



2015; 6(12): 1306-1319. doi: 10.7150/jca.13266

Research Paper

Comprehensive Two- and Three-Dimensional RNAi Screening Identifies PI3K Inhibition as a Complement to MEK Inhibitor AS703026 for Combination Treatment of Triple-Negative Breast Cancer

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Received: 2015.07.16; Accepted: 2015.08.08; Published: 2015.10.28

Abstract

Triple-negative breast cancer (TNBC) is a major cause of death among breast cancer patients that results from intrinsic and acquired resistance to systemic chemotherapies. To identify novel targets for effective treatment of TNBC through combination strategies with MEK inhibitor (AS703026), we used a novel method of combining high-throughput two- and three-dimensional (2D and 3D) RNAi screening. TNBC cells were transfected with a kinome siRNA library comprising siRNA targeting 790 kinases under both 2D and 3D culture conditions with or without AS703026. Molecule activity predictor analysis revealed the PI3K pathway as the major target pathway in our RNAi combination studies in TNBC. We found that PI3K inhibitor SAR245409 (also called XL765) combined with AS703026 synergistically inhibited proliferation compared with either drug alone (P < 0.001). Reduced *in vitro* colony formation (P < 0.001) and migration and invasion ability were also observed with the combination treatment (P<0.01). Our data suggest that SAR245409 combined with AS703026 may be effective in patients with TNBC. We conclude that a novel powerful high-throughput RNAi assays were able to identify anti-cancer drugs as single or combinational agents. Integrated and multi-system RNAi screening methods can complement difference between *in vitro* and *in vivo* culture conditions, and enriches targets that are close to the *in vivo* condition.

Key words: Triple-negative breast cancer, two-dimensional RNAi screening, three-dimensional RNAi screening, SAR245409, AS703026, multicellular tumor spheroid, projection formation.

Introduction

Triple-negative breast cancer (TNBC) (in which estrogen receptor, progesterone receptor, and HER2 are not overexpressed) is characterized by distinct clinical and pathologic features, metastatic behavior, a

high recurrence rate, and poor prognosis compared with other breast cancer types [1-3]. TNBC does not respond to receptor-targeted treatments such as hormonal therapy and trastuzumab, leaving chemotherapy as the mainstay of treatment. It is now increasingly appreciated that combinatorial therapies are an essential path to achieving reliable treatment efficacy in TNBC. To facilitate such a multipronged approach, we need to find drugs that work together to modulate/target multiple hallmarks of cancer cell behavior.

Two-dimensional (2D) high-throughput kinome RNA interference (RNAi) screening has become a useful method for exploring the cellular signaling pathways that underlie cancer cell biology and for identifying target molecules for treatment [4]. Despite the powerful strategy of 2D high-throughput kinome RNAi screening in biological studies [5, 6], 2D monolayer cell growth model lacks the key component of cancer - the tumor microenvironment that can be tested in vivo [7, 8]. Three-dimensional (3D) cell cultures exhibit a higher degree of structural complexity and homeostasis, analogous to tissues and organs [9], and are currently used in a broad range of cell biology studies, including tumor biology studies [10]. Thus, we hypothesized that comprehensive RNAi analysis with 2D viability and 3D morphological assays would identify targets with better significance that can be a representative of clinical target for TNBC treatment. To test this hypothesis, we studied the role of gene knockdown under both 2D and 3D culture conditions to synergistically modulate the cancer growth phenotype in conjunction with a MEK inhibitor (AS703026). MDA-MB-231 cells were grown as either a 3D multicellular tumor spheroid (MCTS) cell culture system or as a 2D monolayer cell culture system. MDA-MB-231 cells demonstrate a unique 3D morphology comprising projections suggestive of epithelial-mesenchymal transition (EMT)-mediated invasion processes that are related to the cell line's spontaneous metastasis capability [11, 12].

The objective of this screen was to identify targets whose silencing by siRNA sensitized the TNBC cells to the MEK inhibitor, enhancing cell killing. Rather than focusing solely on a viability assay in 2D culture, we carried out a parallel screen utilizing a 3D culture that more closely resembled the *in vivo* environment to identify modulators of morphological formation of projections/filopodia. We identified and validated the PI3K pathway as a therapeutic target for combination with MEK inhibitor AS703026, a treatment strategy that is being assessed in phase I clinical trial expansion cohorts for TNBC patients.

Results

Primary Screen

We first analyzed 760 kinases targeted by the human siRNA kinase library to identify gene profiles

for the 2D monolayer cell culture and the 3D MCTS culture (Figure 1). In both types of culture, we identified genes that sensitized the cells to the MEK inhibitor, inducing greater than 50% cell growth inhibition or killing compared to cells treated with the solvent/vehicle.

Classification Results

confusion matrix for the long-projection phenotype classifier is shown in Table 1. The entries in each cell represent the percentage of instances that were predicted by the computer model (divided into columns for each phenotype) and the actual instances (shown by rows). Table 2 shows the contingency table based on examining the intersection between the 2D and 3D screen outputs. We found that 5.1% of the genes conferred a growth inhibition phenotype in 2D culture and the "no-projection" phenotype in 3D culture. The panels of genes whose silencing induced this sensitization varied significantly between the 2D and 3D cell systems. The 3D model was superior as it enabled us to identify a panel of genes that had an effect on invasion as well as viability (Table 3).

Table 1. The confusion matrix representing the performance of the no- vs long-projection phenotype classifier for 3D culture results. rows represent the actual labels (based on human expert annotation) of the cells' morphology, whereas columns represent the labels predicted by the random forest classifier. The accuracy of the classifier was 91%.

Predicted	0 (No)	1 (Long)
Actual		
0 (No)	25.2%	5.6%
1 (Long)	3.1%	66.1%

Table 2. Contingency table based on examining the intersection between 2D and 3D RNAi screening. The entries denote the number of genes that confer the indicated morphological designations under 2D and 3D culture conditions (eg: 48.9% of the genes conferred the "no growth inhibition" phenotype in 2D culture but the "long-projection" phenotype in 3D culture). The determination of 3D morphology was made based on decisions outputted by the morphology classifier shown in Table 1.

