaling. The novel findings of the present disclosure show a significantly high expression of TRAF3IP2 in breast cancer cells while this expression is minimal in non-malignant breast epithelial cells and MSCs.

Interestingly, the data presented here also show that the expression of RAB27A is also significantly higher in breast cancer cells compared to 184A1 cells, a non-malignant breast epithelial cell line, and MSCs, as shown in FIG. 4A. The silencing of RAB27A and TRAF3IP2 decrease the cell proliferation in MDA-MB231 cells, while the silencing of these genes has no effect on MSC replication capacity, as seen in FIG. 4B.

Silencing TRAF3IP2 in MDA-MB231 cells (MDA_XDTRAF3IP2) results in remarkable changes in expression of cytokines. Cytokine array analysis shows that the level of cytokines mostly involved in breast cancer progression and metastasis are significantly reduced in MDA_XDTRAF3IP2 cells, as shown in FIG. 4C.

Silencing TRAF3IP2 results in significant changes in the expression of factors involved in the formation of tumor microenvironment and associated inflammation. The tumor microenvironment is under constant chronic inflammatory pressure. It has been shown that one of the potent regulators of inflammation is TGIF-β which was found to regulate the expression of angiopeptin-like 4 (ANGPTL4) via a Smad3-signaling pathway. The up-regulation of ANGPTL4 in cancer cells when they extravasate into the circulatory system likely explains their inclination toward colonizing lung tissue. The rationale for this is based on the ability of ANGPTL4 to disrupt the integrity of vascular tight junctions, thereby increasing the permeability of the capillaries in the lung to promote the intravasation into the lung tissue.

The present data shows a significant reduction in ANGPTL4 expression in both MDA_XDRA657 and MDA_XDTRAF3IP2 as seen in FIG. 5B. The expression of ANGPT1, which binds to extracellular matrix from carcinogenic cells, is exclusively decreased in MDA_XDTRAF3IP2, while its expression is enhanced in MDA_XDRA657 cells, as shown by FIG. 5A-5D. This is due to the halt in exocytosis in MDA_XDRA657 cells. Electron microscopy indicates abnormal morphology in both MDA_XDRA657 and MDA_XDTRAF3IP2 cells.

Silencing In Vivo

The data presented above strongly suggests that silencing TRAF3IP2 and RAB27A could have potent effects in vivo, and thus, the next step was to deliver silencers to tumor cell using cancer cells that already contained the silencers.

In these experiments, the expression of TRAF3IP2 and RAB27A were silenced in MDA-MB231 cells using lentiviral-based vectors encoding silencer RNA. Female 4-6 weeks old NIHIII mice were injected intramammary with 1x10^5 MDA_XDTRAF3IP2 cells in PBS and Matrigel. Another group of animals were injected with 1x10^5 MDA_XDRA657 cells in PBS and Matrigel. As controls, a group of animals were injected with just 1x10^5 MDA-MB231 cells in PBS and Matrigel, another group was injected with PBS.

Earlier work showed that breast cancer cells exhibit significantly high levels of RAB27A expression and ultimately have higher exocytosis activity, as shown in FIG. 4A. These in vivo studies showed a decreased tumor volume in MDA_XDRA657 up to 30 weeks post-injection. The control group injected with MDA-MB231 cells showed tumor growth within 8 weeks and the animals were euthanized, as seen in FIG. 6A. Animals injected with MDA_XDTRA3IP2 cells survived up to 52 weeks with only limited tumor growth.

Compared to animals injected with MDA-MB231 cells, the survival studies also show a 30 and 52 weeks life span for animals injected with MDA_XDRA657 and MDA_XDTRAF3IP2 cells, respectively, as shown by FIG. 6B. These results demonstrate that reducing exocytosis in breast cancer cells attenuates the release of oncogenic molecules into the tumor microenvironment in both soluble and insoluble forms. Silencing TRAF3IP2 regresses tumor growth by reducing cytokine signaling.

Upon euthanizing the animals, the tumors were isolated and weighted to quantify the effects of the silencers. FIG. 7A shows the tumor weight and FIG. 7B shows the tumor size at 8 weeks post-injection of MDA-MB231 injected cells, at 30 weeks for MDA_XDTRAF3IP2 and MDA_XDRA657 cells, and at 52 weeks for MDA_XDTRAF3IP2.
injected cells. These data indicate a significant decrease in tumor growth following down-regulation of TRAF3IP2 and RAB27A.