3D culture	Long	No		
2D culture	projection	projection		
0 (Growth inhibition)	42.6%	3.4%		
1 (No growth inhibition)	48.9%	5.15.1%		

The entries in each cell indicate the number of genes that have the characteristics in the type of culture shown by the column and row headings.

Table 3. Genes whose silencing led to a no-projection phenotype in the 3D screen, grouped by the outcome of the 2D siRNA screen.

Gene symbol	Accession no.	Gene name
	tion outcome in 2D/ no projection	
ADRBK2	NM_001619	adrenergic, beta, receptor kinase 1
ANGPT4	NM_015985	angiopoietin 4
ATR	NM_001184	ataxia telangiectasia and Rad3 related
CDKN3 CKMT1	NM_005192	cyclin-dependent kinase inhibitor 3
CKMT2	NM_020990 NM_001825	creatine kinase, mitochondrial 1B creatine kinase, mitochondrial 2
CNKSR1	NM 006314	connector enhancer of kinase suppressor of Ras 1
DUSP1	NM 004417	dual specificity phosphatase 1
DUSP10	NM 007207	dual specificity phosphatase 10
DYRK1B	NM_004714	dual-specificity tyrosine-(Y)-phosphorylation regulated kinase 1B
EPHA1	NM_005232	EPH receptor A1
FLJ10761	NM_018208	Ethanolamine kinase 2 (ETNK2)
FLJ34389	NM_152649	Mixed lineage kinase domain-like (MLKL), transcript variant 1
FUK	NM_145059	Fucokinase
MERTK	NM_006343	c-mer proto-oncogene tyrosine kinase
MGC4796	NM_032017	serine/threonine kinase 40 (STK40), transcript variant 3
MGC5601	NM_025247	acyl-CoA dehydrogenase family, member 10 (ACAD10), transcript variant 2
PAG	NM_018440	phosphoprotein associated with glycosphingolipid microdomains 1
PDK3	NM_005391	pyruvate dehydrogenase kinase, isozyme 3 (PDK3), transcript variant 2
PDPK1	NM_002613	3-phosphoinositide dependent protein kinase-1 (PDPK1), transcript variant 1
PFTK1	NM_012395	cyclin-dependent kinase 14 (CDK14)
PIK3C2G	NM_004570	phosphatidylinositol-4-phosphate 3-kinase, catalytic subunit type 2 gamma
PKN3	NM_013355	protein kinase N3
PPP2CB	NM_004156	protein phosphatase 2 (formerly 2A), catalytic subunit, beta isoform
PRKAA2	NM_006252	protein kinase, AMP-activated, alpha 2 catalytic subunit
PAK4	NM_005884	p21 protein (Cdc42/Rac)-activated kinase 4
PRKCN	NM_005813	protein kinase D3
PRKCQ	NM_006257	protein kinase C, theta
PRPSAP1	NM_002766	phosphoribosyl pyrophosphate synthetase-associated protein 1
PTPRR	NM_002849	protein tyrosine phosphatase, receptor type, R
SNRK	NM_017719	SNF related kinase
SSTK	NM_032037	testis-specific serine kinase 6
STK19	NM_004197	serine/threonine kinase 19
STK31 STK35	NM_031414	serine/threonine kinase 31 serine/threonine kinase 35
STK36	NM_080836 NM_015690	serine/threonine kinase 36
STYK1	NM 018423	serine/threonine/tyrosine kinase 1
TEX14	NM_031272	testis expressed 14
TNK1	NM 003985	tyrosine kinase, non-receptor, 1
	n outcome in 2D/ no projection in	
CDC42BPA	NM_003607	CDC42 binding protein kinase alpha
CDK6	NM_001259	cyclin-dependent kinase 6
CDKN1B	NM_004064	cyclin-dependent kinase inhibitor 1B
MYO3A	NM_017433	myosin IIIA
NTRK3	NM_002530	neurotrophic tyrosine kinase, receptor, type 3
PAK6	NM_020168	p21 protein (Cdc42/Rac)-activated kinase 6
PANK4	NM_018216	pantothenate kinase 4
PAPSS1	NM_005443	3'-phosphoadenosine 5'-phosphosulfate synthase 1
PCTK1	NM_006201	cyclin-dependent kinase 16
PCTK3	NM_002596	cyclin-dependent kinase 18
PFKP	NM_002627	phosphofructokinase, platelet
PGK2	NM_138733	phosphoglycerate kinase 2
PHKG2	NM_000294	phosphorylase kinase, gamma 2
PIP5K1A	NM_003557	phosphatidylinositol-4-phosphate 5-kinase, type I, alpha
PRKCE	NM_005400	protein kinase C, epsilon
PRKD2	NM_016457	protein kinase D2
PRKWNK4	NM_032387	WNK lysine deficient protein kinase 4
PTK6	NM_005975	protein tyrosine kinase 6
RAC1	NM_018890	ras-related C3 botulinum toxin substrate 1
SGK2	NM_016276	serum/glucocorticoid regulated kinase 2
SPEC2	NM_020240	CDC42 small effector 2
SPHK1	NM_021972	sphingosine kinase 1
SPHK2	NM_020126	sphingosine kinase 2
STK25	NM_006374	serine/threonine kinase 25
TPK1	NM_022445	thiamin pyrophosphokinase 1
TXK	NM_003328	TXK tyrosine kinase

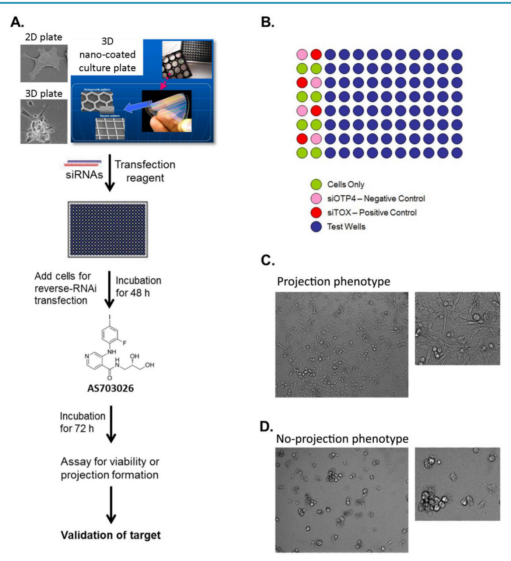


Figure 1. High-throughput RNAi screen using a 3D multicellular tumor spheroid model of breast cancer. (A) The breast cancer cell line MDA-MB-231 was grown on a non-matrix transparent cycloolefin resinous scaffold comprising nanoscale indented patterns with a 500-nm line width, 1-mm line depth, and 10-nm to 500-nm line spacing. Cells grown on this surface form 3D multicellular tumor spheroids with projections. (B) The high-throughput screen was performed in 96-well plates in duplicates. The readouts for the screen were morphology and viability. (C, D) Images from 3D culture plates. Ten images were captured per well for a total of 20 images for each siRNA.

Ingenuity Pathway Analysis

We next analyzed the 5.1% of genes that led to no projections in the 3D MCTS system and were growth inhibition in the 2D monolayer system. The top five pathways that were over-represented based on these genes are listed in Table 4. Based on these five top-ranked pathways, we used the "Overlay" feature in IPA to identify various drugs that could perturb the activity of these molecules. Further, we perturbed these molecules using the MAP feature in IPA (Figure S1). To identify genes that synergize with the MEK inhibitor's action and drive reduction in tumor growth morphology and viability, we used the MAP and upstream drug activity analysis functionality in IPA.[13] Examination of the top-ranking pathways in this set of genes revealed that the PI3K gene is present and the upstream of PI3K gene across the five pathways were related (Figure 3, Figure S1). Based on these observations, we hypothesized that a PI3K inhibitor would synergistically reduce tumor growth in conjunction with MEK inhibitor AS703026.

Drug-Combination Effect in TNBC Cell Lines

To validate whether inhibition of the PI3K and MEK pathways has a synergistic anti-proliferation effect in TNBC, we tested 6 different subtypes of TNBC cell lines (BL1, BL2, IM, MSL, M, and LAR), plus an unclassified (UC) cell line, as defined in previous studies [14, 15]. We evaluated the anti-proliferation index of combination treatment using an ATP-based cell viability assay, CellTiter-Blue reagent. When cells were treated with the combination of SAR245409 and AS703026 for 72 h, all tested cell lines showed IC50 values in the 0.3 – 3.0 μ M range. In particular, we observed a significant shift in the IC50

value of both SAR245409 and AS703026 concentration in some BL2- and MSL-subtype cell lines (Table 5, Figure S2). We next used the CalcuSyn program to analyze the fraction of cells affected by combination treatment with a clinically relevant dose (0.5 - 2.0 μM). Combinational index (CI) and fractional index (Fa) results suggested that SAR245409 and AS703026 enhanced the anti-growth effect in TNBC cell lines, specifically in the BL2 and MSL subtypes (Table 6).

Next, to discover whether the combination of SAR245409 and AS703026 enhances the anti-tumorigenic effect in TNBC cells, we performed *in vitro* tumorigenicity assays using soft-agar colony

formation. We observed that when cells were treated with IC50 concentrations (1 μM) of SAR245409 and AS703026, more than 90% of cells did not form colonies (Figure S3). Thus, at these concentrations, it was difficult to evaluate the synergistic anti-tumor effect of combination treatment. Therefore, we selected doses that were one-tenth or one-hundredth lower than the initial concentrations (0.1 μM or 0.01 μM) for treatment. As shown in Figure 4, colony numbers were significantly lower with combination treatment than with single treatment in most tested TNBC cell lines except BT-20 cell line.

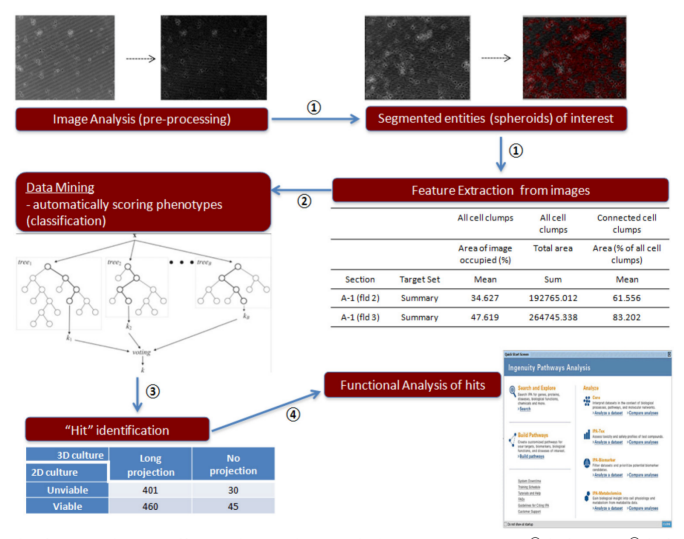


Figure 2. Image analysis procedure of 3D culture siRNA screening outcome. Step 1: Imaging analysis and data extraction, ①; Step 2: Data mining, ②; Step 3: Contingency table, ③; Step 4: Functional analysis of genes, ④