[0116] Thus, the data establishes that silencing TRAF3IP2 and RAB27A in tumor cells prevents tumor growth and/or metastasis in vivo. Injection of MDA-MB231 cells results in metastasis within 8 weeks (data not shown). However, postmortem analysis of animals injected with MDA5DGR2A and MDA5DGR2A(p27) showed no metastasis at 30 weeks and 52 weeks post-injection (data not shown).

ΔTRAF3IP2

[0117] Delivery of gene silencers is one way of shutting down TRAF3IP2 and/or RAB27A, but knockouts are another possibility and also provide a good biological system in which to study the effects of silencing one or both of these genes.

[0118] Using the CRISPR/Cas system, the gene TRAF3IP2 and/or RAB27a can be knocked out. This strategy involves engineering specific nuclease (ex. CAS9-CRISPR) that are designed to create a DNA double-strand break (DS-break) in the e.g., TRAF3IP2 gene, thereby activating the cell’s endogenous homologous recombination repair pathway. Because the DS-break repair mechanisms are not accurate, changes are introduced into the gene by non-homologous end joining (NHEJ), which frequently lead to frame-shift mutations. In this system, CRISPR activation is under strict control of a promoter, such as the IPTG promoter (an analog of lactose). Induction of this promoter activates the TRAF3IP2-specific CRISPR and causes mutations. The delivery of the TRAF3IP2-specific CRISPER lentiviral vectors will be attained by injection to the tumor site. Once the knock-outs are obtained, they can be used in studies to elucidate the biology of this system. Other promoters specific to the respective tumor and under control of tumor-specific, unregulated pathways will also work.

TRAF3IP2 Silencing

[0119] The above experiments were performed in an animal model, which mimicked breast cancer tumors. However, those tumors were not localized in the body, but scattered throughout, and especially subcutaneously. On these experiments, we show that the effect is reproducible in wild type mammary fat tumors.

[0120] Tumors were generated in the mammary fat pad of female immune deficient NIHIII mice. For generating tumors, 5×10⁵ MDA-MB-231 cells were mixed with 50 μl Matrigel and injected into the mammary fat pad. Ten days after injecting the MDA-MB-231 cells, NIH-III mice were randomly divided into two groups. One group of animals was received direct injections of 100 μl lentiviral-vector carrying TRAF3IP2 silencer RNA (in PBS) to the tumor site. The other group (control group), the animals received 100 μl of PBS. Injected tumor volumes were evaluated twice a week by measuring two orthogonal diameters with digital calipers. Tumor volume (V) was calculated using the following equation: \( V = \frac{a \times b^2}{2} \), where “a” is the longer diameter and “b” the shorter diameter. FIG. 9. As can be seen, there was little or no tumor growth in the tumors injected with silencer encoding expression vectors. Although the data is not yet available for RAB27A, we predict the results will be similar.

Antisense Oligonucleotide

[0121] Oligonucleotides are unmodified or chemically modified single-stranded DNA molecules. In general, they are relatively short (13-25 nucleotides) and hybridize (at least in theory) to a unique sequence in cells. Anti-sense oligonucleotides (ASOs) are single strands of DNA or RNA that are complementary to a chosen sequence. In the case of antisense RNA they prevent protein translation of certain messenger RNA strands by binding to them. If binding takes place, this DNA/RNA hybrid can be degraded by the enzyme RNase H. While the oligonucleotide may be susceptible to rapid degradation by nucleases, a 2′-methoxyethyl (2′-MOE) modified or 2′-O-methyl (2′-OMe) modified ASO is resistant to nucleases and has enhanced target binding and pharmacokinetics comparing to DNA. Therefore, the ASOs employed herein can be 2′-MOE or 2′-OMe modified or unmodified.

[0122] The inventors therefore investigated the possibility of silencing TRAF3IP2 with oligonucleotides and especially with more degradation-resistant oligonucleotides, as well as the suitable target binding sites within the gene.

[0123] Several variants of NM_0147200.2 (SEQ ID NO. 8) were investigated, including XM_001153386.2 (SEQ ID NO. 9), NM_0147686.3 (SEQ ID NO. 10), NM_001164281.2 (SEQ ID NO. 7), XM_006715319.4 (SEQ ID NO. 11), and NM_001164283.2 (SEQ ID NO. 12). Highly conserved regions appear in these sequences, as shown in FIG. 17A, suggesting that target binding sequence may be available within the conserved regions. TRAF3IP2 gene has 10 exons, among which exons 9 and 10 are conserved across the known variants. By targeting the most conserved regions across all variants, these oligonucleotides are designed to be able to block most of the TRAF3IP2 activity. Therefore, anti-sense oligonucleotides (ASOs) can be designed to specifically target these exons to obtain more universal applicability.