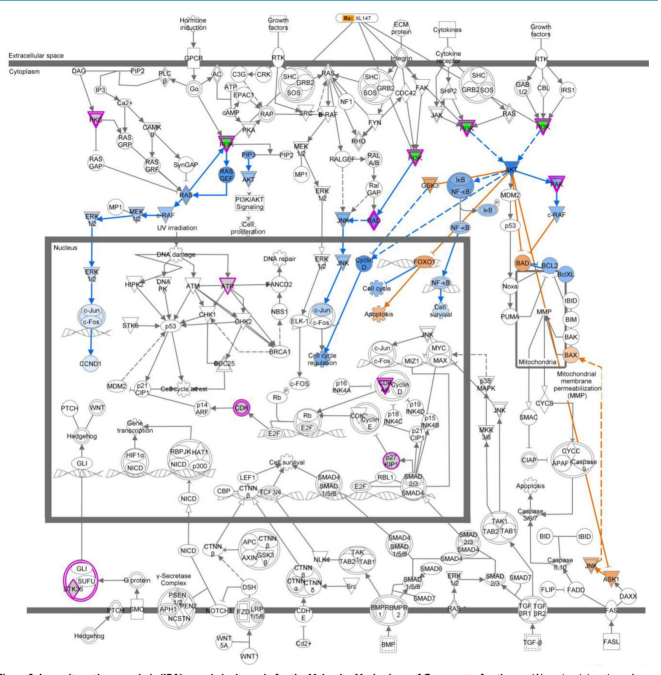


Figure 3. Ingenuity pathway analysis (IPA) revealed a key role for the Molecular Mechanisms of Cancer set of pathways. We analyzed the subset of genes whose knockdown in TNBC cells treated with MEK inhibitor AS703026 led to no projections in the 3D MCTS system and were growth inhibition in the 2D monolayer system (5.1%). Molecular Mechanisms of Cancer was a top-ranking pathway. Output data indicated that PI3K pathway-related genes are the most important target molecules for combination with MEK inhibitor AS703026. Pink color indicated-gene: target gene from 2D/3D siRNA screening analysis. Blue color indicated-gene: negatively regulated downstream molecule by inhibition of pink color indicated-target molecule. Yellow color indicated-gene: possible mechanism on apoptosis by inhibition of target genes from 2D/3D siRNA screening. XL-147: a selective and reversible class I PI3K inhibitor for PI3Kα/δ/γ.

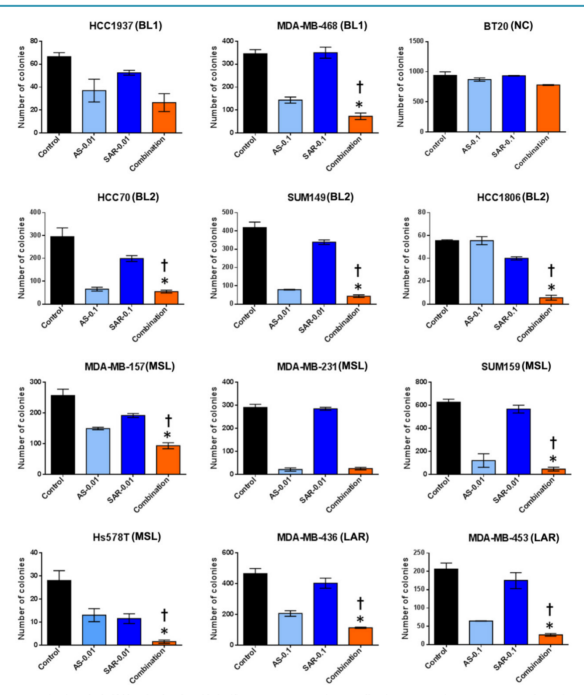


Figure 4. The combination of AS703026 with SAR245409 significantly enhanced antitumor effect in anchorage independent colony formation assay (soft agar assay). Breast cancer cell lines were overlaid onto the bottom agar layer (0.8%) with the indicated drugs (μ M) and incubated for 3-6 weeks. For visualizing, colonies were stained with 200 μ l of MTT (1 mg/ml) solution for 2 h and counted using the GelCount colony-counting system according to the manufacturer's instructions. Statistical significance was evaluated by t test using GraphPad Prism software. Error bars represent the standard deviation of three independent experiments. *, p <0.05 combination compared with SAR245409; †, p <0.01 combination compared with AS703026. BL1: basal-like 1, BL2: basal-like 2, MSL: mesenchymal stem-like, LAR: luminal androgen receptor, UC: unclassified.

Table 4. IPA pathway analysis of the 5.1% of the genes in Table 2 that led to no projection in the 3D MCTS system and a growth inhibition outcome in the 2D monolayer system. The five top-ranked pathways corresponding to the gene hits are listed.

Pathway name	Genes common to the pathway and the list of hits from the 3D screen
Erbb Signaling	PAK4, PAK6, PDPK1, PIK3C2G, PRKCE, PRKCQ, PRKD3
Molecular Mechanisms of Cancer	ATR, CDK6, CDKN1B, PAK4, PAK6, PIK3C2G, PRKCE, PRKCQ, PRKD3, RAC1, STK36
Renin-Angiotensin Signaling	PAK4, PAK6, PI3KC2G, PRKCE, PRKCQ, PRKD3, RAC1
NK Cell Signaling	PAK4, PAK6, PIK3C2G, PRKCE, PRKCQ, PRKD3, RAC1
CCR3 Signaling in Eosinophils	PAK4, PAK6, PIK3C2G, PRKCE, PRKCQ, PRKD3, RAC1

Table 5. IC50 values for single and combinational treatment with AS703026 and SAR245409 in TNBC cells under 2D culture conditions.

		Single treatment		Combinational treatment	
Subtype	TNBC cell line	SAR245409 (IC50,	AS703026 (IC50,	SAR245409:AS703026	
•		μM)	μM)	1:1 mix ratio, (IC50, μM)	
	HCC1937 (BL1)	1.07	> 20	1	
Basal-like	MDA-MB-468 (BL1)	1.98	2.5	1.2	
	HCC1187 (IM)	0.25	1.17	0.3	
	HCC70 (BL2)	0.37	12.7	0.38	
	SUM149 (BL2)	1.7	0.95	0.32 0.9	
	HCC1806 (BL2)	2.2	12.5	0.9	
	HS578T (MSL)	0.5	4.84	0.52	
	MDA-MB-157 (MSL)	6.48	> 20	2.76	
Masanahyimal lika	SUM159 (MSL)	7.3	> 20	2.78	
Mesenchymal-like	MDA-MB-436 (MSL)	0.28	2.7	1	
	MDA-MB-231 (MSL)	1.36	0.6	0.12	
	BT-549 (M)	0.56	> 20	1.2	
Luminal androgen	MDA-MB-453	0.18	6.32	1.04	
receptor	WIDA-WID-455	0.16	0.32	1.04	
Unclassified	BT-20	1.2	9	0.58	

Six different subtypes of TNBC were identified based on gene expression profiles: basal-like 1 and 2 (BL1 and BL2), immunomodulatory (IM), mesenchymal (M), mesenchymal stem-like (MSL), and luminal androgen receptor (LAR) [14].

Table 6. Combinational index (CI), a quantitative measure of the degree of drug interaction, and fractional index (Fa), the fraction of cells affected by the treatment, for the drug combination AS703026 and SAR245409 (1:1 mix ratio).

Cell line and TNBC subtype	CI/Fa values for different concentrations of the drugs (1:1 mix ratio)						
	0.5 μΜ		1.0 μΜ	1.0 μΜ		2.0 μΜ	
	Fa	CI	Fa	CI	Fa	CI	
HCC1937 (BL1)	0.43	0.174	0.45	0.813	0.59	0.724	
MDA-MB-468 (BL1)	0.4	0.795	0.54	0.635	0.68	0.572	
HCC1187 (IM)	0.72	0.405	0.74	0.603	0.81	0.49	
HCC70 (BL2)	0.75	0.302	0.86	0.24	0.94	0.135	
SUM149 (BL2)	0.78	0.079	0.84	0.079	0.89	0.081	
HCC1806 (BL2)	0.51	0.216	0.62	0.263	0.72	0.325	
HS578T (MSL)	0.61	0.517	0.75	0.459	0.87	0.341	
MDA-MB-157 (MSL)	0.23	0.155	0.29	0.233	0.52	0.284	
SUM159 (MSL)	0.32	0.078	0.55	0.076	0.65	0.112	
MDA-MB-436 (MSL)	0.57	0.91	0.84	0.659	0.91	0.595	
MDA-MB-231 (MSL)	0.88	0.025	0.87	0.053	0.96	0.055	
BT-549 (M)	0.51	0.794	0.61	0.899	0.72	1	
MDA-MB-453 (LAR)	0.79	0.44	0.85	0.498	0.92	0.402	
BT-20 (UN)*	0.51	0.317	0.68	0.25	0.82	0.19	

*UN, unstable (those TNBCs that do not fall consistently into one of the six subtypes). CI value: <0.1 indicates very strong synergism; 0.10-0.30, strong synergism; 0.31-0.70, synergism; 0.71-0.85, moderate synergism; 0.86-0.90, slight synergism; 0.91-1.10, nearly additive; 1.11-1.20, slight antagonism; 1.21-1.45, moderate antagonism; 1.46-3.30, antagonism; 3.31-10, strong antagonism; >10, very strong antagonism.

Our Ingenuity pathway analysis revealed that p21-activated kinase (PAK) group II, PAK4 and PAK6, and Rho GTPase RAC1 were the most common genes from the 3D siRNA screening assays. Previous studies indicated that PAK4 and RAC1 are key regulators of not only migration/invasion but also tumorigenicity in breast cancer [16-20]. Therefore, we further tested whether the combination of SAR245409 and AS703026 inhibits TNBC cell migration and invasion. As shown in Figures 5A and 5B, compared with single treatment, combination treatment with SAR245409 and AS703026 (0.5 µM) inhibited both migration and invasion significantly in both TNBC cell lines tested (p < 0.001). PAK and RAC1 have been described as regulatory factors in cellular morphological formation of lamellipodia, membrane ruffling, and filopodia and stress fibers during migration or invasion [21]. We observed that control or single-treatment cells showed well-developed lamellipodia or filopodia, as indicated by the arrows in Figures 5C and 5D; however, combination treatment cells had restricted lamellipodium and filopodium development. These data indicated that combination treatment of SAR245409 and AS703026 inhibited migration/invasion by inhibiting the small GTPase signaling pathway. Also, the conventional assay method successfully reproduced the 3D RNAi screening result.

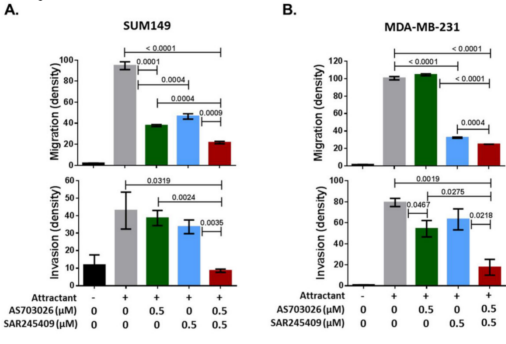
Discussion

In the present study, we used high-throughput 2D and 3D RNAi screening to identify targets whose silencing induced growth inhibition or projection formation of TNBC cells treated with AS703026. Our comprehensive analysis revealed the PI3K pathway as an important target in TNBC cell growth and migration/invasion under AS703026 treatment. We validated that compared with single-agent treatment, the combination of AS703026 and PI3K inhibitor SAR245409 enhanced TNBC cell growth inhibition as

well as inhibition of migration and invasion ability. These data indicate that combination of AS703026 and SAR245409 could be a candidate strategy for inhibiting TNBC cell growth and metastasis.

It has been demonstrated by many studies that 2D monolayer cancer cell culture is a somewhat artificial model and does not fully represent the characteristics of 3D solid tumors [22]. Our group also observed that gene profiling did not perfectly match between TNBC cell line microarray and TNBC patient microarray data using Lehmann and Bauer's analysis algorithm [14, 15]. Li et al reported that most anticancer drugs and molecular targeted therapies have shown promise in preclinical studies; however, cancer

patients have shown a much lower response rate to the drugs compared to *in vitro* chemosensitivity [23]. This low response rate to anti-cancer drugs has been attributed to the tumor microenvironment [24]. The 3D MCTSs are of intermediate complexity between *in vivo* tumors and monolayer cultures, and their growth emulates the heterogeneity of solid tumors; necrosis and radiation-resistant hypoxic regions often occur at the center of solid tumors due to poor vasculature and subsequent lack of oxygen supply [25]. Our unique multicellular spheroid models are an effective 3D cell culture model that takes advantage of the natural tendency of cells to aggregate.