TRAF3IP2 Silencing in Glioblastoma

[0124] The above experiments were performed using breast cancer cell lines, but we also hoped that the method might be applicable to other solid tumors, and thus tested a glioblastoma derived cell line to confirm.

[0125] TRAF3IP2 exhibit significant role in the onset of tumor microenvironment and metastasis in solid tumors including Glioblastoma. TRAF3IP2, a signaling adaptor involved in the regulation of adaptive immunity operates as a positive signaling adaptor in IL-17-mediated cellular immune responses. IL-17 is a dominant ‘signature’ cytokine of TII-17 cells and up regulates neutrophil-mobilizing cytokines, chemokines, and tissue-degrading matrix metalloproteases. IL-17-dependent receptor ligation induces TRAF3IP2 recruitment to the cytoplasmic tail of the IL-17RI. This in turn allows the incorporation of the TNF receptor associated factors (TRAF) TRAF3 and TRAF6 into the signaling complex and the subsequent downstream activation of the MAPK and NF-kB pathway. Accordingly, TRAF3IP2 is not only involved in pathways balancing humoral and cellular immunity, but it also represents a chief link between IL-17 mediated adaptive immune responses and NF-kB as the master regulator of innate immunity controlling the inducible transcription of various pro-inflammatory cytokines.
Previously, our group and others showed that TRAF3IP2 mediates IKK dependent NF-κB activation as well as TLR4 signaling. It has been shown that IL-17 signals exclusively via TRAF3IP2, and TRAF3IP2 gene deletion abrogates IL-17-dependent inflammatory signaling.

We have shown a significantly high expression of TRAF3IP2 in breast cancer cells while this expression is minimal in non-malignant breast epithelial cells and MSC’s. Our data indicate that similar to breast cancer, high amounts of TRAF3IP2 express in glioblastoma cells (data not shown). Herein, we have studied the effect of TRAF3IP2 silencing on in vitro and in vivo characteristics of a glioblastoma cell line (U87).

A human glioblastoma cell line “U87” cells were transduced with lentiviral vector carrying a silencing sequence for TRAF3IP2 and GFP as a detectable marker. As can be seen in FIG. 10, transduced U87 cells with lentiviral delivering silencer sequences for TRAF3IP2. A GFP expressing sequence was used as a reporter gene making transduced cells traceable. In addition, electron microscopic analysis showed morphological changes in U87KDTRAF3IP2 compared to wild type U87 (FIG. 11). These changes include a different cell morphology, which might be related to modified cellular function due to silencing TRAF3IP2.

A cell proliferation assay shown in FIG. 13 shows only a slight decrease in U87KDTRAF3IP2 cell proliferation, as compared with the control cell U87, which suggests that the effect of the silencer is not a direct effect on cell proliferation, but an indirect one on the interaction of the tumor cells with its microenvironment.

When we studied gene and protein expression levels, TRAF3IP2 expression was significantly reduced in both gene and protein levels in U87KDTRAF3IP2 compared to control U87 and U87 transduced with scrambled silencer RNA (up to 92.3%), confirming that the silencer was effective in these cells. The results are shown in FIGS. 14A and 14B which provides the TRAF3IP2 gene and protein expression, respectively.

Cell cycle analysis also showed significant changes in cell cycle profile in U87KDTRAF3IP2 compared to wild type U87. As seen in FIG. 15, silencing TRAF3IP2 caused higher G1 phase and lower populations in S and G2 phases, which might indicate a lower U87 replication rate.

To further test the theory and broaden the possible silencing alternatives, highly effective anti-sense oligonucleotides (ASOs) were designed to target TRAF3IP2. Six (6) exemplary ASOs were designed (SEQ ID NOs. 13-18), for example, ASO1 and ASO2 are shown in FIG. 17B. It is to be noted that the length of the ASOs can vary while achieving similar silencing effect.

The six ASOs were tested for their ability to silence the expression of TRAF3IP2 in glioblastoma cells (FIG. 18) using the lentiviral vector. As shown in FIG. 18, concentration-dependent silencing effects were demonstrated by all six ASOs. Among the six, ASO1, ASO2, ASO4, and ASO5 showed comparable or better silencing capability comparing to the control. The ASO1 significantly suppresses TRAF3IP2 in glioblastoma cells, compared to other possible ASOs.