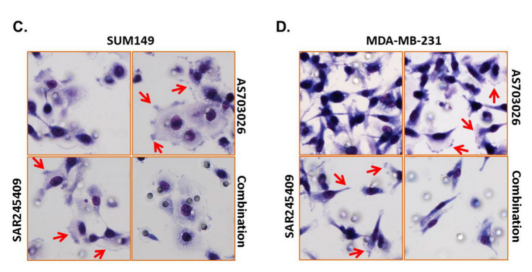


Figure 5. The combination of AS703026 and SAR245409 suppressed SUM149 and MDA-MB-231 cell migration and invasion capability. (A, B) Migration and invasion assay results. P values are indicated between bars; statistical significance was evaluated by t test using GraphPad Prism software. Error bars represent the standard deviation of three independent experiments. (C, D) Images for SUM149 and MDA-MB-231 cells subjected to a migration assay with or without drugs for 6 h. Cells were fixed and stained with Diff-Quik solution. Arrows indicate filopodia or membrane ruffling formations.

The ability of multicellular spheroids to simulate a tumor-like environment, together with the ease by which these models can be applied high-throughput screens [26, 27], has resulted in the use of spheroids to study drug efficacy and penetration as well as 3D cellular interactions [25, 28-36]. For example, several in vitro lines of evidence indicate that the ribosomal S6 kinase (RSK) family promotes head and neck squamous cell carcinoma cell metastasis [37], breast epithelial cell motility [38], and cell adhesion [39]. Since RSKs are activated by the MAPK pathway [40] and large proportion of breast cancer cells show elevated EGFR-MAPK pathway activity [41], RSKs are recognized as a target for inhibiting migration and invasion processing in breast cancer. In contrast to expectations, 2D/3D RNAi screening revealed that the RSK gene is not correlated with TNBC cell projection formation, meaning there was no effect on the cells' migration. We confirmed this result using pan-RSK inhibitor (BI-D1870) with two TNBC cell lines; we did not observe an inhibition effect on migration and invasion ability with this RSK inhibitor (Figure S4). This result indicates that our comprehensive RNAi screening and morphological analysis is able to identify target molecules that might effectively prevent cell growth and migration/invasion in an in vivo setting.

Compared to other breast cancer types, TNBCs have more aggressive behavior that tends to metastasize and survive, resulting in a high tant-recurrence rate [42, 43]. Among the 6 different subtypes of TNBC, the MSL subtype displays unique ontologies related to cell motility, extracellular matrix interaction, inositol phosphate metabolism, EGFR, and ERK1/2 signaling [14]. Owing to prominent EMT and stem cell-like properties, the MSL subtype showed decreased distant-metastasis-free survival compared to other TNBC subtypes [15]. Several articles have also reported that elevated tyrosine kinase PI3K/AKT/mTOR receptor, activated and MAPK/ERK pathways, and mutated PIK3CA are frequently observed in TNBC, specifically in metaplastic breast cancers [44-46]. Of note, the MSL-subtype MDA-MB-231 cells used in the 2D/3D RNAi screening displayed target genes closely related to ErbB and the migration/invasion pathway. We identified PI3K and PAK4/6 pathways as top candidates for combination with AS703026. Current evidence indicates that the PAK4/6 pathway is activated via either the PI3K/AKT or the MAPK/ERK pathway and enhances processes associated with progression of cancer such as proliferation, survival, metastasis, and EMT [47-50]. In the process of EMT, including migration/invasion, PAKs regulate cell polarization, resulting in formation of pseudopodia and membrane ruffles [51]. PAK4 is highly overexpressed in metastatic breast cancer cell lines [52, 53], and our data showed strong inhibition of pseudopodia and membrane ruffles by the AS703026 and SAR245409 combination. These findings suggest that metastatic TNBC with a high PAK4 level may be well suited to a new targeted therapy opportunity with a MEK inhibitor and PI3K inhibitor. Further investigation is needed to translate our result into *in vivo* conditions to determine whether the combination of AS703026 and SAR245409 can suppress tumor growth and metastasis effectively. Such data would provide strong justification for future clinical trials to target metastatic TNBC.

In summary, our results suggest that integrated and multi-system RNAi screening methods and analysis narrow the inherent difference between *in vitro* and *in vivo* conditions, providing powerful high-throughput assays that are able to identify anticancer drugs as single or combinational agents. In testing these methods, we have demonstrated that combining AS703026 and SAR245409 effectively suppressed proliferation and colony formation as well as migration/invasion of TNBC cells. We look forward to advancing the field of therapeutic development with AS703026 and SAR245409 for patients with metastatic TNBC.

Materials and Methods

Therapeutic Agents and Cell Lines

AS703026 and SAR245409 were provided by EMD Serono Research & Development Institute, Inc. and Sanofi Inc., respectively. Human breast cancer cell lines MDA-MB-231, HCC1937, MDA-MB-468, HCC1187, HCC70, BT-549, Hs578T, MDA-MB-157, MDA-MB-436, MDA-MB-231, MDA-MB-453, and BT-20 were purchased from ATCC. Human breast cancer cell lines SUM149 and SUM159 were purchased from Asterand. We authenticated all of the cell lines by genotyping through MD Anderson Cancer Center's Characterized Cell Line Core Facility.