Experiments can also be designed to test the silencing effect of the ASOs across different cell lines from different tissues, in order to validate their efficacy on the TRAF3IP2 variants. It is expected that similar silencing of TRAF3IP2 expression can be achieved because these ASOs are designed to target only the conserved regions across known variants.

It is also contemplated that personalized silencing can be achieved, as the cost of producing the silencing sequences will continue to decline. As such, the need to target only the conserved regions across TRAF3IP2 variants may be obviated. Certain regions within TRAF3IP2 may be even more susceptible to silencing on an individualized basis, and the inventive concept described herein can be readily applied.

In Vivo Tumorigenicity of U87KDTRAF3IP2

The above experiments strongly suggested that TRAF3IP2 silencer might also be effective in glioblastoma tumors, but the results needed to be confirmed in an in vivo system. Therefore, we created U87 glioblastoma-like tumors by injecting these cells into nude mice.

U87KDTRAF3IP2 and wild type U87 cells were injected subcutaneously in the upper portion of the right hind thigh. Tumors were measured with a traceable digital caliper (Fisher Scientific) for calculation of the tumor volume. The tumor size and volume were measured weekly. The animals injected with U87KDTRAF3IP2 showed a significantly smaller tumor size compared to control animals injected with wild type U87. The control animals were sacrificed 8 weeks following injection. The animals injected with U87KDTRAF3IP2, which were sacrificed on week 14 post-injection, showed significantly smaller tumor volume.

This confirms that TRAF3IP2 silencing can also slow glioblastoma tumor growth in vivo.

Treatment of U87 tumors are also under investigation. The efficacy of ASOs will be tested by following a protocol for obtaining and implanting tumors, and for data collection. However, targeting modality will be injections of ASO. After confirming tumors 14 days after initial implantation, animals will be treated with TRAF3IP2-ASO, which we have designed and optimized for targeting efficacy and efficiency) through IV injections in the lateral ventricle (5 μl every 48 h for 40 days; 30 nM concentration). The animals will be followed for up to 32 weeks post-tumor induction. Scrambled-ASO will serve as a control. Preliminary data shows a significant decrease in tumor development and growth on treated animals with lentiviral carrying silencer RNA injected to tumor site.

Therapeutic Significance of Targeting TRAF3IP2 in the Regression of Pre-Existing Glioblastoma

Having demonstrated that TRAF3IP2-silenced malignant U87 glioblastoma cells form significantly smaller tumors, we next determined whether treating existing tumors by lentiviral TRAF3IP2 shRNA regresses their size. In this translationally important strategy, tumors were induced at first by injecting luciferase-labeled U87 cells into the flank region of immunodeficient NIH-III mice. Fourteen days later, when tumors were distinctly quantifiable, lentivirus expressing GFP-tagged TRAF3IP2 shRNA (TRAF3IP2 shRNA-LV) was injected subcutaneously onto the tumors. Scrambled shRNA-LV served as a control. Results in FIG. 19A show a remarkable reduction in tumor size over 50 days post-induction in TRAF3IP2 shRNA-LV-treated animals (versus scrambled shRNA-LV; 0.08±0.03 g
versus 1380±48, respectively) (FIG. 19B). Analysis of residual tumors by IHC revealed a marked reduction in TRAF3IP2, Ki67, IL-8 and VEGF expression (FIG. 19C), but a significant increase in caspase 8 levels (FIG. 19C).

[0140] The results show that treating existing tumors formed by the wild type U87 glioblastoma cells with TRAF3IP2 shRNA significantly reduces tumor size in the flank xenograft model.

TRAF3IP2 Silencing in Glioblastoma Angiogenesis

[0141] It is also reported that TRAF3IP2 contributes to angiogenesis, an important aspect of tumor growth and metastasis. Inhibiting angiogenesis has been widely reported to show hope in tumor treatment. Therefore, silencing TRAF3IP2 could play a major role in controlling or reducing tumor size and reduce metastasis.

[0142] Our results indicate that silencing TRAF3IP2 leads to reduced expression of VEGFA (12-fold reduction), a key growth factor in angiogenesis. Additional in vitro experiments also showed that TRAF3IP2 silencing significantly reduced angiogenesis in U87 cells. It is therefore proposed that silencing TRAF3IP2 could be used to treat tumors expressing high level of TRAF3IP2, as well as preventing its metastasis.

[0143] Future experiments include studies to confirm silencer delivery to mammary tumors, with preferred delivery agents such as MSCs. Of course, clinical studies will be performed eventually to confirm efficacy of these methods in humans, but these experiments are expected to take several years.