High-Throughput siRNA Screening

High-throughput RNAi screens for target identification and synthetic lethality were performed following stringent standard operating procedures at the RNAi screening facility at MD Anderson Cancer Center.[54-57] MDA-MB-231 cells were grown as either a 3D multicellular tumor spheroid (MCTS) cell culture system. MDA-MB-231 cells demonstrate a unique 3D morphology comprising projections suggestive of epithelial-mesenchymal transition (EMT)-mediated invasion processes that are related to the cell line's spontaneous metastasis capability [11, 12]. Both a 2D

monolayer culture and a 3D MCTS culture of MDA-MB-231 cells were transfected, utilizing the reverse transfection procedure with an siGENOME SMARTpool kinome siRNA library comprising a pool of 4 individual siRNA duplexes targeting the individual kinases (Dharmacon). The 3D MCTS were generated by growing the cells on non-matrix transparent cycloolefin resinous sheets comprising nanoscale indented patterns with a 500-nm line width, 1-mm line depth, and 10-mm to 500-nm line spacing [58]. The screen performed in the 2D system was designed to identify genes that affect tumor viability, while the screen performed in the 3D system was designed to identify genes that affect tumor viability as well as modulate the outgrowth of cellular projections, which reflect cell motility and invasive capability. Briefly, for the screen performed utilizing the 2D monolayer cell culture, 10 µl of 200 nM siRNA was mixed with 10 µl of a lipid (0.07 µl of DharmaFECT 1 reagent in 10 µl Opti-MEM medium) for 20 min in a well of a 384-well plate (Corning) and overlaid with 30 µl of cell suspension (5,000 cells) in antibiotic-free cell line-specific media. The final concentration of siRNA in both models was 40 nM per well. siControl ON-TARGETplus siRNA number 4 (siOTP4, Dharmacon) and siTOX ON-TARGETplus (Dharmacon) were used as the negative and positive transfection controls, respectively, in all experiments. In the case of the 3D spheroids, 20 µl of 200 nM siRNA was mixed with 20 μl of a lipid (0.24 μl of DharmaFECT 1 reagent in 20 µl Opti-MEM medium) (Invitrogen) for 20 min in a well of a 96-well NanoCulture plate (SCIVAX Inc.) and overlaid with 60 µl of cell suspension (10,000 cells) in antibiotic-free cell line-specific media. The screen using the 2D monolayer cell culture was performed in triplicate, while the screen using the 3D spheroid cell culture was performed in duplicate. Forty-eight hours after transfection, MEK inhibitor AS703026 was added at a final concentration of 75 nM (the IC20 value) to one set of transfected plates, and a similar volume of DMSO (> 0.1%) was added to the other set of transfected plates. After 72 h of incubation, the cells growing in 3D spheroids were imaged using an In Cell Analyzer 1000 (GE). All images were analyzed for morphological changes following the creation of study-specific programs utilizing GE In Cell Investigator software. The viability of the cells growing in the 2D monolayer or 3D spheroid cultures was determined using the CellTiter-Glo luminescent cell viability assay (Promega) according to the manufacturer's instructions.

Classification of 3D siRNA Culture Outcome

To analyze 3D siRNA screening result, we followed below four classification steps (summarized in

Figure 2).

Step 1: imaging analysis and data extraction.

Since the task of manually annotating the cells into different phenotypic groups is laborious and time consuming, we examined a machine-learning approach to solve this issue: the image-derived features were used to develop a statistical classifier that can predict an expert-annotated class label. A difference in projection type can be used for imaging classifying system. Field-level images were processed using GE IN Cell Investigator software. Briefly, images were pre-processed to adjust contrast and remove artifacts. Subsequently, segmentation routines were used to segment blobs of cells, followed by measurement of morphological parameters such as area, form factor, perimeter, and diameter. The final data output from the analysis tool was a matrix in which each row represents each field within a well, and the columns represented the various morphological features (area, perimeter, diameter, form factor). A board-certified breast medical oncologist (NTU) annotated a subset of images as "long projection", "no projection", and "dead" phenotypes. Based on this annotation, a classification model was built that can discriminate long-projection and no-projection phenotypes. Using the GE IN Cell Analyzer tool, we measured "cell bodies", representing entities after segmentation, and "cell clumps", representing entities before segmentation. These morphological features were computed for each field in the well through the following nine measurements: 1) total area of cell bodies, 2) mean area of cell bodies, 3) number of cell bodies, 4) mean form factor of cell bodies, 5) mean diameter of cell bodies, 6) mean perimeter of cell bodies, 7) mean % area of image occupied by all cell clumps, 8) total area of cell clumps, and 9) mean area of connected cell clumps (% of all cell clumps).

Step 2: data mining.

To automatically score cells based on 'projection' based phenotypes, we used a random forest classifier. A "random forest" is an ensemble of classifiers obtained by aggregating (bagging) several classification trees. Each data point (represented as an input vector) is classified based on the majority vote gained by that vector across all the "trees" of the forest. Each tree of the forest is "grown" in the following way:

- A bootstrapped sample (with replacement) of the training data is used to grow each tree. The sampling for bootstrapped data selection is done individually at each tree of the forest.
- For an M-dimensional input vector, a random subspace of m (≪ M) dimensions is selected, and the best split on this subspace is used to split the

node. This is done for all nodes of the tree.

Before sampling for aggregating process, one-third of the cases are kept "out of the training bag." These out-of-bag (OOB) data are used to obtain an unbiased estimate of the classification error as trees are added to the forest. They are also used to get estimates of variable importance. Several interesting insights into the data are available using random forest analysis. The variable importance plot yields the are variables that most discriminatory classification under the "ensemble of trees" classifier. This importance is based on two measures: Gini index and decrease in accuracy. The Gini index is an entropy-based criterion that measures the purity of a node in the tree, while the decrease in accuracy simply looks at the relative contribution of each variable to the accuracy of the classifier. For our studies, we used the randomForest package for R. The classifier performance on the individual data was assessed via a confusion matrix.

Step 3: contingency table

Further, a contingency table for the genes was built in the following manner:

- For the 2D setting, we examined the viability of TNBC cells with gene knockdown; we compared cells treated with AS703026 vs. untreated cells. Genes that led to a reduction in viability less than vehicle control were designated "growth inhibition", whereas the rest were designated "no growth inhibition".
- For the 3D setting, using the no-projection vs. long-projection classifier, each gene was designated as a "no-projection" gene if its knockdown led to loss of the projection phenotype. Otherwise, it was designated as a "long-projection" gene.

Using definitions from (i) and (ii) above, we populated a contingency table across the 2D (viability) and 3D (no-projection phenotype) settings.

Step 4: Functional Analysis of Genes

Network and functional analysis of the genes inducing the no-projection phenotype was carried out using the Ingenuity Pathway Analysis tool. The tool provides output comprising the top-ranking pathways involving these genes (based on a Fisher exact test). We explored both "Direct and Indirect Interactions" using "Experimentally Verified Predictions" in the Ingenuity Knowledge Base (Genes and Endogenous Chemicals).

Ingenuity Pathway Analysis (IPA)

IPA was used to derive functional and pathway information from the list of genes ("hits") associated

with the growth inhibition and no-projection phenotypes. Further, IPA was also used for network analysis of drug-induced perturbation of pathway activity. More details are in the Supplementary Information. To identify targetable genes that synergize with MEK inhibitor action to drive reduction in tumor growth morphology and viability, we used the Molecule Activity Predictor (MAP) and upstream drug activity analysis functionality in IPA [13]. Briefly, based on the top-ranked pathways corresponding to the gene hits, we used the "Overlay" feature in IPA to overlay various drugs that could perturb the activity of these molecules. Further, we perturbed these molecules using the MAP feature in IPA.

In Vitro Cell Proliferation Assay and Analysis of Combinational Drug Effect

Cells were treated with the drug combination and analyzed using a CellTiter-Blue assay (Promega) according to the manufacturer's instructions. Drug synergy was determined by the isobologram and combination-index methods using CalcuSyn software 2.1 (Biosoft) [59]. Detailed methods are described in the Supplementary Information.

Anchorage independent colony formation assay (Soft Agar Assay)

TNBC cells (2×10^3 cells/well) were resuspended in 2 ml of 0.4% agarose solution in DMEM/F12 or F12 medium (Sigma) and overlaid onto the bottom agar layer (0.8%) in 6-well plates. The plates were incubated for 3 to 6 weeks with the two drugs, and colonies were stained with 200 μ l of MTT solution (Sigma, 1 mg/ml) for 2 h. The number of stained colonies greater than 80 μ m in diameter was counted using the GelCount colony-counting system (Oxford Optronix) according to the manufacturer's instructions. Means and standard deviations were calculated on the basis of the colony counts from triplicate wells.

Migration and Invasion Assay

Migration assays were performed in triplicate using a chemotaxis chamber (24-well inserts, 8- μ m pore size, BD Biosciences). Cells were harvested and resuspended in DMEM/F12 media with 1 mg/ml bovine serum albumin (Sigma). 1×10⁵ cells/350 μ l were plated on upper chambers, and bottom wells were filled with 750 μ l of DMEM/F12 media that included 2% FBS and 100 ng/ml EGF (Invitrogen) as an attractant to induce migration or invasion. Six hours later, membranes were fixed and stained with Diff-Quik solutions (Dade Behring, Inc.). Chemotaxis was quantitated by the ImageJ program. For invasion, 50 μ l of 10% Matrigel solution was coated on the up-

per chamber for 1 h at 37°C. Cells were allowed to invade for 24 h.

Statistical Analysis

For experimental outcomes, descriptive statistics (mean and standard deviation) were summarized for each group. An analysis of variance model was used to compare the mean outcome values among the tested groups. Statistical analyses were performed using an unpaired *t*-test with Prism version 5 (GraphPad Software). *P* values of <0.05 were considered statistically significant.

Supplementary Material

Figures S1-S4. http://www.jcancer.org/v06p1306s1.pdf

Abbreviations

TNBC: Triple-negative breast cancer; 2D: two-dimensional; 3D: three-dimensional; MEK: mitogen-activated protein kinase kinase; HER2: human epidermal growth factor receptor 2; BL1: basal-like 1; BL2: basal-like 2; IM: immunomodulatory; M: mesenchymal; MSL: mesenchymal stem-like; LAR: luminal androgen receptor; RNAi: RNA interference; MCTS: multicellular tumor spheroid; EMT: epithelial-mesenchymal transition; MAP: Molecule Activity Predictor; IPA: Ingenuity Pathway Analysis; PAK: p21-activated kinase; Rac 1: Ras-related C3 botulinum toxin substrate 1; RSK: ribosomal S6 kinase.

Acknowledgments

Naoto Ueno was supported by EMD Serono Research & Development Institute, Inc., the Morgan Welch Inflammatory Breast Cancer Research Program, the State of Texas Rare and Aggressive Breast Cancer Research Program, and the National Institutes of Health (CA123318 and MD Anderson's Cancer Center Support Grant [CCSG], CA016672 [the Characterized Cell Line Core Facility and RNAi Screening Service]). Arvind Rao was supported by the National Institutes of Health (CCSG CA016672 [Bioinformatics Shared Resource] and 5R00CA139006 [PI: Chandra Bartholomeusz]), the Cancer Prevention and Research Institute of Texas High-Throughput Screening Program grant (RP110532), an MD Anderson Institutional Research Grant, and MD Anderson start-up funds. Chandra Bartholomeusz was supported by the National Institutes of Health (R00CA139006) and MD Anderson start-up funds. Geoffrey Bartholomeusz was supported by the Cancer Prevention and Research Institute of Texas High-Throughput Screening Program (RP110532-P3) and the National Institutes of Health (MDACC/CCSG CA016672 PI: Ronald DePinho). We are grateful to Tony Zucca and Robert

Graves of GE Healthcare, who provided help with building the image analysis protocol in IN Cell Investigator. We thank Sunita Patterson of the Department of Scientific Publications at MD Anderson for editorial assistance.

Competing Interests

Samantha Goodstal is an employee of EMD Serono Research & Development Institute Inc. All other authors have no conflicts of interest to disclose.

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