The following reference are incorporated by reference herein in its entirety for all purposes:


[0151] WO2014030602 An agent for treating cancer


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What is claimed is:

1. A pharmaceutical composition for the treatment of a tumor having increased expression of TRAF3IP2, wherein said composition comprises at least one silencing sequence for TRAF3IP2 in a pharmaceutically acceptable carrier in an amount effective for the therapeutic treatment of a tumor, wherein said silencing sequence reduces the expression of the TRAF3IP2 gene by at least two-fold as compared to without the silencing sequence for TRAF3IP2, and wherein said silencing sequence is a modified portion of sense strand of NM_001164281.2 (SEQ ID NO. 7), NM_147200.2 (SEQ ID NO. 8), XM_011535386.2 (SEQ ID NO. 9), NM_147686.3 (SEQ ID NO. 10), XM006715319.4 (SEQ ID NO. 11), and NM_001164283.2 (SEQ ID NO. 12).

2. The composition of claim 1, wherein said composition comprises an expression vector encoding a TRAF3IP2 silencer operably coupled to an inducible promoter.

3. The composition of claim 1, wherein said silencing sequence is an siRNA, an miRNA, an shRNA, an antisense RNA, or an antisense oligonucleotide.

4. The composition of any of claim 1, said silencing sequence encoded by an expression vector hosted in a mesenchymal stem cell (MSC) that targets said tumor.

5. The composition of claim 4, said MSC having been previously exposed to exosomes from said tumor.

6. The composition of claim 3, wherein said silencing sequence is an antisense oligonucleotide that is 13-25 nucleotides in length.

7. The composition of claim 6, wherein the antisense oligonucleotide is complementary to a portion of the sense strand of any one of SEQ ID NOs. 7-12.

8. The composition of claim 6, wherein the antisense oligonucleotide is selected from SEQ ID NOs. 13-18.

9. The composition of claim 1, wherein the pharmaceutically acceptable carrier is a nucleic acid carrier.

10. The composition of claim 1, further comprising a silencing sequence for Rab27a.

11. The composition of claim 1, wherein said composition is formulated for parenteral administration, including direct injection into a tumor or its metastasis site by transeutaneous, intraarticular, intraductal, intravenous, intradermal, intramuscular, intraperitoneal, or subcutaneous administration.

12. The composition of claim 1, wherein the composition is used in treating glioblastoma or breast cancer, or for use in treating any cancer with at least 2-fold increased TRAF3IP2 and/or Rab27a expression.

13. A method of treating at least one tumor in a mammal comprising administering to the mammal an effective amount of the composition of claim 1.

14. The method of claim 13, wherein said tumor is a solid tumor or breast cancer or a glioblastoma or a cancer with at least 2 fold increased expression of TRAF3IP2 or Rab27a or both.

15. The method of claim 13, wherein the composition is injected directly into said tumor and said injection is guided by ultrasound, fluoroscopy, imaging, CT, MM, or visually, in order to enhance the local concentration of the silencer within the tumor.

16. A method to selectively treat a tumor and minimize side effects, by administering an effective amount of a silencer for TRAF3IP2 or Rab27a, or both, to a tumor that expresses at least 10 times the amount of TRAF3IP2 or Rab27a, or both, as compared to a non-tumor cell from the surrounding tissue.

17. The method of claim 16, wherein said silencer is an siRNA, an miRNA, an shRNA, an antisense RNA, or an antisense oligonucleotide.

18. The method of claim 16, further comprising enhancing the selective effect on tumor cells and avoiding effects on non-tumor cells by increasing the local concentration of the silencer within the tumor by injecting said silencer(s) directly into said tumor.

19. The method of claim 16, wherein said silencer(s) is encoded in an expression vector having an inducible promoter, thus enhancing the selective effect on tumor cells and avoiding effects on normal cells by selectively activating the production of the silencer by a switch that activates said inducible promoter.

20. The method of claim 16, wherein said silencer(s) is linked to an antibody, or other targeting substance specific to said tumor.

21. A silencing sequence of TRAF3IP2 for use as a medicament or for use in treating a tumor, or solid tumor, or for use in treating glioblastoma or breast cancer, or for use in treating any cancer with at least 2-fold increased TRAF3IP2 expression, wherein said silencing sequence is a modified portion of sense strand of NM_001164281.2 (SEQ ID NO. 7), NM_147200.2 (SEQ ID NO. 8), XM_011535386.2 (SEQ ID NO. 9), NM_147686.3 (SEQ ID NO. 10), XM006715319.4 (SEQ ID NO. 11), and NM_001164283.2 (SEQ ID NO. 12).